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Bioorthogonally activated probes for precise fluorescence imaging

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Over the past two decades, bioorthogonal chemistry has undergone a remarkable development, challenging traditional assumptions in biology and medicine. Recent advancements in the design of probes tailored for bioorthogonal applications have met the increasing demand for precise imaging, facilitating the exploration of complex biological systems. These state-of-the-art probes enable highly sensitive, low background, *in situ* imaging of biological species and events within live organisms, achieving resolutions comparable to the size of the biomolecule under investigation. This review provides a comprehensive examination of various categories of bioorthogonally activated *in situ* fluorescent labels. It highlights the intricate design and benefits of bioorthogonal chemistry for precise *in situ* imaging, while also discussing future prospects in this rapidly evolving field.

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1. Introduction

Molecular bioimaging^{1,2} using advanced imaging techniques, such as fluorescence imaging,^{3,4} Raman imaging,^{5–7} magnetic resonance imaging (MRI),^{8,9} positron emission tomography (PET),¹⁰ photoacoustic imaging,¹¹ and single-photon emission computed tomography (SPECT),¹² profoundly impacts biomedical research, disease diagnosis and treatment monitoring, and clinical diagnostics. Among these methods, fluorescence imaging is particularly interesting because it offers us the ability to quantitatively visualize and understand molecular



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processes in living systems from cellular organelles to entire organisms with high spatiotemporal resolution. The development of fluorescent probes,³ pivotal tools for fluorescence imaging, plays a key role in advancing fluorescence imaging techniques, enabling a deeper understanding of biological processes. Over recent decades, many fluorescent probes have been successfully developed for detecting and tracking cellular dynamics, molecular interactions, and tissue architecture.

Traditional fluorescent probes,^{3,13} involving the direct use of emissive dyes as well as dye conjugated biomolecules, such as antibodies, have been widely adopted for staining the cells and tissues for fluorescence imaging.¹⁴ These probes, while effective, often encounter challenges associated with nonspecific binding within cellular and tissue environments, resulting in unexpected signals with compromised accuracy and specificity in fluorescence analysis and imaging. The ability to differentiate genuine signals from nonspecific noise is crucial for

acquiring reliable and meaningful information in fluorescence imaging, particularly within the complex context of biological systems.¹⁵ Photobleaching of the fluorophores represents another major concern in conventional fluorescence imaging.¹⁶ Prolonged exposure to light excitation causes irreversible degradation of fluorophores, restricting the duration of imaging experiments for monitoring the evolution of biomarkers.¹⁷ Moreover, autofluorescence¹⁸ originating from endogenous fluorescent molecules within cells and tissues poses another challenge in fluorescence analysis and imaging. The presence of intrinsic fluorophores, such as flavins¹⁹ and lipofuscin,²⁰ produces intense background signals that overlap with those emitted by the fluorescent probes. As such discriminating specific probe signals from intrinsic background autofluorescence is pivotal for achieving reliable imaging outcomes.

There is a growing emphasis on the development of advanced fluorescent probes with enhanced specificity,

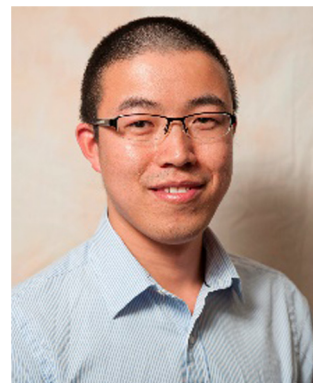


Luling Wu (left) and Tony D. James (right)



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minimized nonspecific interactions, and improved photostability.^{21,22} Innovative strategies explored for the design of new generation probes include the incorporation of responsive elements into probes and the design of alternative fluorophores with superior photophysical properties, such as responsive metal complexes,²³ aggregation-induced emission (AIE) probes,^{24–27} and near-infrared (NIR) II fluorescent dyes.^{28,29} In brief, while fluorescent probes have contributed significantly to the field of bioimaging, recognizing and mitigating the limitations of traditional probes are crucial for developing and validating new probes that can detect and image biomolecules in living cells and organisms.

Bioorthogonal chemistry^{30–32} refers to reactions occurring within biological settings without interruption of biomolecule synthesis or biochemical processes. Pioneered by the Carolyn Bertozzi group in 2003,³³ the term likely stems from the mathematical concept of orthogonality, indicating independent variation. Over the past two decades, bioorthogonal chemistry has evolved into a crucial tool for biological research. The key criteria of bioorthogonal reactions include operation under physiological conditions, selective and high-yield product formation, resilience to biological agents, rapid product formation at low concentrations, and stable product generation. Moreover, bioorthogonal chemistry involves functional groups that are not naturally present in biological systems, ensuring compatibility within complex environments. This methodology has significantly advanced biological studies by facilitating precise investigation of cellular processes while maintaining biological integrity. Notably, bioorthogonal chemistry, exemplified by copper-catalyzed azide-alkyne cycloaddition (CuAAC),^{34–37} Staudinger ligation,³ inverse-electron-demand Diels–Alder (IEDDA) reactions,^{38–40} and 1,2-aminothiol-2-cyanobenzothiazole (CBT) click reactions,^{41,42} has garnered enormous attention for selective biomolecule labeling,⁴³ targeted imaging,⁴⁴ and drug delivery⁴⁵ *in vivo* over recent years.

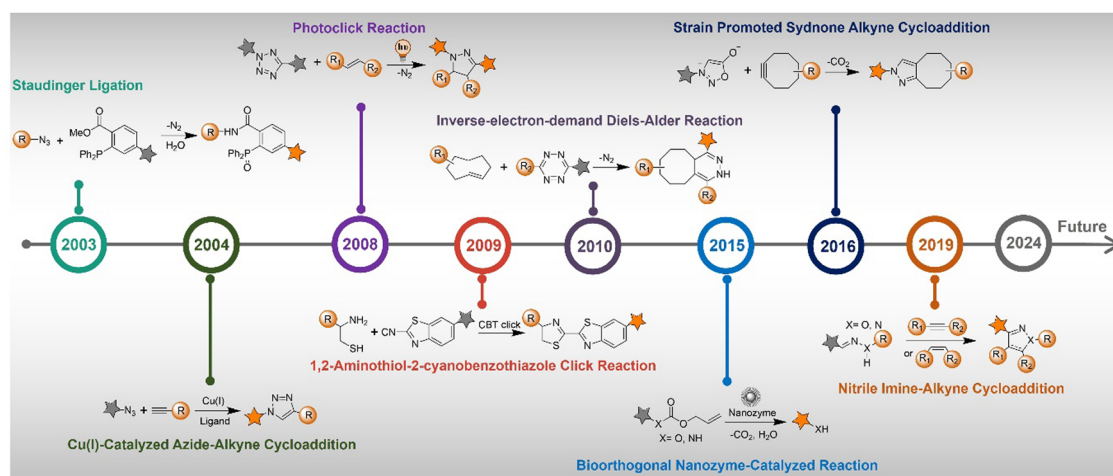
A signalling unit (chemical reporter) and a bioorthogonal click reaction handle are the two key components of bioorthogonally activated probes.⁴⁶ The majority of reported bioorthogonal probes are designed by directly incorporating a click

reaction handle into fluorescent dyes.^{31,43,47,48} Following the bioorthogonal reaction the stringent washing to remove any unreacted bioorthogonal probes is essential to reduce background fluorescence. Bioorthogonally activated probes have been subsequently developed to enhance the imaging precision and streamline sample pre-treatment procedures.^{23,44} These responsive probes exhibit weak fluorescence initially and can be activated by bioorthogonal reactions, resulting in an increase in the fluorescence signal. This kind of selective activation enables the probes to emit fluorescence only in regions where the bioorthogonal reaction occurs. As a result, there is no need for additional washing process in the molecular fluorescence imaging of cells, tissues, and living organisms that are stained with bioorthogonally activated probes. Such bioorthogonally activated probes are particularly useful in scenarios where extensive washing of the samples is not feasible.

In this review, we provide a comprehensive overview of the advancements in bioorthogonally activated fluorescent probe technology over the last decade, with particular focus on applications in precise, wash-free, and *in situ* biological imaging. We summarize the activated probes based on the types of bioorthogonal reactions, providing a succinct introduction to each reaction in each subsection. Subsequently, we outline recent research progress, identify the limitations of each bioorthogonal reaction, and discuss key future directions. This is followed by a comprehensive summary, emphasizing both the challenges and opportunities faced by bioorthogonally activated probes in achieving precise *in situ* biological imaging.

2. Bioorthogonally activated probes generated by different activation mechanisms

The advancement of bioorthogonally activated probes strongly relies on the underlying mechanisms of bioorthogonal activation. Since 2003, eight distinct categories of bioorthogonal activation have been identified and successfully adopted for



Scheme 1 A timeline of development of the bioorthogonally activated probes based on eight activation reactions.



the development of activated probes for precision bioimaging (Scheme 1). These mechanisms of bioorthogonal activation include Staudinger ligation, Cu(i)-catalyzed azide–alkyne cycloadditions (CuAAC), inverse-electron-demand Diels–Alder reactions (IEDDA), 1,2-aminothiol-2-cyanobenzothiazole (CBT) click reactions, strain promoted sydnone alkyne cycloadditions (SPSAC), nitrile imine–alkyne cycloadditions, photoclick reactions, and bioorthogonal nanozyme-catalyzed reactions. In this section, we will provide insights into advances in the development of bioorthogonally activated probes based on the eight bioorthogonal activation mechanisms as well as specific probes for precise fluorescence imaging.

2.1. Staudinger ligation

Staudinger ligation^{49–51} is an established bioorthogonal reaction derived from the classic Staudinger reduction.⁵² This bioorthogonal reaction involves the use of organic azides and triphenylphosphine-functional groups as the key components. The underlying mechanism of this bioorthogonal reaction is depicted in Fig. 1A. Diverging from the traditional Staudinger reduction pathway, the intermediate aza-ylide reacts with an adjacent ester, yielding a phosphorane that subsequently collapses to yield a stable amide linkage and a phosphine oxide. The second-order rate constant (k_2) for this reaction is approximately $\sim 10^{-3} \text{ M}^{-1} \text{ s}^{-1}$ in acetonitrile. During the reaction, the phosphine undergoes oxidation, forming a phosphine oxide, while the ester is cleaved, releasing methanol. These concurrent processes have been strategically exploited in developing advanced phosphine probes, activated *via* Staudinger ligation.

The Staudinger ligation is known for being bioinert and biocompatible since both reactants are entirely nonbiological,

which enables selective ligation across all biological compartments, distinguishing it from aldehyde or ketone condensation reactions. Popularized by Bertozzi's work,^{49,56,57} this reaction has evolved into a standard approach for the development of bioorthogonal probes and the modulation of biological processes. Herein, we examine the latest advancements in using Staudinger ligation-based bioorthogonal probes for precise *in situ* imaging of living organisms.

Chemical reactions that can occur within specific cellular environments contribute to the investigation of biological processes and the engineering of cell functions. A landmark achievement in this realm was reported in 2000 by the Bertozzi group,⁴⁹ introducing a chemical transformation that enables the precise formation of covalent adducts among intricately functionalized biopolymers within the cellular matrix. This transformative ligation, inspired by the Staudinger reaction, facilitates the formation of amide bonds through the coupling of an azide moiety with a custom-engineered triarylphosphine. Both reactive units, *i.e.*, azide and triarylphosphine, are synthetic and chemically orthogonal to endogenous cellular constituents. By harnessing metabolic pathways, azides can be incorporated into cell surface glycoconjugates *via* a synthetic azidosugar. The azide then selectively reacts with a biotinylated triarylphosphine, yielding stable cell-surface adducts. This research provided a new pathway for the simple engineering of cell surfaces using bioorthogonal reactions.

Apart from the glycoconjugate for cell surface modification, cellular glucose uptake plays a pivotal role in energy generation and is implicated in various metabolic disorders. To understand the details of the cellular glucose uptake process, Liang and coworkers⁵³ developed a bioorthogonal fluorescent probe,

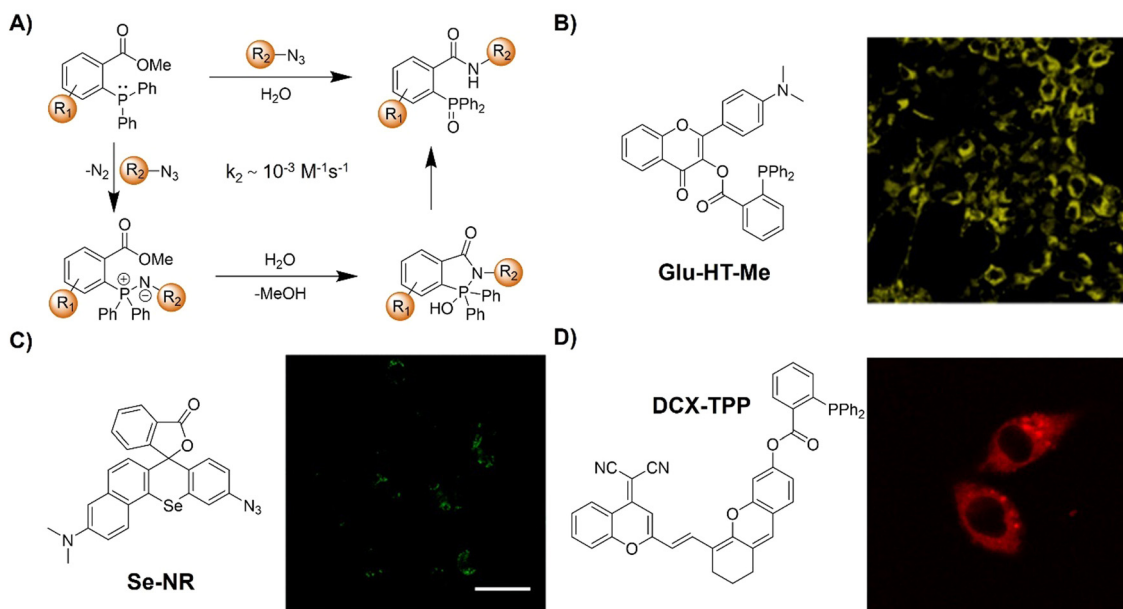


Fig. 1 Fluorescent probes based on Staudinger ligation. (A) Proposed mechanism of Staudinger ligation, (B) chemical structure and cell imaging of Glu-HT-Me, adapted with permission from ref. 53. Copyright 2022, American Chemical Society; (C) chemical structure and cell imaging of Se-NR, adapted with permission from ref. 54. Copyright 2019, Royal Society of Chemistry; (D) chemical structure and cell imaging of DCX-TPP, adapted with permission from ref. 55. Copyright 2019, Science Direct.



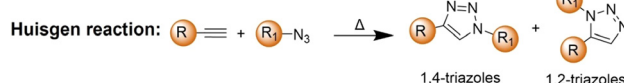
consisting of two parts Glu-HT-Me and azido-glucose, to light up the glucose uptake in live cells (Fig. 1B). The fluorescence of Glu-HT-Me was quenched due to photo-induced electron transfer (PeT) and the fluorescence was significantly increased (20-fold enhancement) when activated by internalized azido-glucose within 25 min. The fluorescence “OFF” to “ON” response allowed for monitoring glucose uptake dynamics in a wash-free manner. More interestingly, this bioorthogonal light-up fluorescent probe was able to discriminate cancer cells from normal cells due to the different glucose flux in anticancer drug treatment, glycolysis, and transport. The potential application of this probe in tracking glucose uptake fluctuations in a doxycycline-inducible K-ras^{G12V} oncogenic cell system was confirmed.

In addition to the bioorthogonal reaction in a cellular environment, bioorthogonal moieties have been used for the development of activatable photosensitizers and fluorescent probes for reactive biomolecule detection. In 2019, the Liu group⁵⁴ synthesized a π -extended Se-rhodamine (Se-NR, Fig. 1C) that is capable of efficiently producing singlet oxygen ($\Phi_{\Delta} = 0.35$) upon irradiation with red light. Leveraging precise control over the spirocyclization of Se-NR, a bioorthogonally activatable photosensitizer (Se-NR-Az) was developed by masking the amine group as an azide. This Se-NR-Az exhibited low cytotoxicity even under red light irradiation. However, the bioorthogonal reaction of Se-NR-Az with a phosphine resulted in the formation of a photocytotoxic compound, *i.e.*, Se-NR. This bioorthogonally activatable photosensitizer was then used for the controlled inhibition of the growth of HeLa cells by producing reactive oxygen species (ROS).

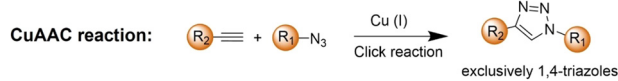
In another example, the triarylphosphine bioorthogonal moiety has been used by Zhang and coworkers for the development of an activatable probe (DCX-TPP) for the detection of nitroxyl (HNO), a type of ROS derived from nitric oxide (NO) through reduction and protonation (Fig. 1D).⁵⁵ The probe DCX-TPP was developed by combining a bioorthogonal HNO-recognition moiety, the diphenylphosphinobenzoyl group, with a NIR fluorophore *via* an ester linker. Initially non-fluorescent, HNO-catalyzed oxidation led to the activation of the probe and the emission of red fluorescence. The probe DCX-TPP could be used to detect 2–80 μ M HNO within 10 min with a detection limit (LoD) of 0.05 μ M and was able to visualize endogenous HNO in living cells.

In summary, the classic Staudinger ligation uses an azide, a small and biocompatible functional group, seamlessly integrated into biomolecules, for selective reactions with phosphine. However, its sluggish kinetics frequently lead to suboptimal labeling efficiency. While electron-rich phosphines can accelerate the bioorthogonal reaction, the rapid oxidation of phosphines under physiological conditions represents a significant challenge to the ligation process. As such, high concentrations of phosphine reagents are often required to ensure efficient bioorthogonal reactions. However, excess phosphine reagents can result in an increase in the background signal during fluorescence imaging, causing false positive results. While the Staudinger ligation remains one of the most

A)



B)



Scheme 2 Illustration of the (A) Huisgen reaction and (B) CuAAC reaction.

important activation mechanisms in the development of bioorthogonal probes, more recent efforts have focused on exploring faster ligation techniques in various scenarios.

2.2. Copper(I)-catalyzed azide–alkyne cycloaddition

The azide and terminal alkyne functional groups, being small and primarily engaged in weak dipolar interactions, exert minimal influence on the functional properties of the molecules to which they are attached. Both azides and alkynes exhibit low reactivity in biological systems, enabling them to traverse intact cells before being triggered by specific conditions or catalysts. The Huisgen 1,3-dipolar cycloaddition⁵⁸ of organic azides and alkynes, yielding 1,2,3-triazoles, is typically too slow for biological applications (Scheme 2A). However, the introduction of Cu(I) catalysts (Scheme 2B), pioneered by Meldal and Sharpless, significantly accelerates triazole formation up to 100 million-fold.^{34–36,59} The Cu(I)-catalyzed azide–alkyne cycloaddition (CuAAC) facilitates the formation of 1,4-disubstituted triazoles exclusively, mimicking native peptide-based connectors in size and geometry and offering superior stability to thermal and chemical cleavage.

CuAAC is modular, compatible with aqueous conditions, and largely by-product-free, facilitating successful click reactions. By using various catalysts, including simple Cu(I) salts, small-molecule Cu-binding ligands, and diverse materials, the application of CuAAC has been confirmed in organic synthesis,⁶⁰ polymer functionalization,⁶¹ drug discovery,⁶² and chemical biology.⁶³ While naturally occurring azides are rare, labelling of proteins, lipids, oligonucleotides, and cell surface moieties with azides or alkynes can be achieved through benign chemical transformations or metabolic incorporation of labeled metabolites, enabling their conjugation to orthogonally labeled partners.

