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Selective catecholamine detection in living cells by a copper-mediated oxidative bond cleavage†

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The development of a new triggered-release system for selective detection of catecholamines in biological samples including living cells is reported. Catecholamines are a class of tightly regulated hormones and neurotransmitters in the human body and their dysregulation is implicated in various neurodegenerative diseases. It is highly challenging to selectively sense and detect catecholamines in a complex biological environment due to their small size, non-specific molecular shape and trivial chemical properties. In this study, a copper-based, catecholamine-triggered oxidation that releases a fluorescent reporter is described. The probe is highly sensitive and selective for detecting changes in catecholamine levels in aqueous buffer, human plasma, and cellular models of neuronal differentiation and Parkinson's disease. This new catecholamine sensing strategy features chemical reactivity as part of small molecule recognition as opposed to the conventional use of a well-designed host for reversible binding.

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Introduction

Catecholamines (*e.g.* dopamine, adrenaline and noradrenaline) are important small molecule neurotransmitters involved in vital physiological processes such as learning and memory, blood pressure regulation, motor control and physical responses.^{1,2} Abnormal catecholamine levels are associated with various neurological disorders such as schizophrenia,³ psychosis,⁴ and attention deficit hyperactivity disorder.⁵ At the cellular level, catecholamine dynamics is tightly regulated to maintain proper cellular functions. In neurons, for example, dopamine is stored in synaptic vesicles and released into the synaptic cleft when an action potential is reached, and the released dopamine could be taken up again and repackaged for further release.⁶ In particular, dopamine depletion in dopaminergic neurons in the mid-brain is attributed to motor deficits which are the main symptoms of Parkinson's disease.⁷ Recognition and detection of catecholamine in cells are therefore critical to track their dynamics for a further understanding of their roles in neuronal development and related diseases, as well as developing diagnostic and therapeutic strategies for catecholamine-related disorders.⁸

Development of selective recognition and detection for catecholamines in biological cells or other biological samples is far from trivial because catecholamines have only minimal structural features and diol/amine functional groups are widely present in other biomolecules. The small size and neutral charge of catecholamines also add extra complexity in designing recognition systems in aqueous environments. Although there are a few sophisticated synthetic hosts that show promising catecholamine binding, they are not biocompatible, sensitive or selective for biological applications.^{9–14} Methods to engineer an easily detectable response from these hosts are also not straightforward. Very recently, several (bio)macromolecular hosts that sense catecholamines by a conformational change of the host upon catecholamine binding have been reported.^{15–17} For example, protein-based sensors have been constructed by fusing a known catecholamine-binding protein with a fluorescent protein.^{15,16} The genetically engineered proteins offer a high selectivity and strong binding to catecholamines but their use may require cell transfection or implantation of genetically modified cells. A few systems based on DNA-wrapped carbon nanotubes or boronic acid-functionalized conjugated polymers have also been reported.^{18–21} Compared with macromolecular sensors, small molecule systems for fluorescent catecholamine detection are operationally simple and applicable in different types of biological samples with minimum preparative procedures. Rational chemical modifications on small molecule probes also offer a facile tuning of photophysical, localization, cell-permeation and other properties to suit the specific need of the experiment. Examples of small molecule probes for direct catecholamine sensing are however scarce, with one example featuring reversible iminium and/or boronate ester formation

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between dopamine/noradrenaline and an aldehyde- and/or a boronic acid-containing probe.^{22–24} The small association constant ($K \sim 10^2$ to 10^4 M⁻¹) of iminium and boronic acid catechol ester may also limit its application to samples with a high catecholamine concentration. Other bioanalytical methods such as liquid chromatography,^{25–27} electrochemical methods^{28,29} and the use of fluorescent false neurotransmitters^{30–34} are either not applicable in living cells or only indirectly studying neuronal activity.

In view of the distinctive difficulty and importance of catecholamine detection, we sought to develop a new strategy for the selective recognition and fluorescence detection of the small molecules, which not only will provide new optical tools for studying the neurotransmitters in different samples and conditions, but also could be versatily applied to other catecholamine-responsive systems for potential controlled-release, triggered response and other applications. Because catecholamine binding *via* reversible boronic acid catechol ester and/or C=N exchange is inherently weak,^{35,36} we turned our attention to include chemical reactivity as part of the recognition.^{37,38} In particular, we are inspired by the oxidative reactivity of dopamine β -hydroxylase (D β H), which is a copper oxygenase featuring a His/His/Met coordination, and reason that a similar catecholamine-triggered, copper-based oxidation can be harnessed for the selective release of a caged fluorescent reporter (Scheme 1).^{39–43} Here we report the design and development of a prototypical catecholamine probe (CAP) that is based on a copper-based oxidative reaction and demonstrate that it can successfully detect changes in catecholamine levels in PC12 models of neuronal differentiation and Parkinson's

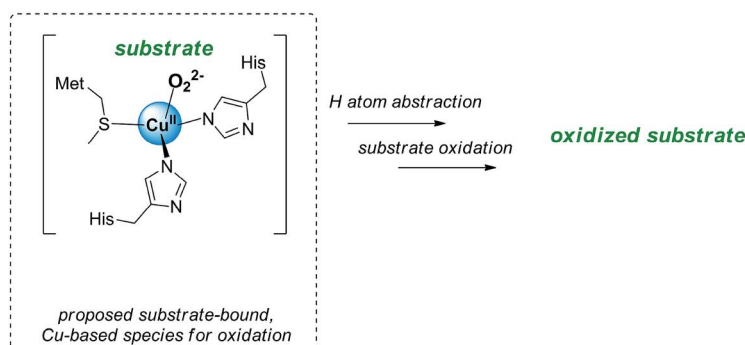
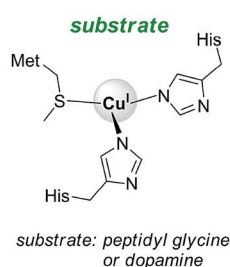
disease. Combined with morphological assessment of the cells in the same imaging experiment, an integrative study of important neuronal processes by morphology and molecular imaging is demonstrated for the first time. On the other hand, connecting the chemical reactivity of a small molecule to that of metal-based oxidation is demonstrated to be an effective strategy for the selective detection of small organic molecules that are challenging recognition targets using conventional host-guest chemistry.

Results and discussion

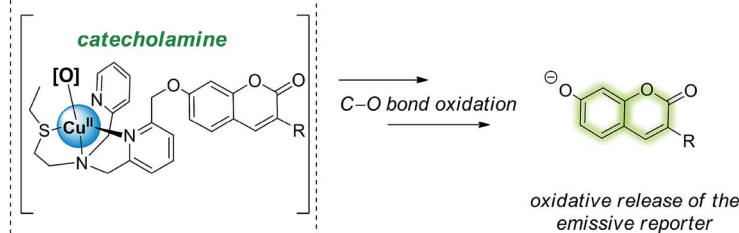
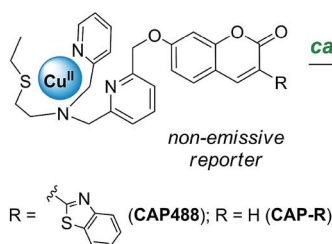
Design, synthesis and initial evaluation of CAP

Our strategy to develop a catecholamine-triggered oxidative release of a caged reporter was inspired by the oxidation (hydroxylation) of dopamine to noradrenaline by dopamine β -hydroxylase (D β H). D β H is a copper monooxygenase with a copper co-factor coordinated by two histidine residues and one methionine residue.^{44,45} Detailed structural, spectroscopic and kinetic studies on D β H and the related peptidylglycine α -hydroxylating monooxygenase (PHM) suggest that substrate oxidation involves hydrogen atom abstraction from the substrate by a copper-peroxo species generated from dioxygenation of the copper and substrate-mediated electron transfer (Scheme S1†).⁴⁶ If a synthetic copper complex in a similar coordination environment could trigger an oxidation upon reaction with catecholamines, a caged reporter could be released to produce a luminescence response. A copper(II) complex supported by two sp² nitrogen and one thioether donor that mimics the His/His/Met coordination in D β H was therefore

suggested mechanism for D β H and PHM



bioinspired design of CAP



Scheme 1 The His–His–Met copper coordination in the mono-copper active site of dopamine β -hydroxylase (D β H) and related enzymes featuring a proposed substrate-bound, Cu^{II}-peroxo species for substrate oxidation inspires the design of the catecholamine probe (CAP) that consists of a non-luminescent reporter caged by an N₃S–Cu^{II} complex, in which dopamine (and catecholamine) binding will trigger O₂ activation at the copper for the oxidative release of the reporter in its luminescent form for catecholamine sensing.