The adoption of CuAAC for bioorthogonal bioconjugation has been significantly advanced by the development of water-soluble Cu(I)-binding ligands, mitigating the limitation of Cu(I)-mediated ROS formation through Fenton reaction with endogenous H₂O₂. These ligands stabilize the Cu(I) oxidation state, accelerating the CuAAC reaction while minimizing ROS production through the Fenton reaction. Ligands with additional features, such as ROS scavenging, have expanded the use of CuAAC for more demanding biological applications, including *in situ* labeling of living cellular surfaces, tagging enzymes or biomolecules in complex cellular mixtures, rapid click reactions within cellular compartments, and versatile dye labeling

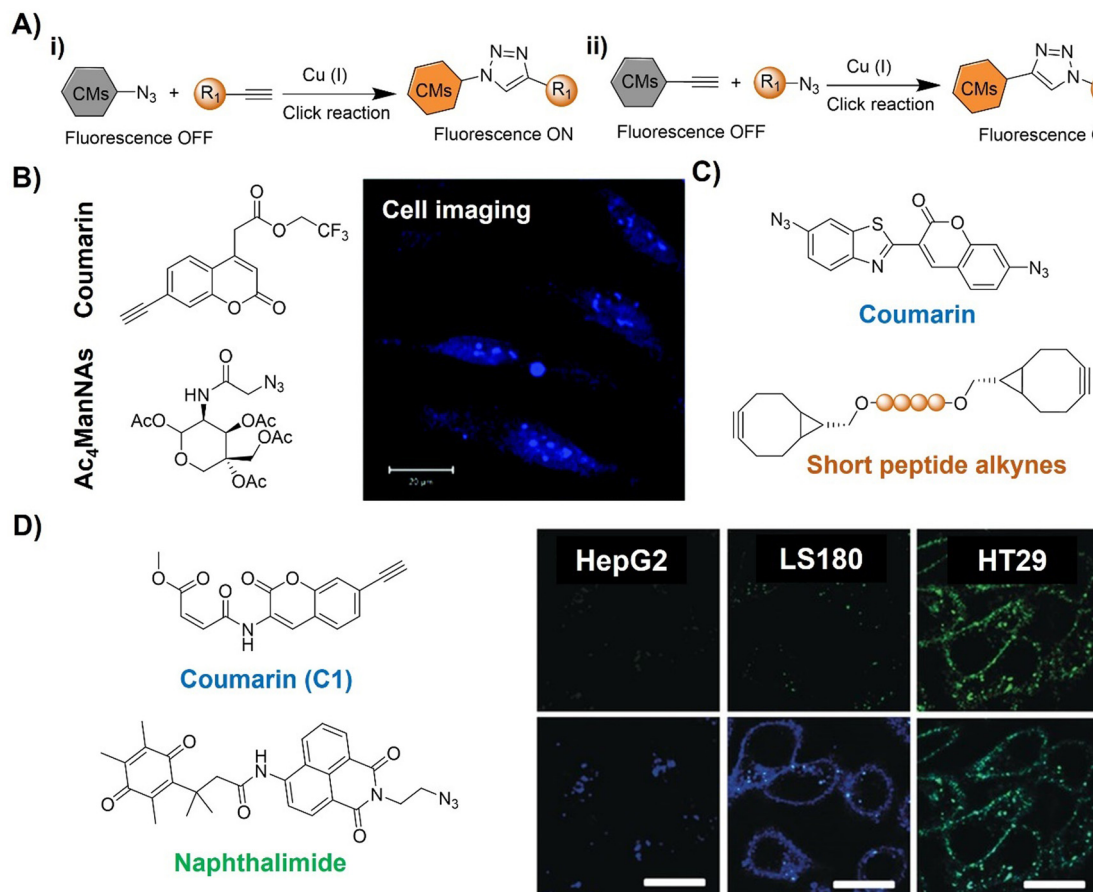


Fig. 2 Coumarin fluorophore-based bioorthogonal probes using copper(I)-catalyzed azide–alkyne cycloaddition. (A) Illustration of the CuAAC activated coumarin probes; (B) the structure and cell imaging of the alkyne modified coumarin, adapted with permission from ref. 64. Copyright 2014, The Royal Society of Chemistry; (C) the structures of bis-azide coumarin and biscyclooctynylated peptides; (D) the structures of the profluorescent coumarin derivative (C1) and azido-naphthalimide and bioimaging in different cell lines, adapted with permission from ref. 65. Copyright 2017, Wiley-VCH GmbH.

of diverse cellular structures for super-resolution microscopy. Herein, we provide a comprehensive overview of CuAAC reaction applications in the realm of *in situ* precision imaging over the last decade, categorized by the types of probes developed.

2.2.1. Coumarins. Zhang and colleagues⁶⁴ have engineered an alkyne-functionalized coumarin fluorogenic glycoproteomic probe to visualize sialoglycoconjugates within living cells (Fig. 2B). Using coumarin's well-established CuAAC fluorogenic properties, an alkyne group was introduced at the 7 position of coumarin, transforming it into a profluorophore. A trifluoroethyl ester-protected terminal carboxyl group at the 4 position is used to enhance water solubility and offers the opportunity for additional modification prior to bioorthogonal labeling. This design enabled a CuAAC reaction with a mannose analogue containing an azido functionality in aqueous solution and HeLa cells. Given the fluorescence “off-on” response after the click reaction, additional washing was not required prior to cell imaging.

A bis-azide fluorogenic probe for two-point bioorthogonal tagging was developed by Kele *et al.*⁶⁶ for labeling biscyclooctynylated peptides (Fig. 2C). This probe exhibited robust fluorescence enhancement with a polar hexapeptide containing two

cyclooctyne motifs. Theoretical computation revealed that the fluorescence “off-on” response is dependent on dihedral angles between the probe's aromatic core. Amongst the screened biscyclooctynylated peptides, a polar hexapeptide demonstrated near-maximal fluorogenicity. These bis-azides offered a new approach to develop bioorthogonal probes for specific, fluorogenic two-point labeling of proteins containing two genetically encoded cyclooctyne motifs.

Similarly, Liu and colleagues⁶⁵ developed a doubly caged heterodifunctional linker, a profluorescent coumarin derivative (C1) with orthogonal dual “click” motifs, for functionalization of protein/antibody bioconjugates (Fig. 2D). The core molecule, C1 contains both CuAAC and Michael addition sites on a coumarin skeleton for a “click” style reaction with azide- and thiol-functionalized precursors, respectively. Click reactions of the ethynyl of C1 with naphthalimide QNAM-N₃ catalyzed by Cu(I) catalysis and the C1's alkenyl amide/ester unit with a thiol functionalized anti-carcinoembryonic antigen (ACEA) produced QNAM-C1-ACEA and switched on coumarin's emission at 435 nm. The quinone-caged naphthalimide exhibited weak fluorescence, while the emission could be switched on ($\lambda_{\text{em}} = 530$ nm) after a quinone oxidoreductase (NQO1)-triggered



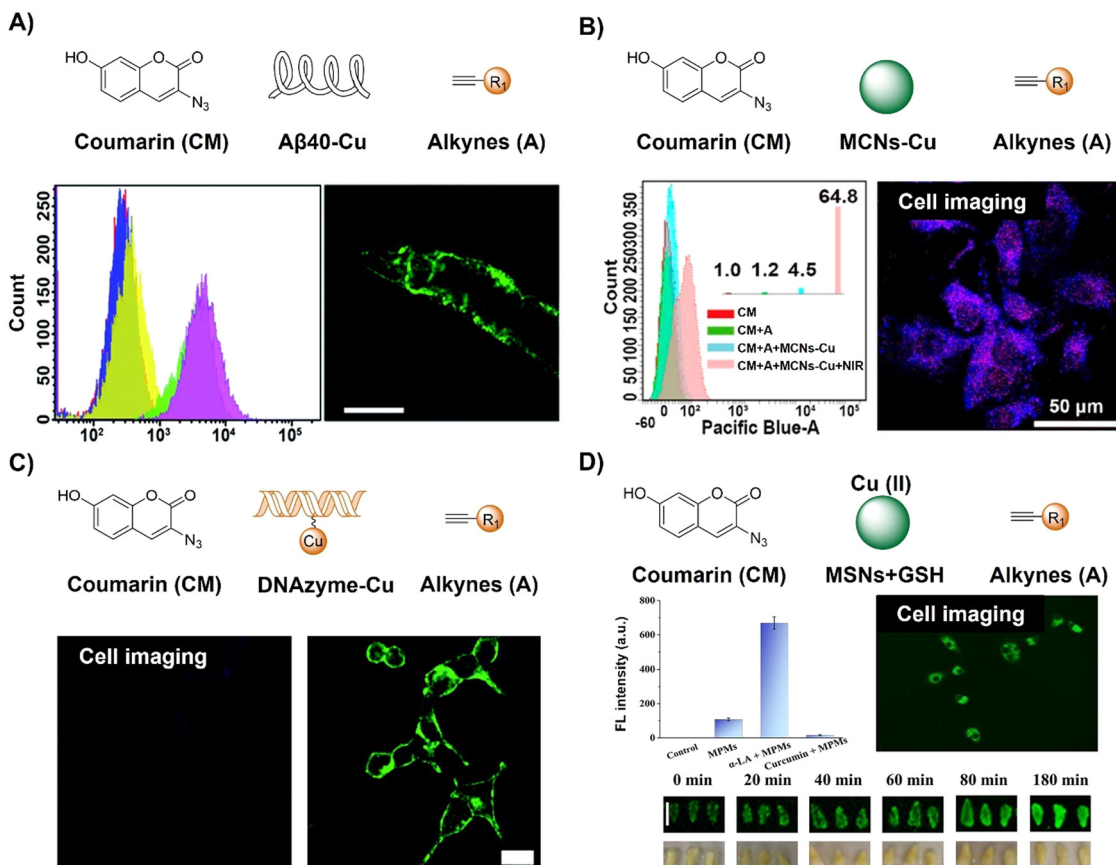


Fig. 3 Coumarin fluorescent probes based on the copper(II)-catalyzed azide–alkyne cycloaddition. (A) The structure, flow cytometry detection and worm *in situ* imaging of the coumarin-A_β-Cu system, adapted from ref. 67, under the license CC BY-NC 3.0 DEED; (B) the structure, flow cytometry detection and *in situ* cell imaging of the coumarin–MCNs–Cu system, adapted with permission from ref. 68. Copyright 2020, American Chemical Society; (C) the structure and *in situ* cell imaging of the coumarin–DNAzyme–Cu system, adapted from ref. 69, under the license CC BY-NC 3.0 DEED; and (D) the structure and *in situ* cell imaging and plumule imaging of the coumarin–MCNs/GSH–Cu(II) system, adapted with permission from ref. 70. Copyright 2021, Elsevier B.V.

cleavage of the quinone cage. Moreover, overlapping of coumarin's emission with the absorption of naphthalimide's allowed FRET upon excitation at 365 nm, facilitating the ratiometric fluorescence response of QNAM-C1-ACEA for analysis and imaging of NQO1. This double-caging strategy and modular/fluorogenic fabrication of functional protein/antibody conjugates offers versatility, potentially integrating with other bioorthogonal reactivity methods.

Cu is an essential element in the body and is linked to neurodegenerative disorders, notably Alzheimer's disease, marked by A_β aggregation and Cu accumulation. In addition to the exogenous Cu catalyst, the Qu group⁶⁷ found that Cu accumulation in A_β plaques can effectively catalyze an azide–alkyne bioorthogonal cycloaddition for fluorescence activation of coumarin and drug synthesis in living cells, a transgenic AD model of *Caenorhabditis elegans* CL2006, and brain slices of triple transgenic AD mice (Fig. 3A). The as-synthesized drug destroyed A_β-Cu aggregates, alleviating paralysis and locomotion defects. This *in situ* approach is a self-triggered drug synthesis method for AD therapy, a pioneering approach utilizing *in situ* Cu for click reactions in physiological environments.

Taking advantage of enhanced stability and biosafety, heterogeneous Cu nanoparticles (CuNPs) have emerged as promising contenders for facilitating the CuAAC reaction. Research conducted by Qu *et al.*⁶⁸ resulted in a NIR light promoted CuAAC reaction using a biocompatible heterogeneous copper nanocatalyst, which was developed for investigating both photodynamic and photothermal effects *in vitro* and *in vivo* (Fig. 3B). Specifically, upon NIR light irradiation, Cu nanoparticle attached mesoporous carbon nanospheres (MCNs-Cu) produce reactive oxygen species (ROS) and heating in a local microenvironment through photodynamic and photothermal ($\eta = 50.6\%$) effects, respectively, which promote the conversion of inert Cu(0) to Cu(I), expediting the catalytic CuAAC process. This synergistic effect resulted in enhanced activation of a profluorophore (3-azido-7-hydroxycoumarin), as observed in cellular and murine models under NIR irradiation. In addition to the use of CuNPs as a catalyst for CuAAC reactions, further research from the same group also used DNAzyme-augmented bioorthogonal catalysis for targeted synergistic cancer therapy.⁶⁹ The design of a DNAzyme leveraged DNA structure and substrate interaction enabled precise imaging

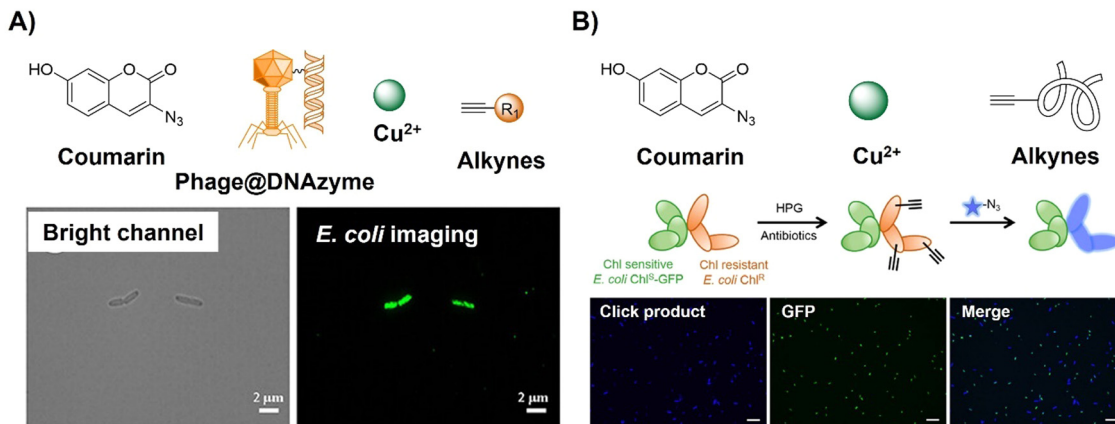


Fig. 4 Coumarin fluorophore-based bioorthogonal probes obtained using the copper(i)-catalyzed azide–alkyne cycloaddition. (A) The structure and *E. coli* imaging of the Phage@DNAzyme probe, adapted with permission from ref. 71. Copyright 2023, American Chemical Society; (B) the structure and selective *E. coli* labeling of the azido-coumarin system, adapted with permission from ref. 72. Copyright 2022, American Chemical Society.

in situ and efficient prodrug activation in cancer cells for synergistic cancer therapy (Fig. 3C).

CuAAC reaction activated 3-azido-7-hydroxycoumarin has also been explored for the development of fluorescent probes for the detection of glutathione (GSH), antimicrobial resistant bacteria, and foodborne pathogens. In 2021, Song and colleagues⁷⁰ devised a GSH-triggered MSN (mesoporous silica nanoparticle) nanoreactor featuring a Cu(ii)-BTC shell, where Cu(i), generated through *in situ* reduction by GSH, serves as a catalyst (Fig. 3D). This catalytic activity expedites the bioorthogonal reaction between an alkyne and azide, thereby enhancing the fluorescence of 7-hydroxycoumarin and 4-ethynylanisole loaded within the MSNs. Evaluation of these nanoreactors for GSH detection, spanning cellular to wheat plumule levels, demonstrated their robust biocompatibility and potential for detecting metal ions in plants. As such, the monitoring of intracellular bioorthogonal reactions could advance precision chemistry in the design of intelligent platforms for biomarker detection as well as targeted drug synthesis for precision medicine.

Antibiotic resistance, stemming from the ongoing overuse of antibiotics, poses a global threat to human and animal health, environmental stability, food and nutrition security, economic development, and societal equity and safety. Detection of pathogens, particularly antibiotic-resistant bacteria, has contributed to

addressing this threat. CuAAC activated probes based on coumarin derivatives have been applied for foodborne pathogen detection. For example, Gan *et al.*⁷¹ reported a simple, sensitive on-site bioassay for foodborne pathogens employing the Phage@DNAzyme probe (3-azido-7-hydroxycoumarin, Fig. 4A). The probe displayed specificity towards bacterial strains (*E. coli* as the model) and catalytic efficiency for fluorescent CuAAC click reactions. Synergistic signal amplification using a gold nanoparticle modified slide and Phage@DNAzyme catalysis enabled rapid, sensitive, on-site bacterial measurement with a limit of detection (LOD) of 50 CFU mL⁻¹ in 30 minutes, which could be detected using a smartphone. Using a CuAAC reaction, Yu *et al.*⁷² developed a rapid bioorthogonal noncanonical amino acid (BONCAT) labelling method for the detection of antimicrobial-resistant bacterial cells in an aquatic environment (Fig. 4B). Using this system the fluorescence of 3-azido-7-hydroxycoumarin was switched on after a CuAAC reaction with the L-homopropargylglycine incubated *E. coli*. In the presence of an antibiotic (e.g., chloramphenicol), the antibiotic-resistant *E. coli* could be easily identified within 20 min of coincubation with 3-azido-7-hydroxycoumarin.

2.2.2. Naphthalimides. Naphthalimide and its derivatives with alkyne or azide functionalization have also been explored for the development of bioorthogonally activated fluorescent probes (Fig. 5A). For example, taking advantage of the specific

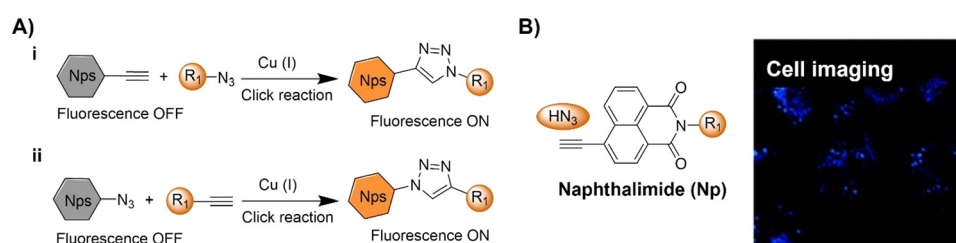


Fig. 5 Bioorthogonal fluorescent probes based on the copper(i)-catalyzed azide–alkyne cycloaddition with a naphthalimide fluorophore. (A) Illustration and (B) chemical structure of the CuAAC activated naphthalimide probe for cell imaging. Adapted with permission from ref. 73. Copyright 2013, The Royal Society of Chemistry.



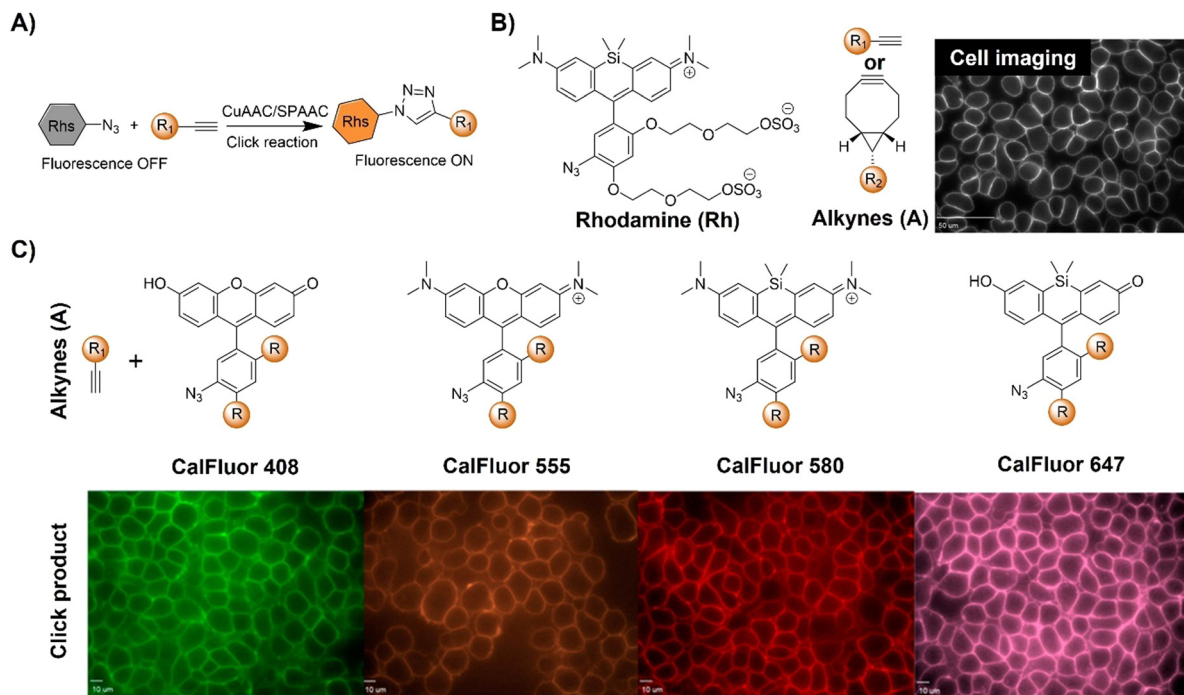


Fig. 6 Bioorthogonal fluorescent probes based on the copper(I)-catalyzed azide–alkyne cycloaddition with rhodamine derivatives. (A) Illustration of the CuAAC activated rhodamine probe; (B) the structure and the wash-free mammalian cell surface labeling of the Si-rhodamine, adapted with permission from ref. 74. Copyright 2014, National Academy of Sciences; (C) structures of CalFluors 488, 555, 580, and 647 and their ability to enable the no-wash labeling of cell-surface glycoproteins on HEK 293T cells, adapted with permission from ref. 75. Copyright 2015, American Chemical Society.

reaction between alkynes and azides, Yao *et al.*⁷³ introduced an alkyne-based naphthalimide fluorescent probe that can be activated through bioorthogonal reaction for the detection of hydrazoic acid (HN_3). The emission of this probe was quenched due to internal charge transfer (ICT) and switched on after a clicking with HN_3 , exhibiting high selectivity over other ROS and reactive nitrogen species (RNS). The probe was used for the fluorescence detection and imaging of HN_3 in living HeLa cells and larval zebrafish (Fig. 5B).

2.2.3. Rhodamine derivatives. Rhodamine derivatives have been widely used for the development of fluorescent probes for bioassays and imaging, such as the development of bioorthogonal probes for the imaging of bacterial peptidoglycan by Bertozzi and co-workers (Fig. 6A and B).⁷⁴ The probe was prepared by functionalization of NIR-emitting Si-rhodamine with an azide. For this probe the azide served as (1) a quencher for Si-rhodamine's emission through PeT and (2) the click reaction moiety for bioorthogonal chemistry. Significantly an up to 48-fold fluorescence increase was obtained upon triazole formation through click reaction of the probe with terminal or strained alkynes. Further modification by glycol substitution generated probes (Fig. 6B) with high water solubility for “wash-free” labelling in live cells and bacteria. Moreover a copper free click reaction was observed between azido-PEG3-carboxyrhodamine 110 and cyclooctyne D-alanine analogs, enabling peptidoglycan (PG) imaging with high sensitivity while the cytotoxic Cu catalyst was not necessary for the click reaction.

In a biological environment, bioorthogonal fluorescent probes spanning the visible spectrum offer precise visualization of metabolically labeled molecules. Toward this end, Bertozzi *et al.*⁷⁵ reported the development of highly fluorogenic azide probes based on the PeT fluorescence response mechanism (Fig. 6B). Activated by click chemistry to form triazoles, these probes exhibited remarkable fluorescence enhancement, up to 283-fold, with a Φ_F of 0.74. Exploiting a similar PeT mechanism, bioorthogonal probes were developed using diverse fluorophores including bodipy, cyanines, and pyrazolines. Termed CalFluors, these probes with emission from green to far-red wavelengths enabled the sensitive detection of biomolecules without washing steps. Through bioorthogonal reactions, visualization of alkyne-labeled biomolecules (glycans, DNA, RNA, and proteins) in various cellular contexts, including zebrafish and mouse brain tissue slices, was achieved. Further modification of rhodamine derivatives enabled Zhang *et al.*⁷⁶ to develop a platform to evaluate protein engineering outcomes *via* quantification of extracellular alkyne-tagged metabolites using a fluorogenic click reaction. This method facilitated the screening and optimization of alkyne-based biosynthetic tools. For example, the probes for JamB, a membrane-bound bifunctional desaturase/acetylenase, exhibited approximately 20-fold enhanced activity in *E. coli*.

2.2.4. Bodipy. Bodipy dyes⁷⁷ have unique emission properties, such as small Stokes shifts, narrow absorption bands, sharp emissions, high fluorescence quantum yields, and excellent chemical and photostability. Using biocompatible bodipy

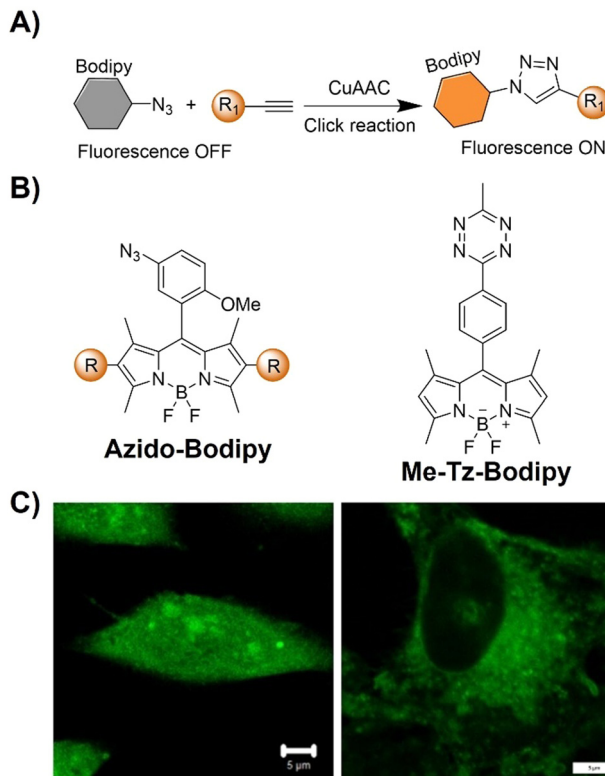


Fig. 7 Bioorthogonal fluorescent probes based on the copper(I)-catalyzed azide–alkyne cycloaddition with Bodipy derivatives. (A) Illustration of the CuAAC activated Bodipy probe for cell imaging; chemical structure (B) and cell imaging (C) of azido-Bodipy and Me-Tz-Bodipy. Adapted with permission from ref. 78. Copyright 2023, Wiley-VCH GmbH.

as the fluorophore, Shie *et al.*⁷⁸ developed a bioorthogonally activated probe for monitoring the intracellular location of metabolic glycan and live cell glycosylation pattern investigation *via* multicolor labeling (Fig. 7). A bodipy probe bearing an azide moiety could label multiple glycoconjugates containing alkynyl groups *via* CuAAC reactions. The fluorescence labelling exhibited a high signal-to-noise ratio, allowing for the effective tracking of sialylation and GlcNAcylation. The probe could be

used to label cell-surface and intracellular sialylated glycoconjugates and exhibited site-specific dual-channel fluorescence imaging of HeLa and PC9 cancer cells. Moreover, the fluorescence labeling of sialylated glycoproteins and glycans by the bodipy bioorthogonal probe was validated using MS-based proteomic analysis, confirming the probes as robust tools for glycan imaging and glycoproteomic analysis of cellular systems.

2.2.5. Aggregation-induced emission (AIE) probes.

Aggregation-induced emission (AIE)^{24,26,79} is an emerging concept, pioneered by Tang's group, where non-emissive fluorophores like tetraphenylethene emit intense fluorescence in aggregated and solid states. Bioorthogonally activated probes are typically designed using blue or green emissive fluorophores with azide or tetrazine (**Tz**) functional groups. To enhance the availability of the bioorthogonal probes, in 2016, Liu *et al.*⁸⁰ introduced red-emitting AIE fluorogens (AIEgens) for the development of bioorthogonally activated probes for cancer cell imaging and ablation (Fig. 8). Initially non-fluorescent in water due to free molecular motion, emission of the probes switched on upon reacting with azide-functionalized glycans on the cancer cell surfaces (Fig. 8B and C). Additionally, the AIEgens served as photosensitizers to produce cytotoxic ROS under visible light ($\lambda = 400\text{--}700\text{ nm}$) irradiation, enabling photodynamic ablation of cancer cells.

Incorporating bioorthogonal chemical groups into synthetic glycans followed by click reactions with fluorescent probes is pivotal in monitoring the metabolic glycoengineering bioprocesses of unnatural glycans in living systems. Following the success of their prior work, the Liu group⁸¹ devised a dual-targeted metabolic precursor, cRGD-S-Ac₃ManNAz, selectively targeting $\alpha_5\beta_3$ integrin-overexpressing MDA-MB-231 cancer cells for internalization through receptor mediated endocytosis and subsequent generation of azide-tagged glycans (Ac₃ManNAz) upon the GSH-mediated cleavage of the disulfide bond on cRGD-S-Ac₃ManNAz. A bioorthogonal fluorescent probe, TPE-BAI, was also developed for specific cancer cell labelling (Fig. 9A). TPEBAI was initially non-fluorescent in aqueous media but exhibited light-up fluorescence upon azide click reaction, enabling selective cancer cell imaging. TPEBAI's

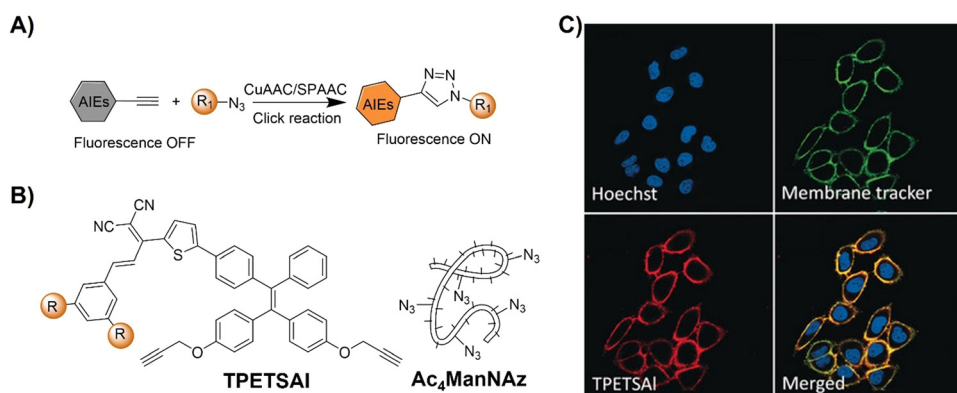


Fig. 8 Bioorthogonal probes for fluorescence imaging based on the copper(I)-catalyzed azide–alkyne cycloaddition with AIEgens. (A) Illustration of the CuAAC activated AIEgen probe for cell imaging; chemical structure (B) and (C) co-stain of TPETSAI labeled HeLa cells with membrane tracker CellMask Green. Adapted with permission from ref. 80. Copyright 2016, Wiley-VCH GmbH.

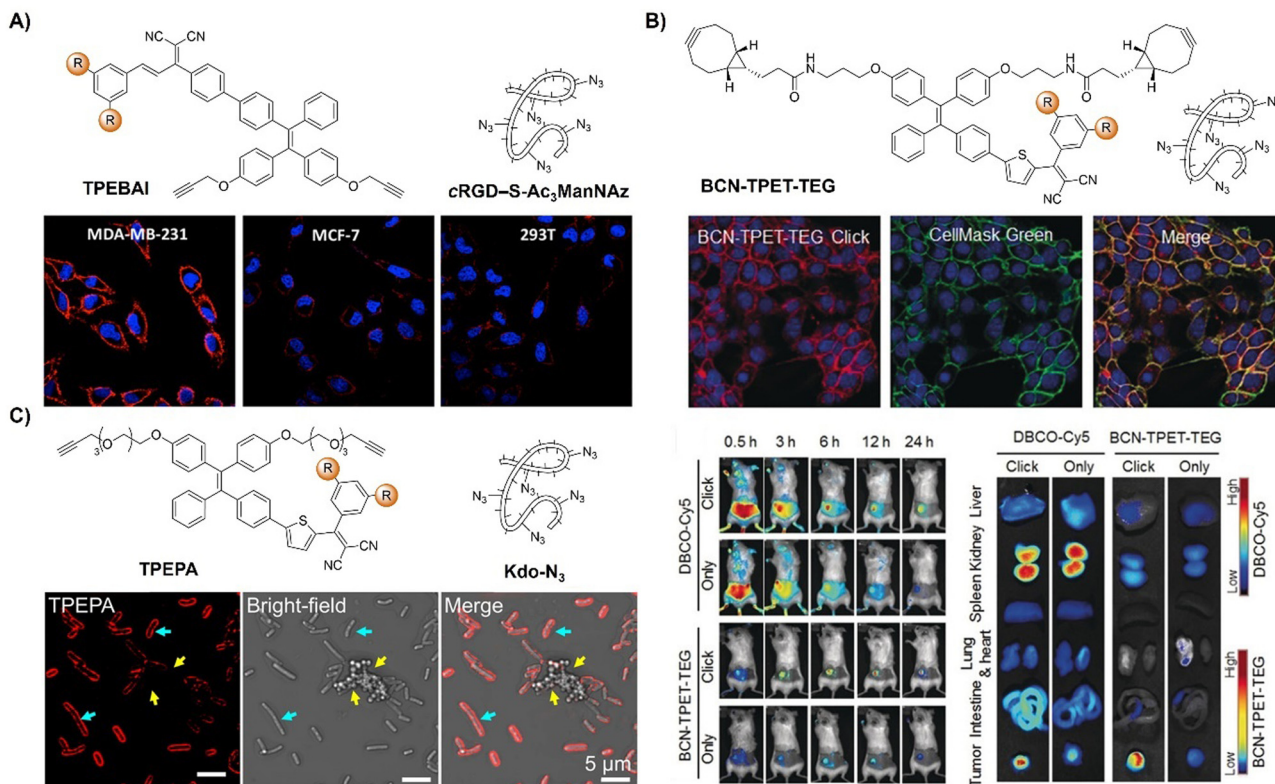


Fig. 9 AIE probes for bioorthogonal reaction based on the copper(I)-catalyzed azide–alkyne cycloaddition. (A) The structure and confocal images of TPEBAI labeled MDA-MB-231/MCF-7/293T cells. Adapted with permission from ref. 81. Copyright 2018, American Chemical Society; (B) the structure, overlay images of 4T1 cells and *in vivo* imaging of BCN-TPET-TEG. Adapted with permission from ref. 82. Copyright 2018, Wiley-VCH GmbH; (C) the structure and TPEPA-selective metabolic labeling of Gram-negative (*E. coli*) and Gram-positive bacteria (*S. aureus*) with Kdo-N₃ addition. Adapted with permission from ref. 83. Copyright 2020, American Chemical Society.

robust $^1\text{O}_2$ production also allowed for image-guided cancer cell ablation under light irradiation. Normal cells that feature low $\alpha_1\beta_3$ integrin expression and intracellular GSH levels showed minimal TPEBAI labeling and negligible toxicity under light irradiation, offering precise cancer cell imaging and ablation capabilities.

As a result of the covalent ligation, bioorthogonal labeling using activated probes generally exhibits a better performance in imaging and drug delivery in tumors than active targeting strategies. For instance, Liu *et al.*⁸² developed a light-up fluorescent probe (BCN-TPET-TEG, BCN = bicyclo[6.1.0]non-4-yne) utilizing a AIEgen for *in vivo* bioorthogonal fluorescence tumor labeling (Fig. 9B). BCN-TPET-TEG exhibited low background fluorescence in aqueous media and minimal non-specific interaction with normal tissues. With a broad visible absorption spectrum (400–600 nm) and bright NIR emission ($\lambda_{\text{em}} = 700$ nm), emission of BCN-TPET-TEG was rapidly switched on after reacting with azide groups present on tumor cells, facilitating swift tumor-specific imaging. Moreover, the photosensitizing capacity enables effective image-guided photodynamic tumor therapy.

In addition to the tumor specific labelling and ablation through PDT, AIE based bioorthogonal probes have been developed for specific bacteria discrimination and biofilm

eradication by Liu's group.⁸³ TPEPA represents an example of a bioorthogonal fluorescence turn-on probe for this purpose (Fig. 9C). This probe exhibited excellent water solubility and minimal fluorescence in water-based environments. Upon reaction with azide groups engineered onto bacteria through metabolic processes, emission of TPEPA at 680 nm was switched on, enabling fluorescence imaging and discrimination of bacteria, *i.e.*, Gram-negative (G^-) *E. coli* bacteria with Kdo-N₃ and Gram-positive (G^+) *S. aureus* bacteria with D-Ala-N₃. ROS generation under light irradiation enabled the bioorthogonally activated probe TPEPA to be used as a photosensitizer for the precise ablation of bacteria. Leveraging cell surface components of pathogenic bacteria for specific G^+ and G^- bacterial discrimination, Liu *et al.*⁸⁴ then devised a fluorescence-based Gram-staining method using bioorthogonal fluorescence turn-on probes TPPB and BPTZ (Fig. 10A). Metabolic incorporation of Kdo-N₃ and Ala-CP into bacterial surface membranes allowed for selective labeling *via* strain-promoted alkyne–azide cycloaddition (SPAAC) reaction and Diels–Alder reaction with inverse electronic demand (DAR_{inv}). TPPB targets characteristic LPS in G^- bacteria, while BPTZ labels both G^+ and G^- bacteria, enabling clear differentiation with high selectivity and sensitivity (Fig. 10B).