conjugated to 3-(2-benzothiazolyl)-7-hydroxycoumarin ($\lambda_{em} = 488$ nm) to give **CAP488**. When caged as an ether, the coumarin is only weakly fluorescent, but a C–O bond cleavage will release the fluorophore in its phenol/phenolate form to produce a fluorescence turn-on (Scheme 1 and Fig. S1†).^{43,47} It is worth noting that related O₂ dependent oxidations based on iron, cobalt and copper have been developed as selective fluorescent probes for the metals and reactive oxygen species, and have been shown to be useful imaging tools for many different biological samples including live cells, tissues and animals.^{48–51} The use of an organic molecule to trigger such a metal-based oxidation for sensing is however not known.

CAP488 was obtained from coordinating a Cu²⁺ ion to the corresponding ligand–fluorophore conjugate that was synthesized in three steps from commercially available materials (Scheme S2†). Due to the modular probe design, variants of **CAP488** of other properties, for example the ratiometric **CAP-R**, can be easily obtained by simply linking the ligand to other reporters. An X-ray structure of **CAP-R** prepared from CuCl₂ shows the expected N₃S–Cu^{II} coordination with a fifth chloride ligand (Fig. S83†).⁵² Initial evaluation of **CAP488** in PBS buffer (pH 7.4) showed a significant 70-fold fluorescence enhancement after treating with 1 eq. of dopamine for 1 h. A *ca.* 15-fold enhancement of the intensity ratio I_{455}/I_{385} was also observed when **CAP-R** was treated under the same conditions (Fig. 2b). When a pyridine donor in **CAP488** is replaced by a phenyl ring (N₂S-cyan) or a quinoline (N_{3q}S-cyan), or the thioether is replaced by a pyridine (N₄-cyan), the fluorescence response towards dopamine was significantly reduced or became negligible (Fig. 1), showing that the observed dopamine reactivity of **CAP488** is specific to the N₃S-coordinated Cu²⁺.⁵³ In addition, similar fluorescence responses were observed when **CAP488** and **CAP-R** were treated with adrenaline and noradrenaline (Fig. S2†), showing that the N₃S–Cu^{II} complex could serve as a general catecholamine-responsive trigger.

Catecholamine reactivity, fluorescence response and selectivity

The fluorescence response of **CAP488** was evaluated in more detail. As discussed above, **CAP488** (10 μ M, PBS, pH 7.4) showed

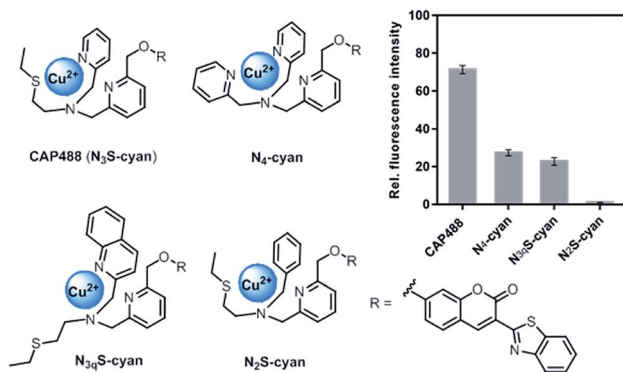


Fig. 1 Structures and fluorescence response towards 1 eq. of dopamine of Cu(II) complex–fluorophore conjugates with combinations of N,S-donors.