2.2.6. Others. In addition to the above-discussed systems, other fluorescent molecules like cresyl violet, benzothiazolium



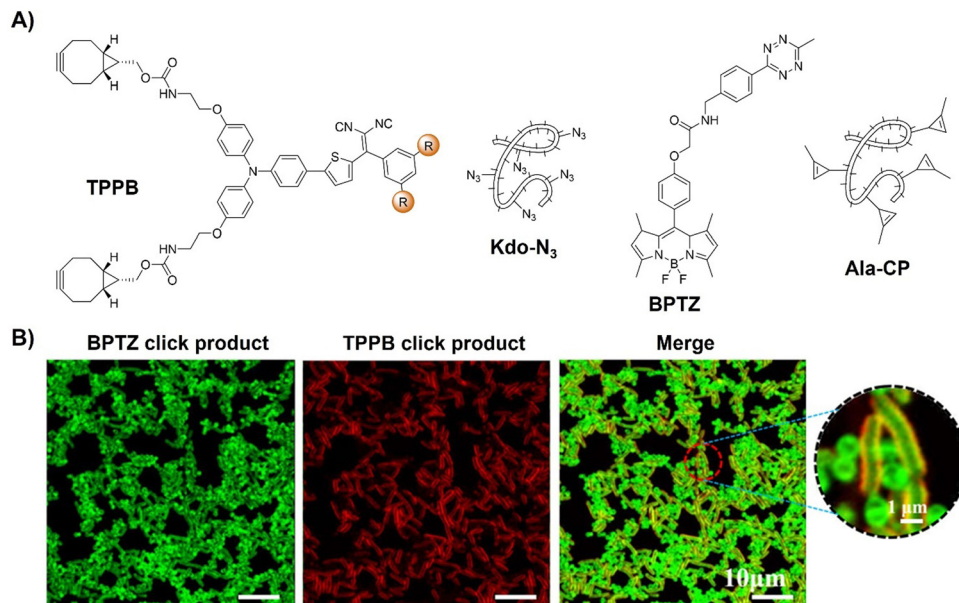


Fig. 10 Bioorthogonal probes for bacterial imaging based on the copper(I)-catalyzed azide–alkyne cycloaddition with an AIEgen. (A) The structures of TPPB, Kdo-N₃, BPTZ and Ala-CP, (B) metabolic dual labeling of G[−] bacteria (*E. coli*) and G⁺ bacteria (*S. aureus*) after being co-cultured with solution and then treated with TPPB, Kdo-N₃, BPTZ and Ala-CP. Adapted with permission from ref. 84. Copyright 2021, American Chemical Society.

and 8-aminoquinoline have been used to develop CuAAC-activated probes for precise *in situ* imaging. For example, using cresyl violet as the azido fluorophore, Duan *et al.*⁸⁵ developed an azide–alkyne cycloaddition based probe for the ratiometric fluorescence labeling and imaging of live cells (Fig. 11A). Initially the azido cresyl violet (Acv) emits at 566 nm but upon click reaction an emission at 620 nm emerged due to the reduced electronic density in the triazole ring, enabling ratiometric fluorescence monitoring. The probe was used for the ratiometric fluorescence imaging of propargylcholine- and dibenzocyclooctyne-labeled MCF-7 cells.

In 2015, Boons and colleagues⁸⁶ introduced Fl-DIBO, a dibenzocyclooctyne derivative fused with a cyclopropenone moiety, capable of undergoing rapid strain-promoted cycloaddition reactions without catalysts (Fig. 11B). Fl-DIBO reacted rapidly with azides, nitrones, nitrile oxides, and mono- and disubstituted diazo derivatives. While reactions with certain compounds resulted in products of low quantum yield, mono-substituted diazo reagents produced 1*H*-pyrazole derivatives exhibiting about 160-fold fluorescence enhancement over Fl-DIBO, with over a 10 000-fold increase in brightness. Fl-DIBO was then used for the precise background-free fluorescence labeling of diazo-tagged proteins, demonstrating the potential of diazo derivatives as biomolecule tags.

Red-emitting fluorescent benzothiazolium dimethine dyes with azide groups have also been employed for bioorthogonal conjugation to alkyne targets by Kele *et al.*⁸⁷ Upon the CuAAC reaction of weakly fluorescent probes with alkyne-modified oligonucleotides, the corresponding triazolic products exhibit visible excitation and NIR emission with large Stokes shifts and high quantum yields (a Φ_F of up to 0.7). For the alkyne functionalized targets, incorporating DNA with the

benzocyclooctyne motif through rigid (ethynyl) or flexible (ethyl) linkers enabled copper-free click reaction labelling of DNA strands. *In vitro* evaluation using HeLa cells transfected with cyclooctynylated-DNA confirmed cellular uptake without toxicity, enabling specific tagging of target DNA with negligible background fluorescence, facilitating fluorescent cell imaging (Fig. 11C). Similarly, another two bioorthogonal probes with quinolinium units and azide or Tz motifs on DNA-sensitive frames (TOA and TOT respectively) were developed by the Németh group⁸⁹ to detect DNA–protein interactions (Fig. 11E).

Bioorthogonal activation and precise imaging *in vivo* are important for the investigation of bioprocesses but remain challenging due to the limited tissue penetration of excitation light. In this context, Zhu *et al.*⁸⁸ developed a series of azide-based light-up bioorthogonal probes, leveraging a weakly fluorescent 8-aminoquinoline (AQ) scaffold for *in vivo* two-photon fluorescence imaging (Fig. 11D). These probes, upon click reaction with alkynes, exhibited remarkable fluorescence enhancement (up to 1352-fold). By introducing various styryl groups into the AQ scaffold, the probes exhibited excellent two-photon properties ($\delta = 542$ GM at 780 nm). Extensive bioimaging experiments validated the versatility of these probes for live cell/zebrafish imaging, eliminating the need for washing steps. Notably, *in vivo* two-photon imaging experiments showcased the superior performance of these bioorthogonal probes over conventional fluorophores, boasting a high signal-to-noise ratio and deep tissue penetration.

In summary, the CuAAC reaction has been widely employed for developing bioorthogonally activated probes using fluorophores like coumarin, naphthalimide, rhodamine, bodipy, and AIEgens, facilitating precise *in situ* imaging in bacteria and live cells. However, its translation into *in vivo* applications faces



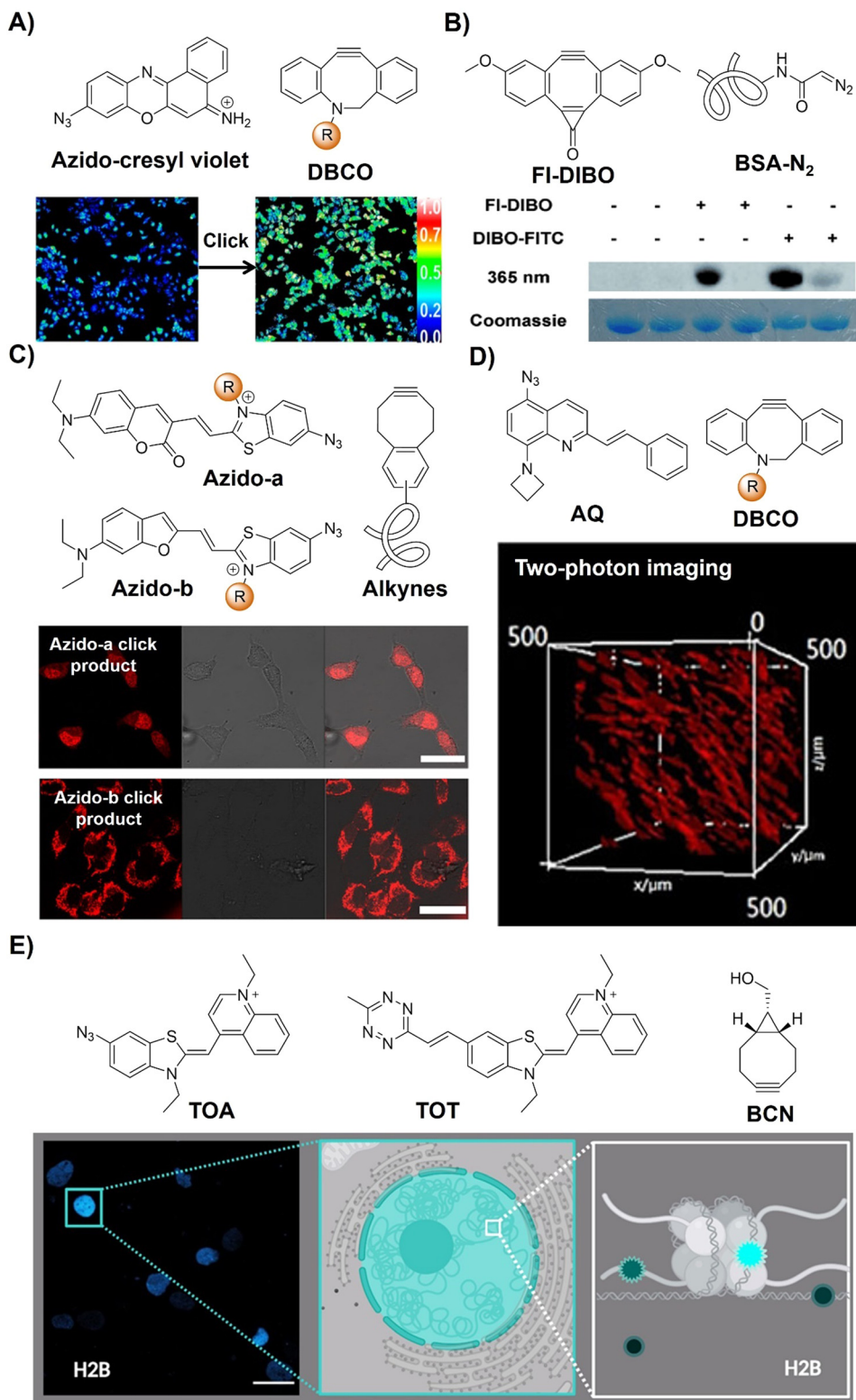


Fig. 11 Bioorthogonal probes based on the copper(I)-catalyzed azide–alkyne cycloaddition with other fluorophores. (A) The structure of azido-cresyl violet and ratiometric fluorescence imaging before and after click reaction. Adapted with permission from ref. 85. Copyright 2015, American Chemical Society; (B) the structure of FI-DIBO/BSA-N₂ and target protein labeling. Adapted with permission from ref. 86. Copyright 2015, Wiley-VCH GmbH; (C) the structure and cell imaging of azido-a/b. Adapted with permission from ref. 87. Copyright 2016, American Chemical Society; (D) the structure and two-photon cell imaging of AQ. Adapted with permission from ref. 88. Copyright 2020, Wiley-VCH GmbH; and (E) the structures and nucleus imaging of TOA and TOT, adapted from ref. 89, licensed under CC-BY 4.0.

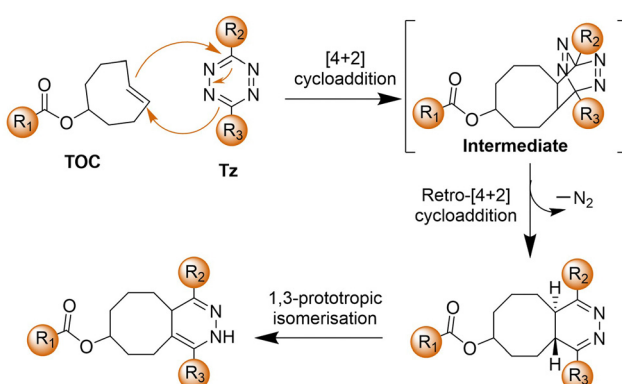
challenges. Direct utilization of Cu(i) in living systems can induce the generation of ROS, leading to cellular toxicity, which restricts the application of CuAAC bioorthogonal activation for *in vivo* use. Efforts to enhance biocompatibility focus on developing water-soluble ligands to accelerate CuAAC while scavenging ROS to mitigate cytotoxicity. Although Cu(i) remains the preferred catalyst, exploration of alternative transition metals such as Ru,⁹⁰ and Rh⁹¹ has been pursued, predominantly in organic solvents, impeding their biological applicability.

Efforts to bypass transition-metal catalysts entirely have gained momentum. Notably, the development of Cu-free SPAAC reactions^{30,31,48,92–95} forming triazole products has emerged. Incorporating alkynes within strained cyclooctyne systems achieves Cu-free modification, enhancing biocompatibility, albeit with reaction rates comparable to Staudinger ligation and slower than CuAAC. Improving reaction kinetics involves integrating electron-withdrawing fluorine atoms into cyclooctyne reagents, exemplified by difluorinated cyclooctyne (**DIFO**),⁹⁶ significantly enhancing reactivity for *in vivo* bioorthogonal imaging of azide-labeled sugars. Initial *in vivo* bioimaging applications in organisms such as *Caenorhabditis elegans* and zebrafish embryos were studied using **DIFO** and azides.⁹⁷ Further development of cyclooctyne analogues with improved water solubility and kinetics aims to enhance azide–alkyne cycloaddition efficacy in living organisms.

Despite addressing slow kinetics in Staudinger ligation and the potential toxicity of CuAAC, SPAAC's performance remains suboptimal due to reduced reaction rates, susceptibility to side reactions, and reliance on sterically larger connecting groups that compromise functionality. Consequently, research persists in identifying fully biocompatible accelerating ligands for CuAAC, in order to facilitate rapid intracellular reactions without inducing biological stress or interfering with endogenous copper-binding biomolecules.

2.3. Inverse-electron-demand Diels–Alder reaction

The bioorthogonal Tz ligation^{38,39,98,99} describes a chemical reaction involving a 1,2,4,5-Tz (Tz) and an alkene or alkyne dienophile, yielding a dihydropyridazine or pyridazine conjugate *via* a [4+2]/retro [4+2] cycloaddition sequence (Scheme 3).



Scheme 3 The proposed mechanism of the IEDDA reaction.

This reaction, known as the IEDDA reaction,¹⁰⁰ stands out for its rapid kinetics, occurring without the need for catalysis and yielding nitrogen gas as the sole byproduct. Initially utilized in organic synthesis for their swift reactivity with strained dienophiles, Tzs found application in bioorthogonal reactions in 2008.³⁸ This breakthrough involved reactions with derivatives of *trans*-cyclooctene (TCO) and norbornene by the groups of Fox³⁸ and Weissleder,¹⁰¹ respectively. Subsequently, there has been a proliferation of chemical diversity for both Tzs and their dienophile reaction partners, allowing for fine-tuning of reaction rates, stability, and the design of 'minimal' reporter molecules. The IEDDA reaction of Tz ligation is accelerated by electron-withdrawing groups on the Tz.^{102,103} Tzs are commonly synthesized through the condensation of Pinner salts or nitriles with hydrazine, often facilitated by Lewis acids, thiols, or sulfur catalysts.¹⁰⁴ Alternatively, milder protocols have been developed for directly introducing intact Tz groups *via* palladium-catalyzed coupling reactions of aryl halides or aryl boronic acids.¹⁰⁴ Carboxylic esters also serve as handles for the preparation of unsymmetrical and monosubstituted Tzs.

Various dienophiles have been developed for Tz ligation. TCO, prepared *via* flow photochemistry, exhibits unmatched reactivity with Tzs, engaging in the most rapid bioorthogonal reactions known under aqueous conditions.¹⁰⁵ Highly strained cyclooctynes such as BCN also react rapidly, yielding aromatic conjugates useful for workflows requiring defined stereochemistry. Cyclopropenes offer a complementary approach, especially beneficial for genetic encoding and metabolic incorporation. Norbornene, known for its good kinetics at a low cost, has proven advantageous for scale-up and materials applications. Smaller dienophiles such as cyclobutenes, azetidines, unstrained olefins, and isonitriles also react efficiently with Tzs, albeit often requiring large excesses of one reagent due to slower kinetics. Triazines can serve as surrogates for Tzs, offering advantages of smaller size and greater stability.^{104,106}

Additionally, Tz also acts as a fluorescence quencher, with its mechanism varying based on the design of the bioorthogonal probes. The common mechanism involves non-radiative dissipation of molecular energy to quench the emission of the fluorophores, where the excited state energy of the fluorophore is released through vibrations, heat, or interactions with the surrounding environment. However, after the IEDDA reaction, these energy release pathways are removed, restoring fluorescence. As a result of the "off-on" fluorescence response, Tz-conjugated bioorthogonal fluorescent probes could be used for wash-free molecular imaging. Varieties of Tz-modified fluorophores use diverse quenching mechanisms, including electronic energy transfer (EET) such as Förster resonance energy transfer (FRET), Dexter, and through-bond energy transfer (TBET), monochromophoric design, and new fluorophore formation post-IEDDA. FRET exploits donor–acceptor excited state interactions, requiring spectral overlap. Dexter-type EET involves electron hopping between molecules, needing donor–acceptor wavefunction overlap. TBET occurs *via* a rigid π-system linker. Monochromophoric design relies on dark-state Tz quenching. Post-reaction, optically forbidden transitions shift

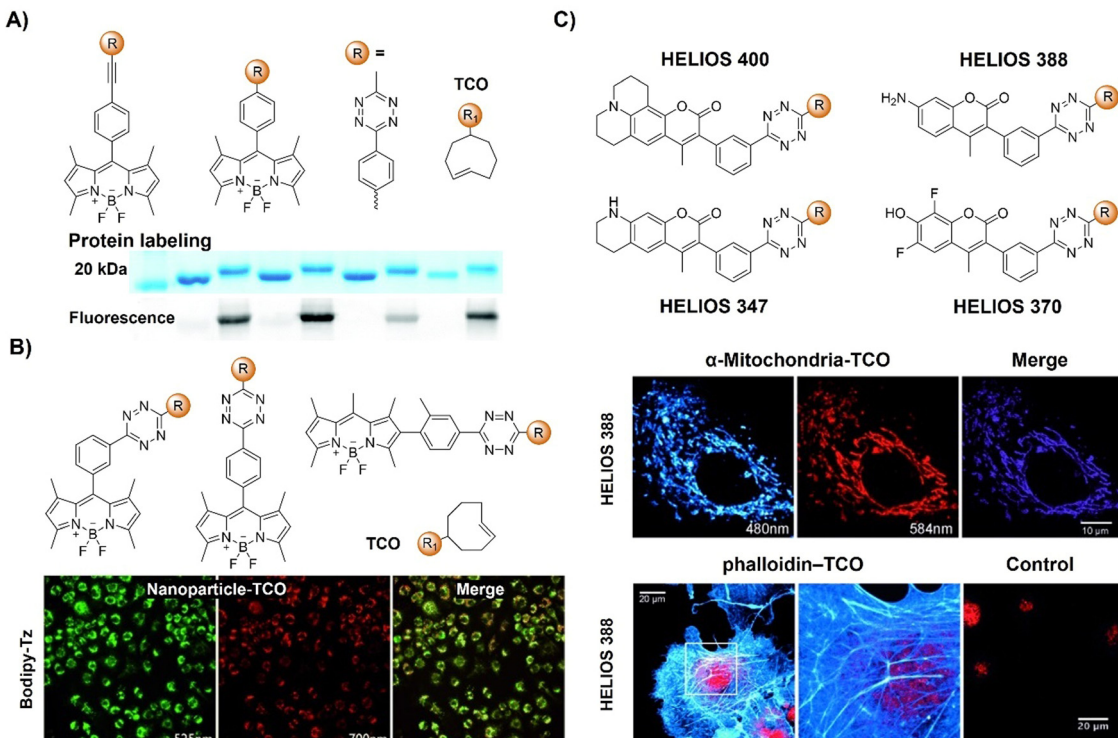


Fig. 12 Fluorescent probes based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure and protein labeling of the Tz modified Bodipy, adapted with permission from ref. 107. Copyright 2014, The Royal Society of Chemistry; (B) the structures and cell imaging of the Bodipy-p-Tz derivatives, adapted with permission from ref. 108. Copyright 2013, Wiley-VCH GmbH; (C) the structures of the HELIOS probes and the wash-free fluorogenic imaging of intracellular targets, adapted with permission from ref. 109. Copyright 2014, Wiley-VCH GmbH.

to allowed states, allowing for changes in the fluorescence signal enabling molecular imaging. Research has highlighted that new fluorophore formation post-IEDDA offers high turn-on ratios. Optimizing energy transfer efficiency from a fluorophore to a quencher is crucial in EET strategies. These varied fluorescence response approaches enable flexibility for the development of bioorthogonally activated probes for precise *in situ* imaging.

2.3.1. TBET approach. Using Sonogashira and Stille cross-coupling, alongside CuAAC, 1,2,4,5-Tz fluorophore derivatives with structurally rigid molecular frameworks were synthesized by Wombacher *et al.*¹⁰⁷ as bioorthogonally activated probes for protein labelling (Fig. 12A). These bichromophoric systems exhibited low fluorescence quantum yields, primarily attributed to Tz-mediated quenching *via* TBTE. The extent of fluorescence quenching was contingent upon the separation distance between the fluorophore and the Tz. However, this reduced fluorescence was effectively restored through Tz conversion *via* IEDDA, leading to a fluorescence enhancement of up to 24-fold. Time-resolved spectroscopy unveiled resonance energy transfer between the Bodipy fluorophore and the Tz as the underlying quenching mechanism. Remarkably, these conjugates proved effective in protein labeling experiments (Fig. 12A).

Another Bodipy-Tz platform was reported by Weissleder and colleagues for the development of bioorthogonal turn on probes (Fig. 12B).¹⁰⁸ After reaction with TCO, the π conjugation

of Tz-Bodipy with *para*- and *meta* Tz exhibited a 900 and 1600-fold increase in fluorescence, respectively. The significant enhancement in fluorescence facilitated the use of the probe for bioorthogonal wash-free labelling and imaging of fixed and live cells. Following this success, the same group developed a series of fluorescent probes with emissions ranging from blue to green, showing fluorescence enhancement surpassing 11 000-fold with a Φ_F of up to 0.49 (Fig. 12C).¹⁰⁹ These “off-on” response probes were engineered using the coumarin fluorochrome with the bioorthogonal TCO-Tz chemistry platform. Leveraging highly efficient TBET, these probes exhibit significant enhancement in brightness (2500–11 000-fold enhancement depending on chemical structures and emission of probes). The probes enabled wash-free, fluorogenic imaging of various targets such as cell-surface receptors in cancer cells, mitochondria, and the actin cytoskeleton in a few seconds, featuring minimal background signal and negligible non-specific binding.

To address the limitation of synthetic methods for preparation of Tz-based bioorthogonal probes, Devaraj *et al.*¹¹⁰ introduced an elimination-Heck cascade reaction for the *in situ* synthesis of (*E*)-3-substituted 6-alkenyl-1,2,4,5-Tz derivatives, including 24 compounds with Tz conjugation (Fig. 13A). These derivatives, including fluorescein and Bodipy-based fluorescent probes, are suitable for live-cell imaging, exhibiting strong fluorescence turn-on response (up to 400-fold) upon reacting

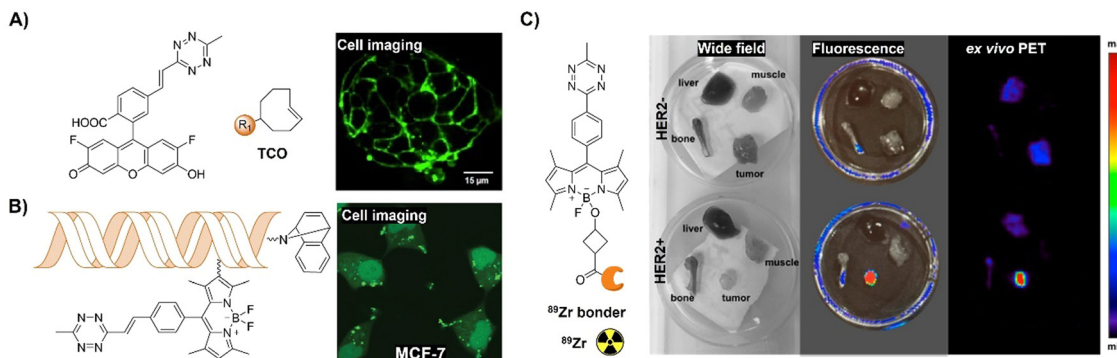


Fig. 13 Fluorescent probes based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure and LS174T cell imaging of the alkenyl-Tz modified fluorescein-based fluorescent probe, adapted with permission from ref. 110. Copyright 2014, Wiley–VCH GmbH; (B) structures of the Tzs and dienophiles used and live-cell miRNA detection in the human cancer MCF-7 cell line. Adapted with permission from ref. 111. Copyright 2014, American Chemical Society; (C) the structure of the bioorthogonal fluorogenic PET probe and the wide field, fluorescence and *ex vivo* PET images acquired for the muscle, liver, bone, and tumor, adapted with permission from ref. 112. Copyright 2016, American Chemical Society.

with dienophiles like cyclopropenes and **TCO**. In another research conducted by the same group,¹¹¹ a fluorogenic Tz-mediated transfer (TMT) reaction employing the 7-azabenzonorbornadiene derivative was reported for the detection of oligonucleotides with high sensitivity and sequence specificity (Fig. 13B). The key requirement for achieving signal enhancement is the template-driven turnover of antisense probes that was facilitated by spontaneous diaxine release post-Tz ligation. Using a highly quenched alkenyl-fluorogenic Tz achieved an over 100-fold fluorescence increase *via* the TMT reaction. RNA-templated transfer reactions with antisense probes enabled mir-21 detection with high sensitivity at low-picomolar levels, suitable for distinguishing single-base mismatches.

Multi-modality imaging, such as fluorescence-PET, using multiple functionalized probes is an emerging approach for better understanding the biological processes in living organisms. Linking deferoxamine (DFO) with Bodipy-Tz, Weissleder *et al.*¹¹² reported a bioorthogonal labeling method to create PET/fluorescence mAb imaging probes. The chelate linker, utilizing **TCO-Tz** chemistry, integrated DFO for ⁸⁹Zr PET imaging and a bodipy for fluorescence imaging. Fluorescence activation (<3 min) during biomolecule–fluorophore–chelator conjugation allowed for real-time monitoring and precise quantification *via* absorbance measurements (Fig. 13C). In mice with tumor xenografts, the probe exhibited high target specificity in PET and robust *ex vivo* fluorescence imaging.

A series of water soluble fluorogenic xanthene Tz derivatives (Fig. 14A) spanning a broad emission spectrum (521–666 nm), encompassing green (fluorescein Oregon Green), yellow (tetramethyl-rhodamine), and far-red (Si-rhodamine), were synthesized by Wombacher *et al.*¹¹³ as bioorthogonal probes for wash-free fluorescence imaging of proteins in live cells. Using these fluorescent probes for selective protein labelling achieved up to 109-fold enhancement in fluorescence upon bioorthogonal IEDDA reaction (DA_{inv}), allowing for wash-free fluorescence microscopic imaging of the actin cytoskeleton in

fixed cells and visualization of mitochondrial and nuclear proteins in living cells.

In addition, the bifunctional rhodamine fluorophore has also been employed for the development of bioorthogonally activated probes. For instance, Wombacher's group reported a bifunctional rhodamine-based fluorescent probe for a DNA-templated assay (Fig. 14B).¹¹⁴ The bifunctional probe (Rhod-Tz-NHS) was prepared and the proximity effect on the reactivity of DAinv-bioorthogonal chemistry was investigated using different dienophiles with different reactivities. Stable dienophiles with poor kinetics gained reactivity through DNA-templated proximity, generating fluorescence signals exclusively from proximity-induced reactions. Up to 34-fold enhancement of the fluorescence signal was obtained within 40 min, with negligible background reactivity at micromolar concentrations. This proximity concept addressed stability challenges in bioorthogonal chemistry, maintaining crucial rapid reaction rates for cellular applications. A subsequent study conducted by the same research group uncovered a dark intermediate in the fluorogenic reaction between Rhod-Tz and **TCO**, which will be discussed below.¹¹⁵

The Tz unit can quench emission from luminophores *via* FRET, TBET, or PeT, with quenching efficiency influenced by factors like the distance between the donor and acceptor, dipole orientation, spectral overlap, and electron transfer thermodynamics. Unlike the above probes with fluorescence emission from the singlet excited state (S₁), metal complexes like iridium(III) and ruthenium(II) complexes emit from the triplet excited state (T₁) characterized by large Stokes shift, prolonged emission lifetime, and excellent photostability.^{116,117} Therefore, a number of responsive probes have been successfully developed for luminescence bioassays and bioimaging. For example, Lo's group¹¹⁸ developed a bioorthogonal luminescence probe based on a luminescent iridium(III) complex bearing a single built-in Tz for protein labelling and cell imaging (Fig. 15A). The probe was prepared by coordinating iridium(III) with a Tz-containing pyridyl ligand, resulting in a weakly luminescent

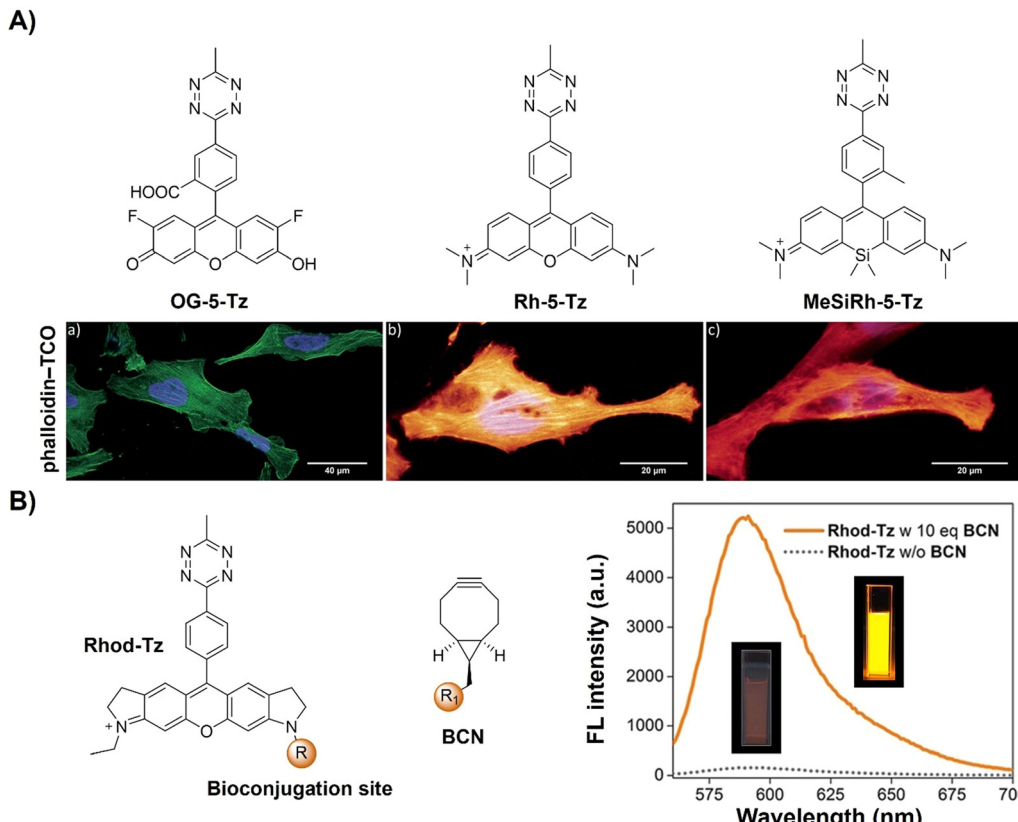


Fig. 14 Fluorescent probes based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structures of green- to far-red-emitting fluorogenic xanthene Tzs and wash-free fixed cell confocal imaging of actin cytoskeleton, adapted with permission from ref. 113. Copyright 2017, The Royal Society of Chemistry; (B) the structures of Rhod-Tz and BCN and emission spectra of Rhod-Tz with/without BCN addition, adapted with permission from ref. 114. Copyright 2017, Wiley-VCH GmbH.

iridium(III) complex with a 60-fold higher reactivity than the ligand itself. These complexes were stable upon co-incubation with biothiols but exhibited significant emission enhancement upon treatment with BCN-OH. HaloTag technology facilitated protein conjugation, with BCN-C6-Cl enabling phosphorogenic coupling. For example, GST-HaloTag treated with BCN-C6-Cl prior to incubation with complex 3 exhibited increased emission intensity, as evidenced by a luminescence band only present in the BCN-C6-Cl-loaded SDS-PAGE gel. The application of complex 3 was demonstrated for detection of BCN-modified endogenous HaloTag proteins in HeLa cells.

Reactions with nonfluorescent raw materials to produce highly fluorescent products through bioorthogonal activation are interesting, particularly for fluorescence labelling and *in situ* imaging of living organisms. Vrabel's research¹¹⁹ revealed how the conformation of the non-fluorescent TCO dienophile directly influences the fluorescent product formation in IEDDA reactions with 1,2,4,5-Tzs (Fig. 15B). Among the evaluated dienophiles, rapid reaction with the axial TCO isomer led to the formation of fluorescent 1,4-dihydropyridazine products with up to 91-fold enhancement. By adjusting the Tz's substitution pattern, the photochemical properties of the corresponding products could be easily tuned to form a new type of fluorophore with intriguing emission

properties. This fluorogenic reaction was then applied for rapid labeling of the intracellular compartments of live cells, indicating its potential for advancing IEDDA applications in various fields, including live-cell imaging.

Fluorescence labelling and imaging of cellular organelles, like the mitochondria, are highly desirable for understanding bioprocesses at a subcellular level. To achieve this goal, Wang *et al.*¹²⁰ reported a molecular guided approach for delivering functional molecules to mitochondria *via* intracellular bioorthogonal reactions (Fig. 15C). Oligo(*p*-phenyleneethynylene) (OPE) that was modified with Tz groups for bioorthogonal click reaction exhibited weak emission. Under laser irradiation and upon bioorthogonal reaction with Mito-TCO bearing triphenylphosphine (TPP) for mitochondrial targeting, a fluorescent Intra-Mito product formed in the cytoplasm could accumulate in the mitochondria, enabling precise mitochondrial labelling and imaging. Unlike most commonly used fluorescent dyes, such as rhodamine 110 and fluorescein for cell labelling, this bioorthogonal approach and the fluorescent product exhibited high stability for the long-term fluorescence imaging of live cells. Another example of TPP guided mitochondrial imaging post-bioorthogonal reaction was reported by Vrabel and colleagues (Fig. 15D).¹²¹ In this system, Tzs with electron-donating substituents were linked to the strained bicyclononyne dienophile *via* a

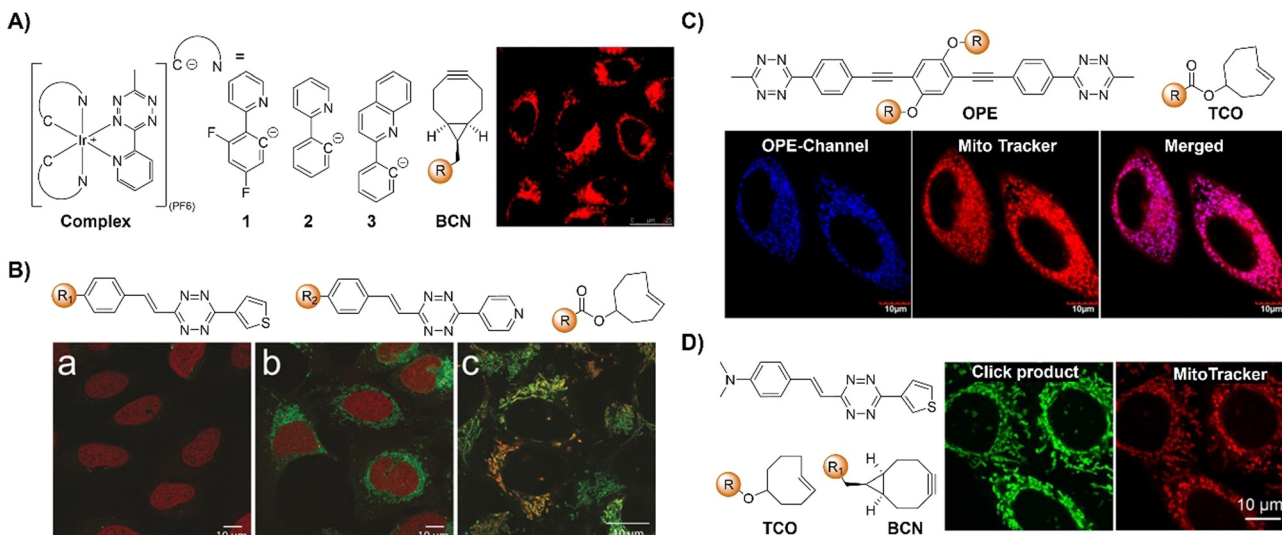


Fig. 15 Bioorthogonal probes based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structures of the Ir complex and BCN and LSCM images of nontransfected HeLa cells, adapted with permission from ref. 118. Copyright 2017, The Royal Society of Chemistry; (B) the structures of a microtubule-selective Taxol-Tet probe and a mitochondria-selective TPP-Tet probe and confocal microscope images of U2OS cells, adapted with permission from ref. 119. Copyright 2017, Wiley-VCH GmbH; (C) the structure of OPE and colocalization analysis of OPE-Mito (top) and OPE (bottom) with Mito Tracker in MCF-7 cells, adapted with permission from ref. 120. Copyright 2018, American Chemical Society; and (D) the structures of electron-donating substituents linked via a π -system fluorophore-Tz and strained dienophiles and fluorogenic labeling of live HeLa cells, adapted with permission from ref. 121. Copyright 2019, Wiley-VCH GmbH.

π -system. The produced fluorescent pyridazine dyes resulted in 100–900-fold enhancement of fluorescence intensity upon bioorthogonal reaction. The resulting fluorophores exhibited large Stokes shift and relatively high fluorescence quantum yield for fluorescence labelling and imaging. Reaction of the bioorthogonal probe with TPP linked bicyclononyne enabled the fluorescent product to accumulate in the mitochondria for fluorescence imaging.

The majority of fluorophore-Tz conjugate (FLTz) strategy-based bioorthogonal probes have been developed using a donor–acceptor-type energy transfer mechanism. However, the effective switching on of fluorescence after click reaction and shifting of a probes' emission wavelength are challenging. To address this, Park and coworkers¹²² introduced a monochromophoric design strategy to expand the emission color range of **FLTzs** while maintaining a high fluorescent turn-on/off ratio (Fig. 16A). Indolizine-based emission-tunable Seoul-Fluor (**SF**) was used as a model fluorophore system for these probes. By enhancing electronic coupling between **Tz** and π -conjugated systems of the indolizine core, the emission of **SF-Tz** conjugates (**SFTzs**) was efficiently quenched. Subsequently an over 1000-fold fluorescence enhancement with a Φ_F of up to 0.68 after the IEDDA reaction with **TCO** was observed. Colorful **SFTzs** with consistent turn-on/off ratios were then developed across various emission wavelengths. Demonstrating the applicability of the bioorthogonal probes, fluorescence bioimaging of microtubules and mitochondria in live cells was achieved in a wash-free manner, offering a reliable strategy for developing versatile bioorthogonal fluorogenic probes with excellent turn-on ratios, regardless of the emission wavelength.

Activation of AIEgen-based bioorthogonal probes allows for the ROS (*e.g.*, singlet oxygen $^1\text{O}_2$) generation for photodynamic

therapy (PDT).^{80–83} Previous research conducted by Weissleder and colleagues has demonstrated the formation of highly luminescent probes after bioorthogonal Bodipy/Tz activation.¹⁰⁸ Taking advantage of halogenated Bodipy's photosensitizing performance, Vázquez and colleagues¹²³ developed a series of Bodipy bioorthogonal turn-on probes for precisely generating $^1\text{O}_2$ by modifying the 4,4-difluoro-4-bora-3a,4a-diaza-s-indacene core (Fig. 16B). These halogenated Bodipy-Tz probes exhibited non-toxicity to HeLa cells and “off-on” fluorescence response in cell imaging as well as effective photosensitization ($k \approx 3 \times 10^{-3} \text{ s}^{-1}$, $\Phi_\Delta \approx 0.50$) for inhibiting cancer cell growth and were obtained through intracellular IEDDA with suitable dienophiles. Another example of site specific bioorthogonal photosensitization for PDT was reported by Lo and colleagues.¹²⁴ In this system, the bodipy-based dye, distyryl boron dipyrromethene (DSBDP), was equipped with two functional tags, including 1,2,4,5-Tz and alkyne groups (Fig. 17A). These tags could selectively bind to **TCO** and azide groups, respectively, which were introduced onto the membrane of A431 cells through pre-targeting with **TCO-GE11** or metabolic glycoengineering with **Ac4ManNAz**. Upon bioorthogonal conjugation, the photosensitizer exhibited increased cellular uptake, ROS generation, and photocytotoxicity. The bioorthogonal reaction was confirmed in tumor-bearing nude mice, illustrating the potential of tumor treatment through bioorthogonal activation and light irradiation.