a 70-fold emission enhancement centered at 488 nm after a 1 h reaction with dopamine (Fig. 2). Adrenaline (84-fold) and noradrenaline (72-fold) also gave comparable fluorescence turn-on. The counter anion, which probably is dissociated from the Cu²⁺ in solution, does not affect the catecholamine response and similar fluorescence enhancements were observed when **CAP488** was prepared using different copper(II) salts (Fig. S4†).⁵⁴ The observed fluorescence turn-on was due to the proposed C–O bond cleavage and the uncaged fluorophore, ligand fragments and other oxidized products could be identified by LCMS and ESI-MS analysis of the solution mixtures after the reactions (Fig. S17–S19†). The yield of the released coumarin from **CAP488** by dopamine was *ca.* 5% as determined by the LC analysis. Albeit being a seemingly low cleavage yield, the emission intensity at 488 nm of 0.05 μ M 3-(2-benzothiazolyl)-7-hydroxycoumarin is indeed found to be about 70-fold stronger than that of a 10 μ M **CAP488** solution, consistent with the observed 70-fold fluorescence enhancement for dopamine (Fig. S5†).

The fluorescence response of **CAP488** is selective to catecholamines. In addition to dopamine, adrenaline and noradrenaline, L-DOPA, a precursor to dopamine, also showed a turn-on response. On the other hand, other biologically

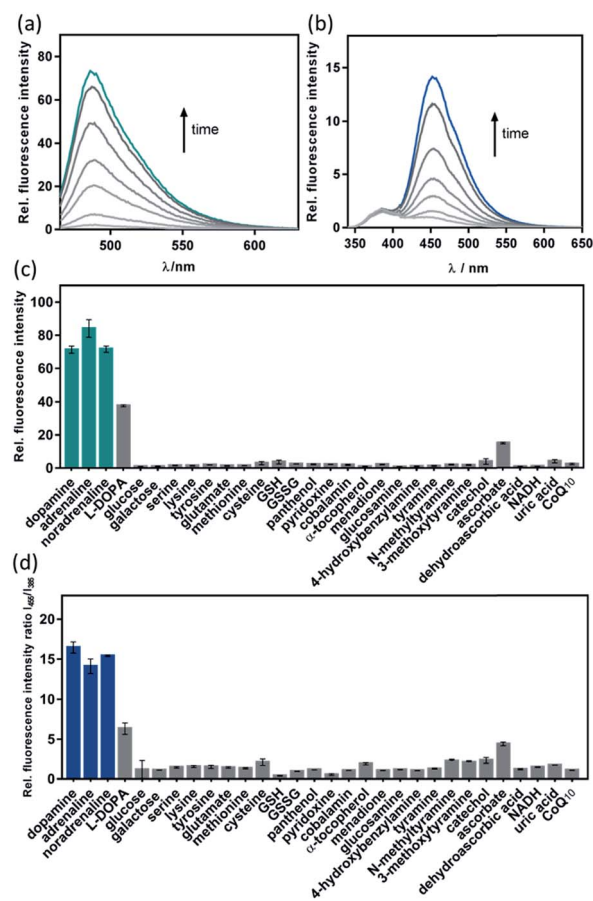


Fig. 2 Time-dependent fluorescence response (0, 1, 3, 5, 10, 30 and 60 min) of 10 μ M (a) **CAP488** and (b) **CAP-R** towards 1 eq. of dopamine; and selectivity of (c) **CAP488** and (d) **CAP-R** against other biomolecules. Error bars are \pm SD ($n = 3$).