In addition to bioorthogonal activation for photosensitization, bioorthogonal prodrug activation for tumor imaging and treatment was achieved by Wu and colleagues.¹²⁵ In this system, a vinyl ether-masked camptothecin (**CPT**) and a **Tz** tethered to a NIR fluorophore (**NR**) were prepared and then



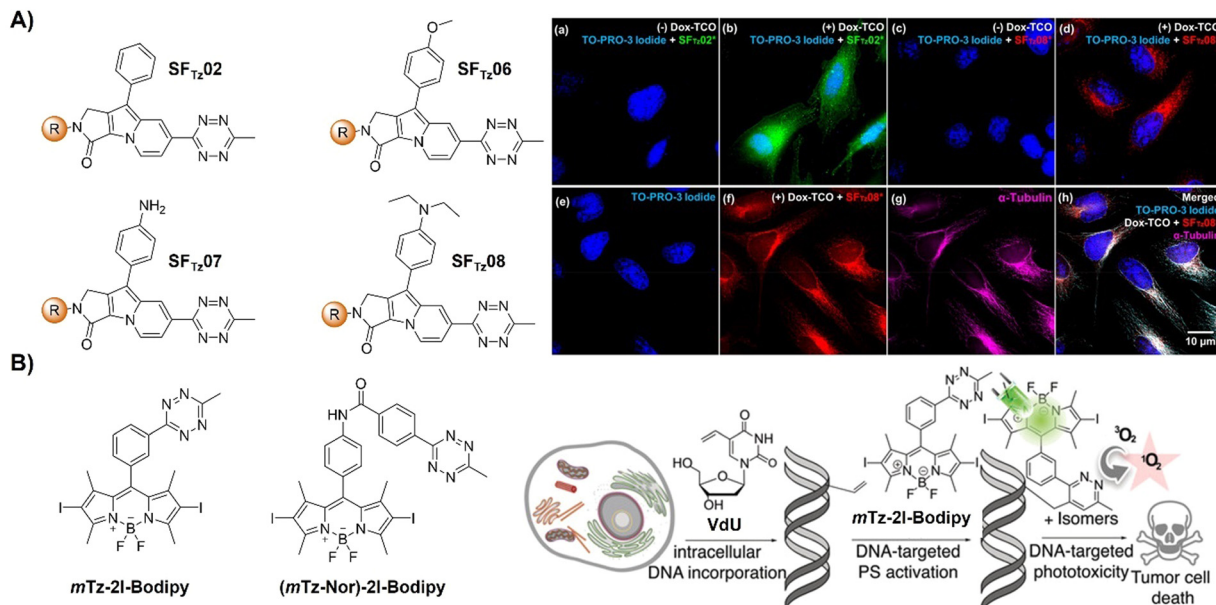


Fig. 16 Bioorthogonal probes designed based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) Molecular structures of SF-Tz conjugates (SF-Tzs) and fluorogenic bioorthogonal imaging of innate microtubules with SF-Tz under fixed-cell conditions, adapted with permission from ref. 122. Copyright 2018, American Chemical Society and (B) the structures of the bioorthogonal, activatable, halogenated Bodipy-Tz derivatives and outline of the DNA-targeted bioorthogonal strategy for activating the phototoxicity, adapted with permission from ref. 123. Copyright 2019, Wiley-VCH GmbH.

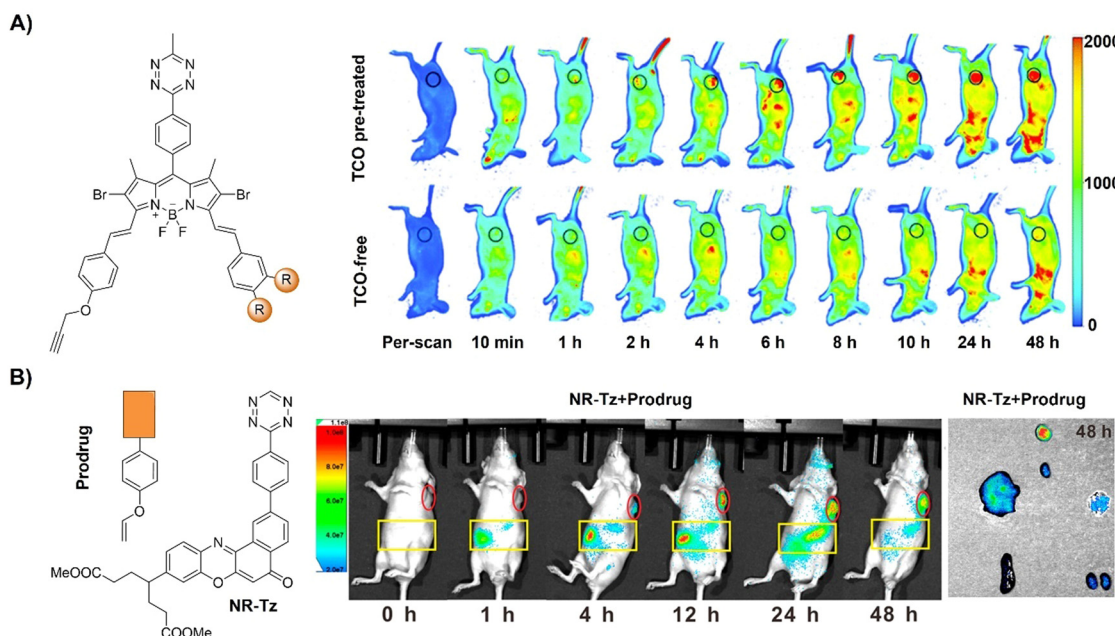


Fig. 17 Fluorescent probes developed based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure of DSBDP and fluorescence images of A431 tumor-bearing nude mice, adapted with permission from ref. 124. Copyright 2019, The Royal Society of Chemistry and (B) the structures of NR-Tz and the prodrug and *in vivo* fluorescence images of tumor and abdominal regions of tumor-bearing mice. *Ex vivo* images of dissected tumors and major organs at 48 h, adapted with permission from ref. 125. Copyright 2019, American Chemical Society.

loaded into lipid nanoparticles separately (Fig. 17B). Accumulation of these lipid nanoparticles in tumors and the subsequent release of the CPT prodrug and NIR-Tz molecules by the tumor triggered IEDDA reaction facilitated chemotherapy and

formation of NIR-Dz with the removal of TBET quenching for tumor fluorescence imaging. This nanosystem is particularly interesting as it allows for monitoring drug release and tumor treatment simply using a bioorthogonal reaction.

Achieving precise signal transduction requires control over the spatial and temporal dynamics of the production of signalling molecules. Phosphatidic acid (PA), a versatile lipid second messenger, exhibits diverse actions influenced by upstream stimuli, biosynthetic origins, and production sites. However, elucidating how cells regulate local PA production to modulate signalling outcomes remains a challenge, as PA biosynthesis sites lack subcellular precision in visualization. In the research lead by Baskin,¹²⁶ a rapid chemoenzymatic method for imaging physiological PA production by phospholipase D (PLD) enzymes was reported (Fig. 18A). Bulky TCO-containing alcohols served as the nucleophilic site in a transphosphatidylation reaction of PLD's lipid substrate, making TCO-containing lipids suitable for fluorescence labelling through an IEDDA reaction with a fluorogenic Tz reagent. Fluorescent reporter lipids were

initially generated in the plasma membrane, and rapid internalization of the fluorescent reporter was observed, suggesting potential applications in probing intracellular phospholipid transport mechanisms. By focusing on the initial 10 s of the IEDDA reaction, precise subcellular localization of endogenous PLD activity triggered by physiological G protein-coupled receptor and receptor tyrosine kinase agonists was possible, implying the capability to track lipid trafficking pathways and the physiological and pathological effects of localized PLD signalling.

The concept of a clickable photocage, involving light to uncage the fluorophore, is an interesting research area for fluorescence bioassays and imaging. A proof-of-concept study for the bioorthogonal click reaction-activatable vinyl Tz-derivatized coumarin photocage system was presented by Kele

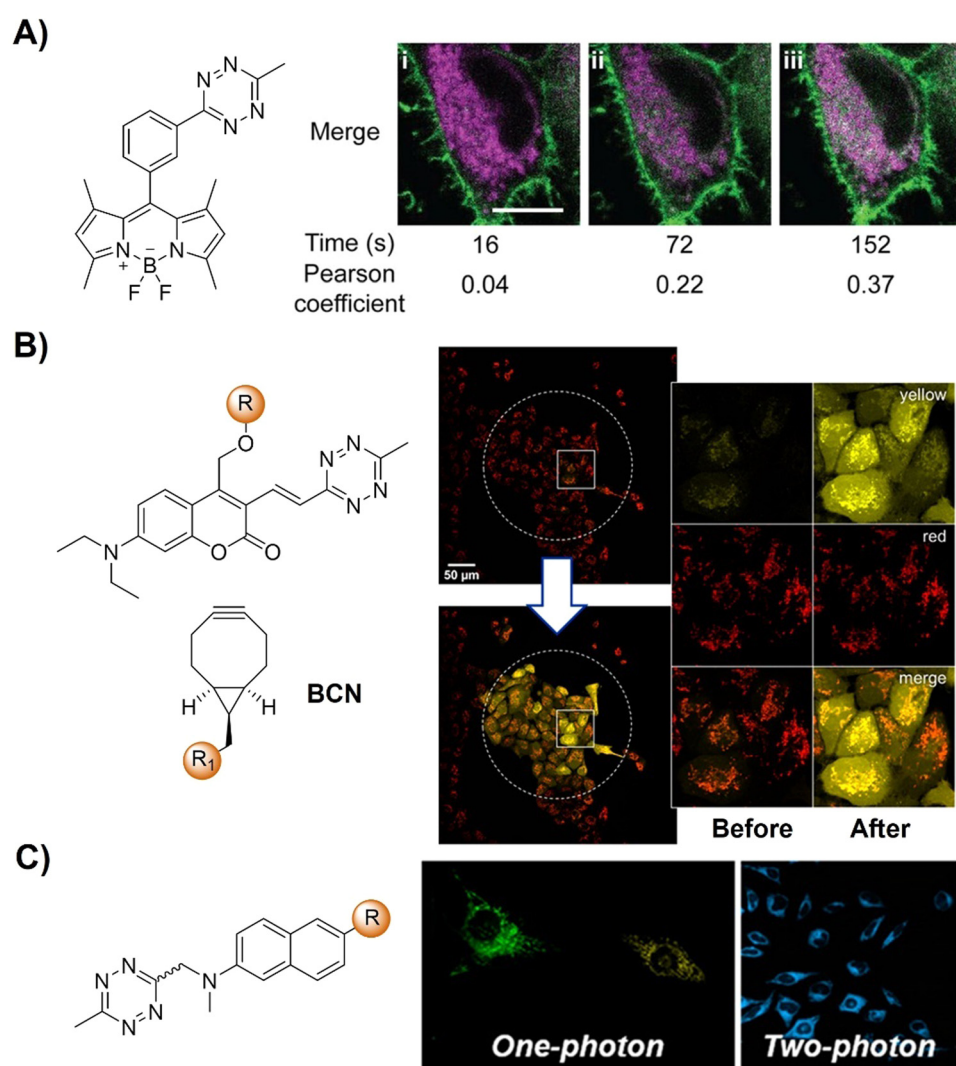


Fig. 18 Bioorthogonal probes developed for fluorescence imaging based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure of the fluorogenic Tz–Bodipy conjugate and representative images of HeLa cells, adapted with permission from ref. 126. Copyright 2019, National Academy of Sciences; (B) the structures of the model photocages and BCNs and cell imaging after irradiation of the central area (marked by a dotted circle) with the built-in blue lamp (460–500 nm, 5 s), adapted with permission from ref. 127. Licensed under CC-BY; and (C) the structural modification of naphthalene-based two- photon fluorogenic dyes and one-photon/two-photon fluorogenic imaging, adapted with permission from ref. 128. Copyright 2020, American Chemical Society.

and coworkers in 2020 (Fig. 18B).¹²⁷ Both experimental data and theoretical calculations indicated that the presence of the bioorthogonal **Tz** motif effectively quenched the excited state of coumarin, necessary for photolysis, thereby disabling photo-responsivity (both photocaging and fluorescence). Upon transformation of the **Tz** moiety in a bioorthogonal click reaction, the sensitivity to light was fully restored. Leveraging bioorthogonal reactions for highly specific cellular targeting, these conditionally activatable photocages offer additional spatial and temporal control over caged compound release. The application of this probe was demonstrated in live cells using a fluorogenic, conditionally activatable construction, which became light-sensitive only when cells were pretargeted with a mitochondria-directed, complementary bioorthogonal function.

Two-photon fluorescent probes^{129,130} that can be excited by two NIR photons have been widely used in fluorescence biosensing and imaging, including the development of two-photon bioorthogonally activated probes.⁸⁸ D-A-type naphthalene fluorophores with moderate-to-good multicolor fluorescence in both one- and two-photon excitation processes are the most widely explored fluorophores for the development of two photon fluorescent probes. Building upon these fluorophores, Park's group¹²⁸ developed a series of **Tz** conjugated multicolor fluorogenic two photon probes (Fig. 18C). These probes exhibited robust turn-on fluorescence response (up to 70-fold) and rapid kinetics with **TCO** in both one- and two-photon fluorescence imaging processes, akin to other FRET-based quenched fluorophores with **Tz**. Modification of either the naphthalene fluorophore with morpholine or **TCO** with TPP allowed for the development of bioorthogonal probes for the precise wash-free fluorescence imaging of lysosomes and mitochondria, respectively, using one-photon fluorescence microscopy.^{120,121} TPP-

TCO-assisted two photon microscopic (TPM) imaging of HeLa cells exhibited rapid turn-on wash-free fluorescence in mitochondria.

Biomarker detection plays an important role in the diagnosis of various diseases, while traditional methods based on ELISA and western blot techniques are time consuming and labor intensive. The fast affinity induced reaction sensor (FAIRS) technique introduced by Wang's group¹³¹ has the potential to address the limitations of traditional methods (Fig. 19A). The probes are comprised of a sandwich ELISA antibody pair, where one antibody is conjugated with a **Tz** based bodipy fluorophore and another is conjugated with azabenzonorbornadiene (AN) for fluorogenic click reaction. Used for IL-6 detection in blood serum and the cell culture medium, FAIRS achieved high sensitivity within 6.5 ± 1.0 minutes, promising broad applicability in medical and clinical diagnostics for rapid biomarker detection.

Super-resolution imaging techniques like single-molecule localization microscopy (SMLM) require active control of fluorescent probes' emissive and dark states. This could be achieved by high power laser illumination, together with redox reagents or UV exposure. Despite high efficiency, the use of UV light irradiation causes phototoxicity for living cells and organisms, thus not suitable for microscopic imaging. Using an established spontaneously blinking silicon rhodamine (**SiR**) scaffold, Wombacher and coworkers¹³² introduced a unique fluorescent probe combining fluorogenic and pH-mediated self-blinking properties, enhancing live-cell localization for SMLM (Fig. 19B). The **Tz** moiety quenches the fluorescence before reaction, facilitating fluorogenic bioconjugation to dienophile-modified cellular targets (e.g. **TPP-BCN**) for mitochondria. The probe, *f*-HM-**SiR**, boasted superior cell permeability, brightness, and photostability, crucial for live-cell imaging. The self-blinking

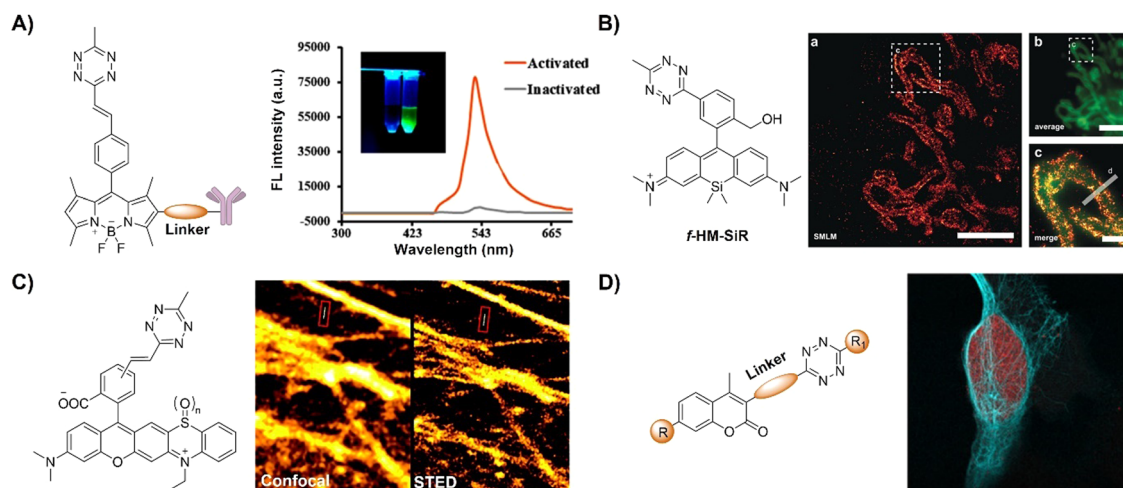


Fig. 19 Fluorescent probes designed based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure of the **Tz** conjugated capture antibody and fluorescence emission spectra in the inactivated/activated state, adapted with permission from ref. 131. Copyright 2020, American Chemical Society; (B) the structure of the *f*-HM-**SiR** and live-cell SMLM imaging, adapted with permission from ref. 132. Copyright 2020, Wiley-VCH GmbH; (C) the structures of **Tz**-functionalized rhodaphenothiazines and confocal/stimulated emission depletion microscopy (STED) super-resolution imaging, adapted with permission from ref. 133. Copyright 2020, The Royal Society of Chemistry; and (D) the general formula of coumarin-**Tz** probes and live cell confocal imaging, adapted with permission from ref. 134. Copyright 2020, Wiley-VCH GmbH.

properties of SiR enabled live-cell SMLM without stabilizing buffers or high excitation power, facilitating long-term, high-resolution intracellular dynamics imaging.

In addition to pH-dependent spontaneous blinking, Kele and colleagues¹³³ devised bioorthogonally applicable **Tz**-functionalized rhoda phenothiazines, elucidating their light-assisted photooxidation-based fluorogenic properties for stimulated emission depletion microscopy (STED) super resolution imaging (Fig. 19C). The experimental findings confirmed efficient sensitization of $^1\text{O}_2$ generation upon green or orange light illumination. This led to self-oxidation of the probes, yielding intensely fluorescent sulfoxide products. Notably, bioconjugated pyridazine exhibited significantly enhanced photo-oxidizability compared to free **Tzs**, resulting in a two-orders of magnitude fluorescence increase. This enabled the selective photoactivation of conjugated probes that were used in actin filament labeling with minimal background fluorescence under wash-free conditions. The “ClickOx (clicked, oxidized)” probes used the same green excitation laser for both probe photooxidation and product excitation, offering added value for STED super-resolution microscopy by excitation at 552 nm with a 660 nm depletion laser.

Through-bond energy transfer (TBET)-based **Tz** probes typically exhibit exceptional fluorescence turn-on responses with various dienophiles, often achieving several thousand-fold enhancement of fluorescence response. For instance, HELIOS (hyperemissive ligation-initiated orthogonal sensing) probes, a series of coumarin-based **Tzs**, exhibited an outstanding 11 000-fold signal enhancement with **TCO** dienophiles.¹⁰⁹ Despite these impressive properties, systematic studies on the design of these coumarin-based probes are essential for developing high quality bioorthogonal probes for precise fluorescence imaging. Towards this aim, Vrabel and coworkers¹³⁴ synthesized and systematically investigated the photophysical properties of seventeen coumarin-**Tz** probes that fluoresce upon reaction with **TCO**- and **BCN**-containing molecules (Fig. 19D). The obtained data confirmed that attaching an azetidine or *N,N*-dimethylpiperazine moiety to the coumarin scaffold improves the fluorescence quantum yield. Click reaction with **TPP-TCO**, **TPP-BCN** or **TCO**-docetaxel and **BCN**-docetaxel allowed for imaging of mitochondria and microtubules in cancer cells.

Abiotic chemical reactions in specific organelles exhibit promise for advancing drug delivery and cell biology and can be achieved by the development of site-specific bioorthogonal probes. In 2021, Vrabel *et al.*¹³⁵ reported a structurally redesigned bioorthogonal **Tz** reagent, enabling the spontaneous accumulation within live mammalian cell mitochondria (Fig. 20A). Optimizing attributes like lipophilicity and positive charge facilitated efficient mitochondrial targeting. Compared to **TCO**-caged prodrugs, these compounds exhibited a 50-fold potency increase upon mitochondrial release and over 100-fold enhancement *versus* unmodified compounds. The transformation of common bioorthogonal reagents into organelle-specific probes in this study thus offers potential for subcellular-level prodrug activation and biological process manipulation using purely chemical methods.

The extensive photophysical characteristics of iridium(III) polypyridine complexes have been employed for biomolecular sensing and imaging. Typically, these complexes possess prolonged triplet excited states, making them exceptional photosensitizing agents for $^1\text{O}_2$ generation for PDT. Expanding on their prior research, Lo *et al.*¹³⁶ introduced iridium(III) **Tz** complexes with diverse phosphorogenic responses and photosensitization attributes upon bioorthogonal reactions with strained alkynes and alkenes (Fig. 20B). These systems were employed for organelle-specific staining and applications involving controlled photocytotoxicity. In addition to commonly used Bodipy, coumarin, and rhodamine frameworks, a series of naphthalimide **Tzs**,¹³⁷ such as LysoNpTz and MitoNpTz, were developed by New as bioorthogonal probes for fluorescence imaging of lysosome and mitochondria using traditional morpholine and TPP as the targeting units (Fig. 20C).

Considering the tissue penetration of light, highly fluorogenic **Tz** bioorthogonal probes emitting at NIR wavelengths are desirable for biomedical imaging. To this aim, Wu *et al.*¹³⁸ introduced fluorogenic Huaxi-Fluor probes where the **Tz** moiety serves as a bioorthogonal partner, quencher, and core of the fluorophore precursor (Fig. 20D). The reaction products, pyridazine derivatives with efficient ICT *via* a thiophene-containing π -bridge, exhibit a large Stokes shift and high quantum yield *in situ*. Modulating the conjugation length and electron pull/push capability fine tunes the emission maximum from 556 to 728 nm. Upon bioorthogonal reaction, a hundred-fold fluorescence increase was achieved, enabling wash-free fluorescence cell imaging. Attaching morpholine and TPP with **BCN** enabled NIR fluorescence imaging of subcellular structures, such as lysosomes and mitochondria, respectively. The application of this bioorthogonal probe was further demonstrated in tumor imaging of a mice model.

Conjugation of Bodipy-**Tz** with hemi-cyanine led to greater than 221 nm redshift of the emission wavelength for NIR emission. Xu and colleagues¹³⁹ thus synthesized an efficient NIR turn-on probe, emitting at 731 nm through indolium incorporation into the Bodipy-**Tz** core (Fig. 20E). A norbornene-modified glucosamine derivative was also prepared for metabolic glycoengineering. Combined together, these derivatives enable rapid and sensitive cell imaging *in vitro* and *in vivo*. The IEDDA reaction leads to a significant fluorescence increase, enabling tumor-specific imaging in tumor bearing mice. Cyanine dyes generally feature NIR emission with high quantum yields. In 2022, Wang and coworkers¹⁴⁰ reported a “torsion-induced disaggregation (TIDA)” phenomenon in a NIR **Tz**-based cyanine probe (Fig. 21A). Upon **Tz**-TOC ligation, the cyanine undergoes TIDA, shifting its heptamethine chain from *S-trans* to *S-cis* conformation. This shift prevents the aggregation of the products of the probe’s bioorthogonal reaction, leading to fluorescence enhancement. In addition to imaging in live cancer cells, the **Tz**-cyanine probe has been used for sensitive tumor imaging in a mouse model. On the other hand, another work conducted by Wu and colleagues¹⁴¹ introduced a molecular design strategy to create **Tz**-based far-red/NIR fluorogenic dyes (Fig. 21B). By



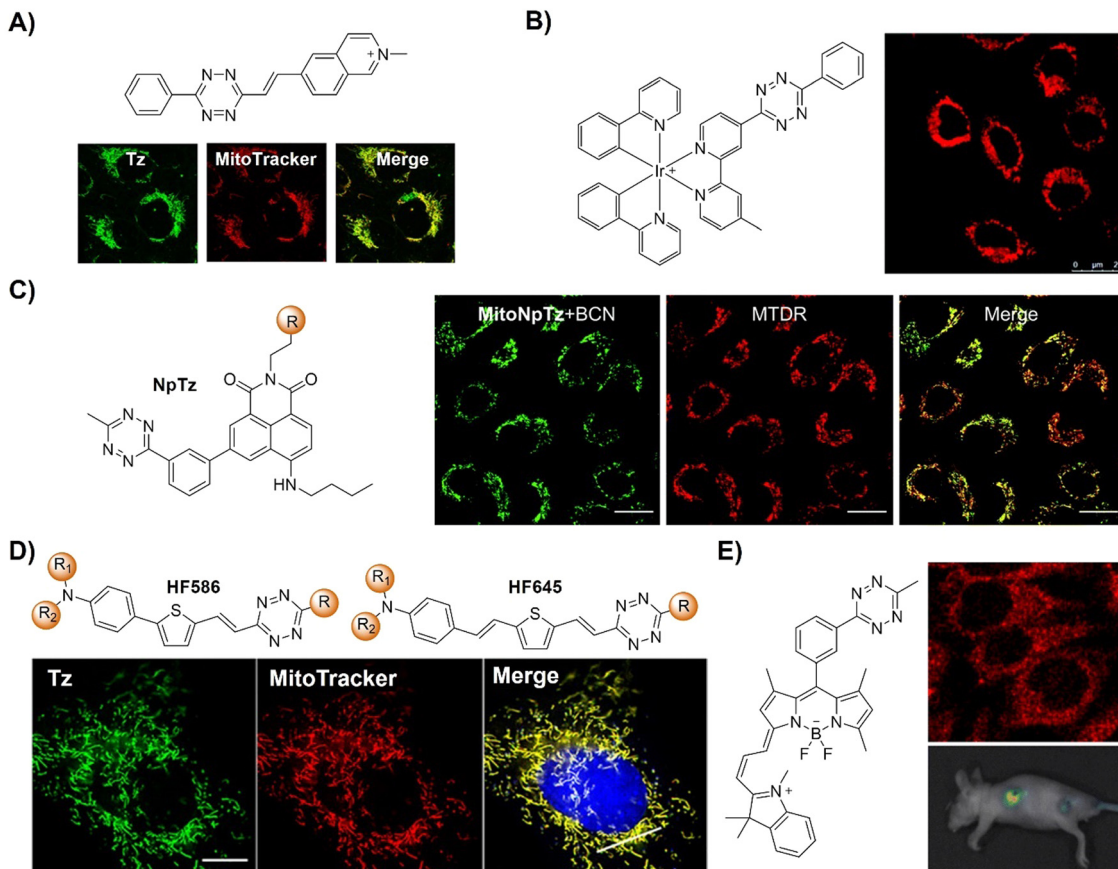


Fig. 20 Fluorescent probes based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure and Mito imaging of the redesigned **Tz** probe, adapted with permission from ref. 135, under the license CC-BY-NC-ND; (B) structures and confocal cell imaging of **Ir-Tz**, adapted with permission from ref. 136. Copyright 2021, The Royal Society of Chemistry; (C) the structures of the naphthalimide tetrazines (Np-Tzs) and representative images of A549 cells, adapted from ref. 137, under the license CC BY-NC 3.0 DEED; (D) the structures of Huaxi-Fluor probes and their application in *in situ* cell imaging, adapted with permission from ref. 138. Copyright 2021, Wiley-VCH GmbH; and (E) the structure of the bodipy-**Tz** core and its application in cell and mice imaging, adapted with permission from ref. 139. Copyright 2022, The Royal Society of Chemistry.

substituting the amino-**Tz** group into the meso-positions of various fluorophores, including rhodamine, Si-rhodamines and cyanine, NIR fluorogenic dyes with remarkable turn-on ratios (≥ 100 -fold) were obtained by harnessing the PeT mechanism. These probes exhibit excellent biocompatibility and photostability, and are well-suited for imaging specific intracellular targets and tumors *in vivo*.

Detection and imaging of specific reactive biomolecules like nitric oxide (NO), one of the key RNS and gasotransmitters, in a localized area in biological samples is essential for understanding their biological functions in their native environment. On the basis of the bioorthogonal concept, *i.e.*, the IEDDA reaction between **Tz** and strained alkyne **BCN**, Xiao and colleagues¹⁴² proposed a versatile toolbox for **NO** detection and imaging of targeted organelles in live cells and zebrafish (Fig. 21C). In their system, rhodamine-*o*-phenylenediamine with a **Tz** unit was used to create the general probe **TMR-Tz-NO**, in which *o*-phenylenediamine served as the **NO** sensing unit and **Tz** served as the bioorthogonal moiety for anchoring the probe in targeted organelles like mitochondria, lysosomes,

and membranes through tethering the **BCN** moiety to various targeting groups (**TPP**, morpholine, and Ac_4ManN). This toolbox enabled **NO** imaging in subcellular locations, as demonstrated by zebrafish experiments for tracking inflammation-induced **NO** production *in vivo*.

Survival, growth, and differentiation of live cells are heavily influenced by the uptake of exogenous nutrients, and the cellular internalization of these exogenous species can be monitored using bioorthogonal probes. For example, employing IEDDA chemistry, the Kasteren group¹⁴³ recently reported a bioorthogonal dual fluorogenic probe for monitoring internalization of extracellular nutrition (Fig. 21D). The non-fluorescent carboxyfluorescein-diacetate-succinimidyl ester (CFSE) was conjugated with **Tz**, a secondary quencher. In this probe system, the diacetate esters can be cleaved by esterase after the probe's internalization into live cells, the succinimidyl ester served as the protein anchoring motif for cytosolic retention, and **Tz** could stain the internalized substrate through the IEDDA reaction, generating enhanced fluorescence for imaging. This advancement facilitated the monitoring of uptake

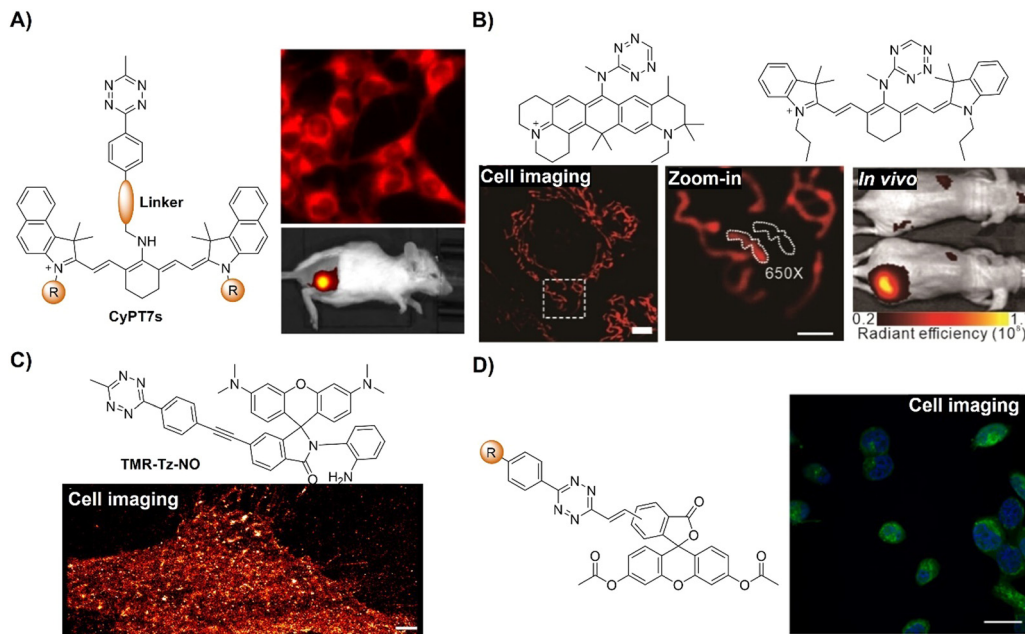


Fig. 21 Bioorthogonal probes based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure of the **Tz**-cyanine core and its application in cell and mice imaging, adapted from ref. 140, under the license CC BY 4.0 DEED; (B) molecular structures and wash-free bioimaging, mice imaging of the representative tetrazine-based probes, adapted with permission from ref. 141. Copyright 2022, Wiley-VCH GmbH; (C) the structure of **TMR-Tz-NO** and its application in plasma membrane super-resolution imaging in living HeLa cells, adapted from ref. 142, Copyright 2022, American Chemical Society; and (D) the structure of the **CFSE-Tz** probe and its application in *in situ* cell imaging, adapted with permission from ref. 143. Copyright 2024, Wiley-VCH GmbH.

of diverse dienophile-containing nutrients in live primary immune cell populations through flow cytometry and live-cell microscopy.

Bioorthogonal chemistry also promoted lipid research by enabling the investigations of lipid uptake and processing in living systems while minimizing detectable pendant group-related biological effects. For example, Kasteren and colleagues¹⁴⁴ introduced sterculic acid, a 1,2-cyclopropene-containing oleic acid analogue, as a bioorthogonal probe for studying unsaturated free fatty acids in live cells (Fig. 22A). The cyclopropene moiety reacted readily with Tz-fluorophore conjugates, facilitating fatty acid localization *via* confocal microscopy. This reaction remained orthogonal to subsequent copper catalyzed Huisgen ligation (CCHL) reactions, enabling the simultaneous study of multiple biomolecules. Further research using sterculic acid as a PTM (post-translational modification) for protein modification suggested that this probe is suitable for investigating protein oleoylation *via* SDS-PAGE and chemical proteomic analysis. This approach, combined with a broadly applicable synthesis of cyclopropene-containing fatty acids, enabled visualization of unsaturated lipids using live-cell microscopy and proteomics.

Selective bioorthogonal labelling of nucleic acids contributes significantly to investigating their chemical biology *in situ*, while steric hindrance from densely packed bioorthogonal functional groups in chromatin could block the chemical modification of nucleic acids in living cells. To overcome this obstacle, Luedtke and coworkers¹⁴⁵ introduced a dual

enhancement strategy for nucleic acid-templated reactions (Fig. 22B). This approach employed a fluorogenic intercalating agent capable of undergoing **IEDDA** with DNA containing 5-vinyl-2'-deoxyuridine (VdU) or RNA containing 5-vinyl-uridine (VU). Reversible high-affinity intercalation of the acridine-Tz conjugate “PINK (Probe for Imaging Nucleosidic AlKene)” dramatically accelerated the reaction rate of **Tz**-alkene **IEDDA** on duplex DNA by 60 000-fold ($590 \text{ M}^{-1} \text{ s}^{-1}$) in comparison with non-templated reactions. Moreover, loss of **Tz**-acridine fluorescence quenching enabled highly fluorogenic detection under wash-free conditions, facilitating dynamic live-cell imaging of acridine-modified nucleic acids during cell division. Using this bioorthogonal labelling approach, imaging of nucleic acids in HeLa cells was achieved.

Apart from DNA, the development of bioorthogonal probes for RNA labelling is particularly promising for advancing disease diagnosis and treatment monitoring. To facilitate the RNA labelling in live cells, which was not achieved by conventional methods, Qian and colleagues¹⁴⁶ introduced a fluorogenic system for precise mRNA labelling and imaging in live cells by combining hybridization chain reaction (HCR) with proximity-induced bioorthogonal chemistry (Fig. 22C). This strategy offered spatial resolution of low-expression targets, activating **Tz**-quenched fluorophores through HCR-directed bioorthogonal reactions for fluorescence turn-on and amplification. Additionally, a biocompatible nucleic acid delivery system derived from black phosphorus nanosheets (BPNSS) was employed for efficient imaging of cytosolic mRNAs in live



cells. The platform's flexibility in targeting RNA of interest facilitated its use as a versatile tool for RNA tracking in living cells.

Most **Tz**-based bioorthogonally activated fluorogenic probes encounter aggregation-caused quenching (ACQ) issues. To address these issues, AIEgens have been explored for the development of bioorthogonal probes for fluorescence imaging using IEDDA. For example, Kim and colleagues¹⁴⁷ found that **Tz** could quench the emission of the Kaleidolizine (**Klz**) AIEgen through TBET, and thus bioorthogonally activated probes **Klz_{Tz}** were designed and synthesized (Fig. 22D). Bioorthogonal click reaction with **TPP-TCO**, together with AIE, facilitated wash-free fluorescence imaging of mitochondria in live cells using **Klz_{Tz}**. In another example, Tian and coworkers¹⁴⁸ devised a series of novel **Tz**-modified tetraphenylenes (**TPEs**) as bioorthogonally activated AIE fluorogenic probes (Fig. 22E). Both fluorescence and AIE properties were quenched by **Tz** *via* a TBET mechanism and then reactivated upon **Tz**-to-pyridazine conversion through the IEDDA reaction. The resulting cycloadducts exhibited significant fluorescence enhancement, large Stokes shift, high fluorescence quantum yield, and AIE activity. Adjusting the length and position of the π -linker finely tunes a probe's photophysical properties, while an excessively planar π -linker induced AIE-to-ACQ transformation. Bi-tetraethyl-substituted probes exhibited higher turn-on ratios due to a "double-quenched" function. Using double-clickable linkers, fluorescent macrocycles

were formed, enabling organelle-specific bioorthogonal wash-free imaging in live cells suitable for applications in biomedicine.

Most naphthalimide derivatives, particularly unsubstituted naphthalimides, exhibit ACQ, while some naphthalimide derivatives with appropriate substitution exhibit AIE properties. For example, Tian and coworkers¹⁴⁹ introduced a range of TBET-based naphthalimide-**Tz** probes with broad bioorthogonal applicability and fluorescence activation properties (Fig. 22F). The **IEDDA** cycloadducts exhibited exceptional photophysical traits, demonstrating AIE characteristics. Modifying the π -bridge linker between the naphthalimide and **Tz** enabled precise control over emission, spanning from blue to red, while also influencing the AIE attributes. These **Tz**-based probes facilitated effective wash-free fluorogenic protein labeling and high signal-to-noise ratio mitochondrial imaging (click reaction with **BCN-TPP**) in live cells.