relevant diols (*e.g.* sugars), amines (*e.g.* amino acids), hydroquinones/quinones (*e.g.* α -tocopherol, menadione or CoQ₁₀) and glucosamine did not result in significant fluorescence responses (Fig. 2c). Reactions of **CAP488** with catechol, 4-hydroxybenzylamine, tyramine, *N*-methyltyramine and 3-methoxytyramine or dopamine derivatives that are masked as silyl ethers and/or trimethylammonium also resulted in a low to negligible fluorescence response, suggesting that both the catechol and amine functional groups in catecholamines are essential for the reaction (Fig. 2c and S6†). Of note, developing an activity-based recognition that targets both amine and diol while avoiding cross reactivity is non-trivial because these functional groups are also widely present in a biological matrix. Because of the necessity of the C–O bond cleavage for the fluorescence response, **CAP** differentiates catecholamines from other competitive species that react with the N₃S–Cu^{II} complex differently.^{55–59} In fact, synthetic copper complexes supported by ligands with primary N₃S donors have been intensively studied as models of D β H and PHM.^{52,60,61} Dioxygenation of the corresponding Cu^I complexes has been shown to generate a copper-oxygen species (*e.g.* superoxo and peroxy) that is reactive towards the oxidation of model substrates. It has also been demonstrated that the weak thioether donor can raise the Cu^{III/I} reduction potential to result in a more electrophilic copper-superoxo species for better hydrogen atom abstraction.^{52b} On the other hand, other studies have demonstrated that the stability and reactivity of a copper-oxygen species are also affected by interactions in the secondary coordination sphere.^{52c} A correct combination of the ligand donors (in both the first and second coordination sphere), terminal oxidant and substrate is therefore critical for useful copper-based oxidations. Different from the bioinorganic model studies or previous ROS/metal sensing in which the metal-oxygen oxidant is generated directly from oxygenation or ROS coordination,^{39,41–43} **CAP488** is not reactive towards O₂ until being activated by catecholamines, therefore representing a new way to activate copper-based, O₂-dependent oxidation.

With the strong catecholamine-dependent fluorescence enhancement, **CAP488** was further studied in a real biological matrix. Linear fluorescence responses to the three catecholamines in both aqueous buffer and human plasma solutions were observed, showing that the probe could be applied in real biological samples (Fig. S7–S9†). Detection limits of **CAP488** for the three catecholamines are around 0.2–0.5 μ M as determined by the 3 σ method. The fluorescence response has a reasonable kinetics in which the fluorescence signal is saturated under 30 min, which is of a timescale similar to those of other activity-based probes that have been successfully applied in cell and animal imaging.^{62,63} Some cellular processes with a change in intracellular catecholamine levels such as neuron differentiation also take place in a much longer time frame,^{64,65} suggesting that the probe could be applied in these biological processes.

Characterization of the copper-based oxidation

Before further evaluation in biological systems, the oxidative bond cleavage was studied in more detail. First, O₂ was found to be the

terminal oxidant, but not other ROS, for the oxidative cleavage of **CAP488**. No fluorescence enhancement was observed when **CAP488** was treated with the catecholamines under anaerobic conditions (Fig. S10†). In addition, reaction of **CAP488** with H₂O₂, O₂[–] and other ROS also gave no significant fluorescence turn-on (Fig. S11†), and that control reactions of **CAP488** and the catecholamines in the presence of catalase or superoxide dismutase showed that the fluorescence responses were not affected (Fig. S12†). The reaction of **CAP488** with the catecholamines also produced no detectable amount of ROS as revealed by LCMS analysis of the **CAP488**/catecholamine reaction mixtures in the presence of a H₂O₂ probe,⁶⁶ an O₂[–] probe³⁹ or a general ROS probe (Fig. S20–S22†).⁶⁷ The ROS probes were found to remain unreacted along with the release of the coumarin and other oxidation products. Indeed, a computational study has shown that ROS generation from a Fenton-like reaction involving copper is significantly inhibited in the presence of catecholamines.⁶⁸ The presence of common transition metal ions like Fe²⁺, Fe³⁺ and Zn²⁺ also showed no effect on the catecholamine-induced fluorescence response (Fig. S13†). All these observations are consistent with a copper-based oxidative bond cleavage. Furthermore, HR-ESI-MS analysis of an immediate mixture of **CAP488** and dopamine showed two new peaks at $m/z = 881.1278$ and 657.1074 that are assignable to a **CAP488**-dopamine adduct and the one-electron reduced form of the probe respectively (Fig. S23–S27†). Similar species were also observed when adrenaline or noradrenaline was used (Fig. S28–S31†).⁶⁹ These results suggest that catecholamine binding may lead to a reduction of **CAP488**, and subsequent dioxygenation would lead to the formation of a copper-based oxidant for the bond cleavage. Indeed, aqueous Cu²⁺ is known to oxidize adrenaline to adrenochrome with itself being reduced,⁷⁰ and adrenochrome can also be identified from the reaction of **CAP488** and adrenaline (Fig. S18†). This oxidative cyclization is also consistent with the observation that both the catechol and amine groups in the catecholamines are necessary for inducing the bond cleavage. Interestingly, dioxygenation of the Cu(I) analogue of **CAP488** did not lead to a significant fluorescence turn-on (Fig. S14†). One possible reason could be the further participation of the catecholamine after the initial electron transfer. In fact, CV studies showed that the direct **CAP488** reduction by dopamine may not be thermodynamically feasible (Fig. S34†). Preliminary kinetic studies also showed a fractional reaction order for the catecholamines (Fig. S35, and Tables S1 and S2†).⁷¹ Since the catecholamine-induced cleavage showed a strong solvent dependence (Fig. S15 and S33†), stabilization and characterization of the reactive copper-based oxidant at low temperature are not trivial.⁵² Nevertheless, it is clear that the C–O bond cleavage in **CAP488** involves a copper-based oxidation that is triggered by catecholamines and further mechanistic studies are warranted.

Imaging of dopamine accumulation in a neuronal differentiation model using **CAP488**

Given the promising analytical performance of **CAP488**, it was further employed in live cell fluorescence imaging. First, only negligible emission was observed when a lysate solution of HeLa cells was treated with **CAP488**, showing that common



intracellular components do not have a significant effect on the bond cleavage (Fig. S15[†]). A cellular model of neuron differentiation using adrenal pheochromocytoma PC12 cells was then studied.^{72–74} Upon stimulation with nerve growth factor (NGF), PC12 differentiates and resembles the phenotype of sympathetic ganglion neurons with an increase in tyrosine hydroxylase (TH) activity and dopamine biosynthesis.^{75–79} While the differentiation process is conventionally assessed by monitoring the cell morphology (*e.g.* length and number of neurites),^{64,65,80} changes in the intracellular molecular content have only been separately determined by HPLC or electrochemical methods on cell lysates.^{73,78,81,82} Molecular imaging of PC12 cells using **CAP488** will therefore establish a direct correlation of the dopamine dynamics with the NGF-stimulated differentiation process. As expected, significant neurite outgrowth was observed upon stimulating the cells with NGF, which is a characteristic morphological change of neuron differentiation of PC12 (Fig. 3g).⁷⁸ By staining the cells with **CAP488**, a significant increase in the intracellular fluorescence was also observed, which is consistent with the previously reported dopamine accumulation due to an increased TH activity (Fig. 3c). In addition, blocking the NGF receptor TrkA with the inhibitor K252a, or treating the undifferentiated cells with only K252a showed no significant fluorescence enhancement,^{83,84} further confirming that the observed fluorescence enhancement in NGF-treated cells was due to an increase of intracellular catecholamines (Fig. 3b). Furthermore, LC/MS/MS analysis of cell lysates from PC12 cells that were stimulated with NGF under the same conditions also showed a significant increase in the dopamine level and no detectable amount of adrenaline or noradrenaline, further confirming that the fluorescence increase in NGF-stimulated PC12 cells observed using **CAP488** is due to dopamine accumulation in the cells (Fig. S36 and S37[†]). Altogether, these results demonstrate that **CAP488** is a new imaging tool for the direct catecholamine imaging in cells. While matching cell morphology to different stages of

neuronal development may require extensive technical experience, fluorescence imaging by measuring emission intensity from a fluorescent probe could be more straightforward that provides additional information at the molecular level.

CAP488 is non-cytotoxic under the imaging conditions. Nuclear staining with Hoechst 33342 and MTT assays showed that the cells were viable throughout the experiment (Fig. S38[†]). Due to the strong binding of Cu^{2+} to the N_3S ligand ($\log K \approx 16$),⁸⁵ the Cu^{2+} complex will remain intact and impose no significant cytotoxicity similar to many other copper-based bioimaging probes.^{86–89} Our fluorescence studies also showed that no detectable amount of free ROS would be generated (see above), further demonstrating the biocompatibility of **CAP488**.

Imaging of dopamine depletion in a cellular Parkinson's disease model using **CAP488**

CAP488 was also examined in a cellular model of Parkinson's disease (PD), which is a major neurological disorder characterized by death of dopaminergic neurons in the midbrain.⁹⁰ Previously, the decrease in dopamine levels in PD has been characterized by techniques involving tracers that mimic the uptake and retention of dopamine.^{91–93} To directly study changes of intracellular dopamine in a PD model, **CAP488** was first differentiated with NGF and then challenged with 1-methyl-4-phenylpyridinium (MPP^+), which is a common neurotoxin for generating cellular and animal models of PD.^{94,95} The neurotoxicity of MPP^+ is multifold and includes dopamine redistribution and efflux.⁹⁶ As expected, a change in the cell morphology and integrity of the neurites was observed, consistent with the expected neurotoxicity of MPP^+ .⁹⁶ Fluorescence imaging with **CAP488** also showed a significant decrease in the intracellular fluorescence in MPP^+ -treated cells, consistent with the expected depletion of the neurotransmitter as a neurotoxic effect of MPP^+ (Fig. 4d). Moreover, MPP^+ -treatment on undifferentiated cells under the same conditions resulted in no statistically different fluorescence as compared with that of

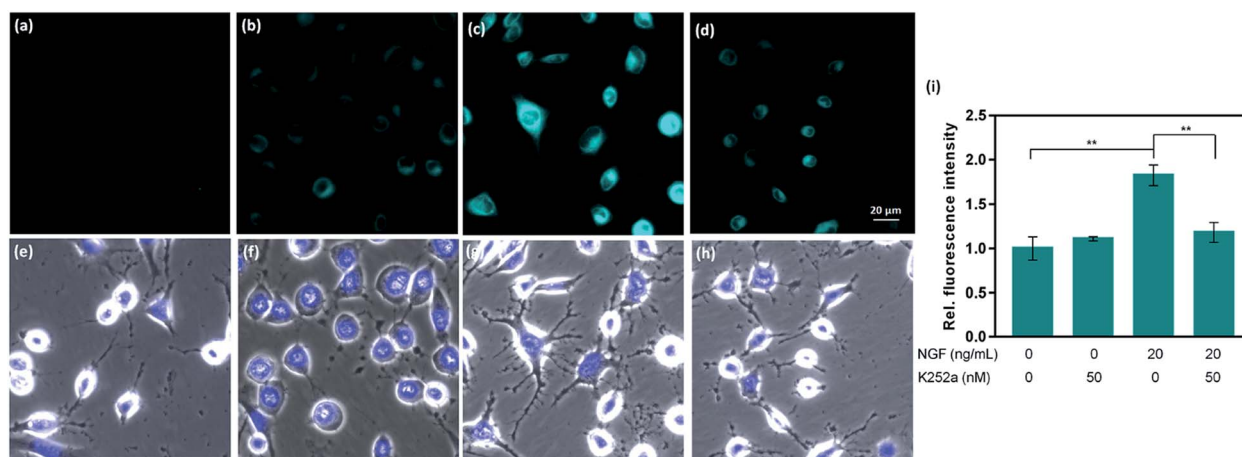


Fig. 3 Representative fluorescent images of PC12 cells stimulated with NGF and stained with **CAP488**. (a) Control cells; and cells treated with (b) 50 nM K252a; (c) 20 ng ml⁻¹ NGF; (d) 20 ng ml⁻¹ NGF and 50 nM K252a; (e–h) overlays of bright field and Hoechst 33342 in (a–d); (i) average fluorescence intensity of PC12 cells treated under conditions in (a)–(d) measured across the entire field from triplicate experiments. Scale bar = 20 μm . Error bars are $\pm\text{SD}$ ($n = 3$). Statistical analysis was performed with one-way ANOVA followed by the Tukey test. **: $p < 0.01$.



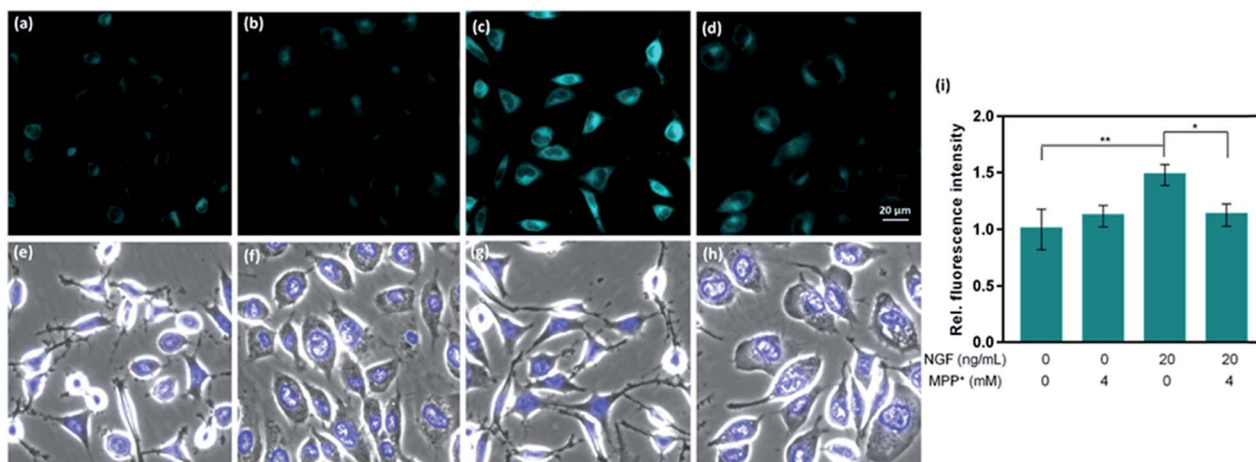


Fig. 4 Representative fluorescence images of PC12 cells treated with NGF/MPP⁺ and stained with CAP488. (a) Control cells; and cells treated with (b) 4 mM MPP⁺; (c) 20 ng ml⁻¹ NGF; (d) 20 ng ml⁻¹ NGF and then 4 mM MPP⁺; (e–h) overlays of bright field and Hoechst 33342 in (a–d); (i) average fluorescence intensity of PC12 cells treated under conditions in (a)–(d) measured across the entire field from triplicate experiments. Scale bar = 20 μ m. Error bars are \pm SD ($n = 3$). Statistical analysis was performed with one-way ANOVA followed by the Tukey test. *: $p < 0.05$; **: $p < 0.01$.

control cells, consistent with the selective vulnerability of dopaminergic neurons to MPP⁺ (Fig. 4b).⁹⁶ Again, these experiments have established CAP488 as a convenient and effective imaging tool for the direct evaluation of dopamine content in a cellular PD model, and reinforced that the complementary use of the molecular and morphological imaging can be an integrative approach for studying neuronal processes.

Conclusions

In summary, the bioinspired development of CAP488 as a first generation of activity-based catecholamine fluorescent probes is described. The probe features a catecholamine-triggered, copper-based oxidative cleavage to release a fluorescent reporter with high sensitivity and selectivity. The combination of small molecule reactivity and metal-based oxidation has significantly expanded the available chemical reactivity for designing effective activity-based recognition and detection systems for challenging targets. This cleavage-based approach is highly modular and the catecholamine-responsive N₃S-Cu²⁺ complex can be installed on other reporters to give other catecholamine probes (e.g. CAP-R) of desired analytical, photo-physical or biological properties. In addition to aqueous buffer and plasma, CAP488 can also be applied as an imaging agent in live cells for the direct visualization of catecholamine content in models of neuronal differentiation and Parkinson's disease, showing for the first time that molecular and morphology imaging can be performed in a single imaging experiment to complementarily correlate changes of intracellular catecholamine to different stages of neuronal development. Detailed mechanistic studies on the catecholamine-triggered, copper-based oxidation and further development of catecholamine probes with improved selectivity, sensitivity, reaction kinetics and tailored bioanalytical properties for specific applications are underway.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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