Two-photon excitation, by shifting absorption to the red, significantly enhances the signal-to-noise ratio and reduces photodamage, enabling imaging approximately 10 times deeper than with confocal microscopy. Despite this, effective two-photon excitable fluorogenic probes are scarce. Mahuteau-Betzer *et al.*¹⁵⁰ introduced fluorogenic probes comprising a two-photon excitable fluorophore and a **Tz** quenching moiety (Fig. 23A). These probes exhibit impressive kinetics (up to $1.1 \times 10^3 \text{ M}^{-1} \text{ s}^{-1}$) and emit in the red with moderate to high turn-on ratios. The best candidate achieved a substantial 13.8-fold

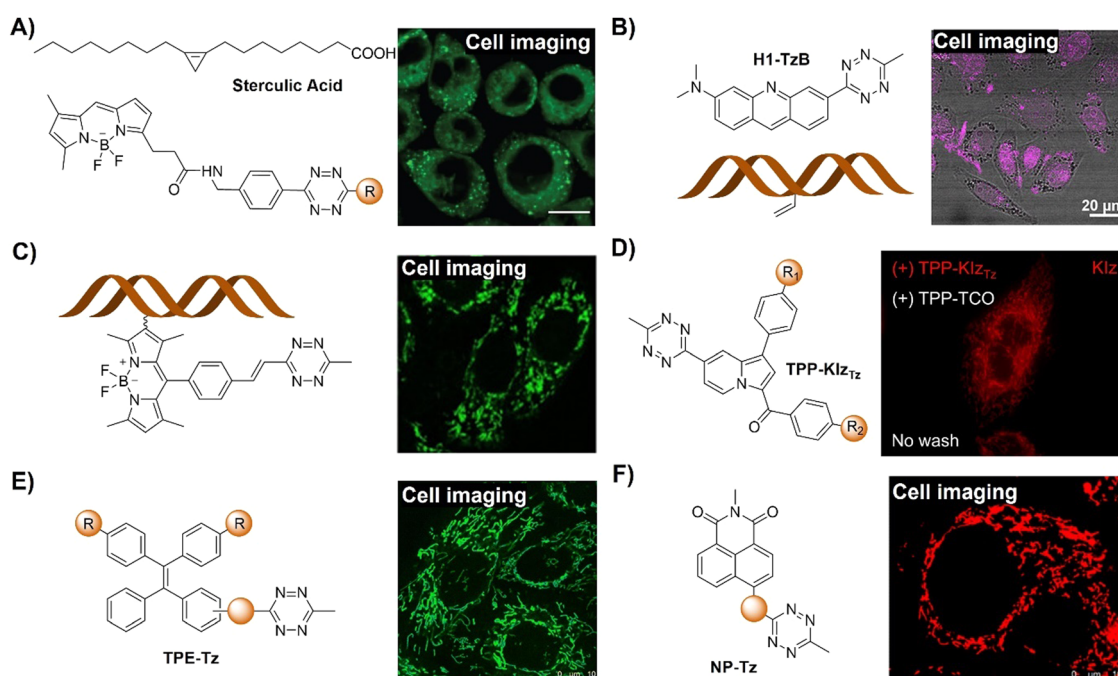


Fig. 22 Bioorthogonal probes based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure of **Tz**-fluorophore conjugate/sterculic acid and its application in cell imaging, adapted from ref. 144, under the license CC BY-NC 4.0 DEED; (B) the structure of acridine-**Tz** conjugate (**H1-TzB**)/vinyl-DNA and wash-free imaging of nucleic acids in HeLa cells, adapted with permission from ref. 145. Copyright 2022, Wiley-VCH GmbH; (C) the structure of the **Tz**-quenched fluorophore and its application in RNA labelling in live cells, adapted from ref. 146, Copyright 2022, American Chemical Society; (D) the structure of the **Klz_{Tz}** probe and its application in wash-free cell imaging, adapted with permission from ref. 147. Copyright 2021, Elsevier B.V.; (E) the structure of the **Tz**-modified tetraphenylenes and its application in wash-free cell imaging, adapted from ref. 148, Copyright 2022, The Royal Society of Chemistry; and (F) the structure of the naphthalimide-**Tz** probe and its application in bioimaging in live HeLa cells, adapted from ref. 149, Copyright 2022, The Royal Society of Chemistry.

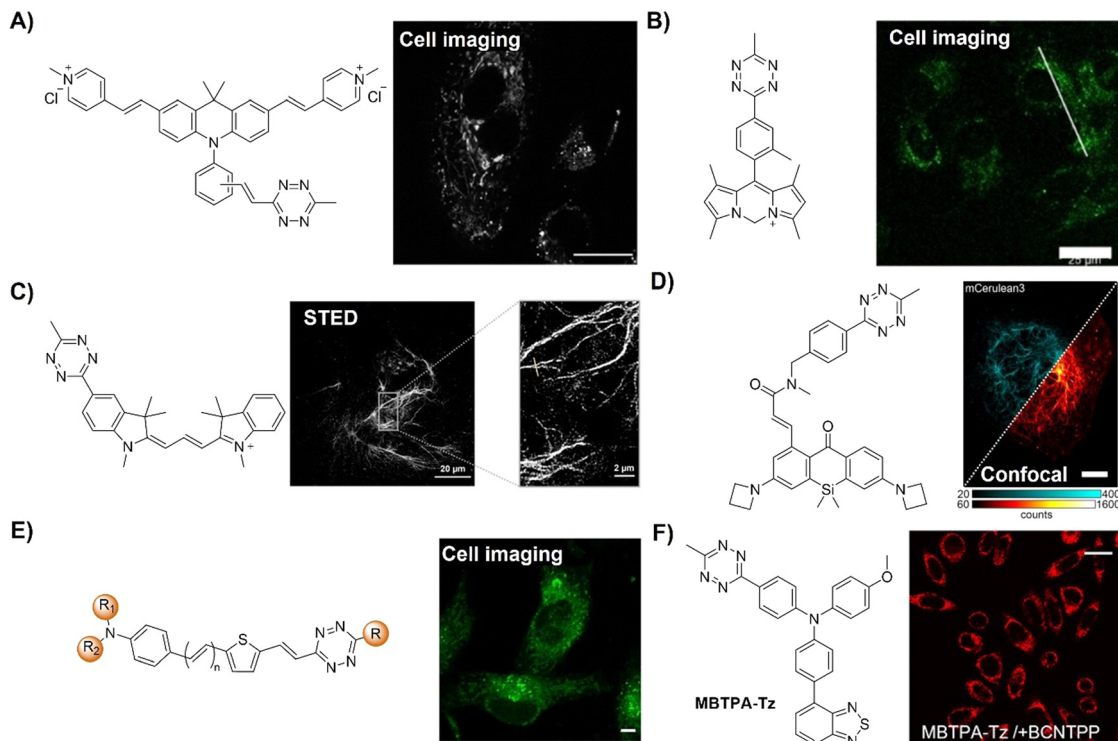


Fig. 23 Inverse-electron-demand Diels–Alder reaction-based bioorthogonal probes using the TBET fluorescence response mechanism. (A) The structure of a two-photon excitable **Tz**-fluorophore conjugate and its application for the imaging of live A549 cells, adapted from ref. 150, under the license CC BY 3.0 DEED; (B) the structure of a C-DIPY-**Tz** conjugate and wash-free imaging in live cells, adapted with permission from ref. 151. Copyright 2023, Wiley-VCH GmbH; (C) the structure of the **Tz**-functionalized Cy3 derivative and its application in STED imaging in live cells, adapted from ref. 152, under the license CC BY 3.0 DEED; (D) the structure of the **Tz**-xanthone (**PaX**) core fluorophore probe and its application in confocal/super resolution imaging, adapted from ref. 153, under the license CC-BY 4.0; (E) the structure of the **Tz**-isonitrile probe and its application in wash-free cell imaging, adapted from ref. 154, Copyright 2024, Wiley-VCH GmbH; and (F) the structure of the AIE-**Tz** probe (**MBTPA-Tz** as an example) and its application in bioimaging, adapted from ref. 155, Copyright 2024, under the license CC BY 4.0 DEED.

turn-on fluorescence response under one-photon excitation, exceeding the 3-fold threshold for high contrast wash-free live-cell imaging. Notably, under two-photon excitation, an 8.0-fold turn-on was observed. The high two-photon brightness (>300 GM) of the resulting adduct allowed for the use of low laser power suitable for *in vivo* imaging.

Carbon-dipyrromethenes (C-DIPYs) are a new class of fluorophores exhibiting intense emissions from green to NIR.¹⁵⁶ Using C-DIPY as a signaling unit, Ng and colleagues¹⁵¹ synthesized a water-soluble bioorthogonal probe *via* a two-step procedure from a known carbonyl pyrrole dimer (Fig. 23B). Similar to other probes, the emission of this probe was quenched by a **Tz** moiety at pH 7.4 PBS by TBET and could be switched on after click reaction with **BCN** derivatives, resulting in up to 6.6-fold restoration of fluorescence. This bioorthogonal activation was demonstrated in U-87 MG human glioblastoma cells, with fluorescence intensity increasing by 8.7-fold post-incubation with the **BCN** derivative. Interestingly, in contrast to previous reports on C-DIPYs that are mitochondria targeting due to their cationic nature, this probe preferential localizes in lysosomes, which could be attributed to an energy-dependent endocytic cellular uptake pathway.

Cell surface and intracellular labelling and imaging can be achieved by the development of bioorthogonally activated

probes that are cell membrane impermeable and permeable, respectively. Kele and coworkers¹⁵² synthesized three series of **Tz**-functionalized Cy3 derivatives using varied design approaches and explored how the bioorthogonal unit influences the fluorescence of yellow-emitting indocyanines (Fig. 23C). The employed strategies included direct conjugation, phenylene-linked units (TBET design) and vinylene-conjugated systems (π -extended direct conjugation), each comprising membrane-permeable and non-permeable derivatives. Fluorogenic characteristics upon reaction with **BCN** were assessed *via* IEDDA and the data indicated that **Tz** quenching efficiency is dependent on the distance between **Tz** and the fluorophore, confirming the efficacy of internal conversion-based modulation over through-bond energy transfer. The directly conjugated probes effectively labeled genetically bioorthogonalized extra- and intracellular proteins in live cells for STED super-resolution imaging of target structures.

Harnessing **Tz** dyads' inherent excited state quenching to control fluorophore activation, Hell and colleagues¹⁵³ reported a series of highly compact, click compatible and photoactivatable bioorthogonal probes for super resolution imaging (Fig. 23D). In this probe system the quencher and bioorthogonal click unit **Tz** was conjugated with the photoactivatable xanthone (**PaX**) core fluorophore. These probes were able to



be used in most nanoscopies, including STED, PALM, and MINFLUX. Leveraging MINFLUX's ultimate resolution, precise visualization of vimentin filament substructure in cellulose was achieved through incorporating clickable unnatural amino acids in the filament monomers' head domain with molecular-scale precision.

Creating fluorogenic probes that can simultaneously label multiple targets in live cells is of great importance for a better understanding of the complex cellular processes. The emerging [4+1] cycloaddition between **Tz** and isonitriles offers potential as a bioorthogonal tool for multiplex labelling and fluorescence cell imaging. For instance, the Wu group¹⁵⁴ introduced a range of **Tz**-functionalized bioorthogonal probes for multiplexing and live cell bioimaging (Fig. 23E). By integrating pyrazole adducts into the fluorophore frameworks, the post-reacted probes exhibited impressive fluorescence turn-on ratios, achieving up to 3184-fold enhancement in fluorescence. These modification strategies are adaptable for the development of probes using other fluorophores, enabling the design of probes showing a broad emission spectrum ranging from 473 to 659 nm. In imaging applications, these probes were found to be able to simultaneously label multiple targets in live cells for wash-free fluorescence imaging.

The impact of molecular aggregation on emission properties was included in the design of **Tz**-functionalized bioorthogonal probes with increased emission enhancement ratio (I_{AC}/I_{BC} , where I_{AC} and I_{BC} stand for fluorescence intensities after and before click reactions, respectively) for labeling and imaging of multiple cellular organelles. As an example of AIEgen-based probes, Tang and coworkers¹⁵⁵ demonstrated that an exceptionally high I_{AC}/I_{BC} can be achieved in aggregated systems when **Tz** is combined with AIEgens (AIE-**Tz**, Fig. 23F). **Tz** enhanced its quenching efficiency upon aggregation, remarkably reducing background emissions. Subsequent click reactions induced changes in **Tz**'s chemical structure and prompted AIE, resulting in significantly enhanced I_{AC}/I_{BC} . The potential of the ultra-fluorogenic systems for selective imaging of multiple organelles was confirmed in the mitochondria, plasma membranes, and lipid droplets of live cells.

Previous research conducted by the Lo group has confirmed that the iridium(III) complex with a built-in **Tz** moiety could serve as a bioorthogonal probe for luminescence biolabeling and imaging.¹¹⁸ Together with the excellent photo- and sonosensitizing performance of metal complexes, Cai and colleagues created activatable “off-on” sonosensitizers with enhanced sonodynamic activity and immunogenic cell death (ICD) induction capability for precise and potent tumor sonotherapies (Fig. 24A).¹⁵⁷ The non-luminescent iridium(III) complexes that are loaded into pH sensitive human serum albumin (HSA) nanoparticles could be released in an acidic microenvironment found in tumors and then activated and anchored to cancer cell membranes after click reaction with BCN receptors. With ultrasound irradiation, cytotoxic ROS was produced for tumor cell ablation through oxidative stresses. Membrane damage induced highly immunogenic PANoptosis, boosting ICD to elicit robust systemic antitumor immunity

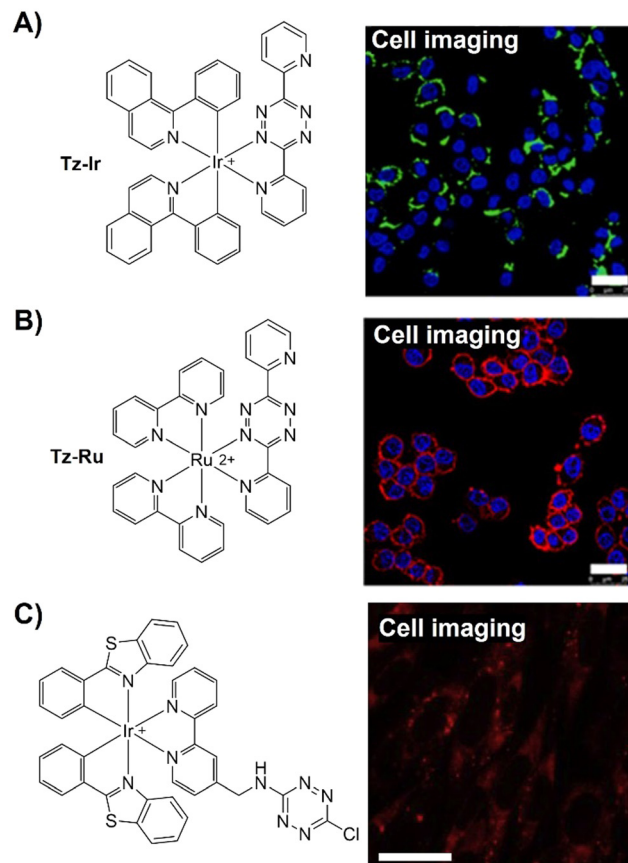


Fig. 24 Transition metal complex-based bioorthogonal probes based on the inverse-electron-demand Diels–Alder reaction using the TBET fluorescence response mechanism. (A) The structure of a **Tz**–Ir complex and its application in *in situ* cell imaging, adapted with permission from ref. 157. Copyright 2024, Elsevier Ltd; (B) the structure of a **Tz**–Ru complex and wash-free imaging in live cells, adapted with permission from ref. 158. Copyright 2024, American Chemical Society; and (C) the structure of the **Tz**–Ir complex and its application in cell imaging, adapted with permission from ref. 159. Copyright 2016, Elsevier Ltd.

against primary/distant tumors and lung metastasis. Based on a similar approach, a ruthenium(II) complex with a built-in **Tz** moiety was developed by the same group for bioorthogonal activation and sonodynamic therapy with boosted antitumor immunity (Fig. 24B).¹⁵⁸

2.3.2. FRET approach. In addition to the TBET approach, the FRET approach has also been used for the design of IEDDA probes based on luminophores including metal complexes, bodipy, coumarin, rhodamine and fluorescein. For example, Lo's group¹⁵⁹ introduced a versatile synthetic method for iridium(III)-**Tz** complexes, serving as phosphorogenic biomaterials for bioorthogonal labeling and multicolor imaging (Fig. 24C). Unlike typical luminescent cyclo-metallated iridium(III) polypyridine complexes, these **Tz**-bearing complexes exhibited minimal emission due to efficient FRET and/or PeT from the excited iridium(III) polypyridine unit to the attached **Tz** moiety. However, upon interaction with the strained alkyne BCN-OH and its BSA conjugate BCN-BSA, the iridium(III) complexes were activated, achieving an emission enhancement up



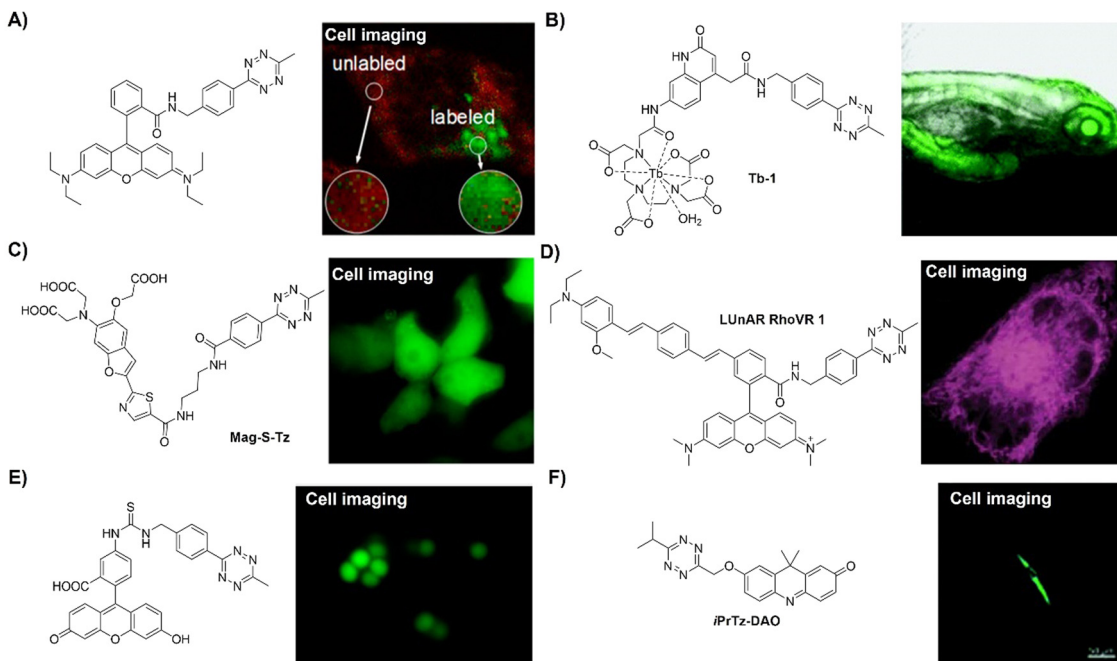


Fig. 25 Inverse-electron-demand Diels–Alder reaction-based bioorthogonal probes using the FRET fluorescence response mechanism. (A) The structure of the **Tz**-rhodamine derivative and photoluminescence lifetime confocal microscopic images of living HeLa cells, adapted with permission from ref. 160. Copyright 2019, American Chemical Society; (B) the structure of the **Tz**-terbium complex and luminescence imaging in zebrafish, adapted with permission from ref. 161. Copyright 2020, The Royal Society of Chemistry; (C) the structure of **Mag-S-Tz** and its application in cell imaging, adapted with permission from ref. 162. Copyright 2016, American Chemical Society; (D) the structure of the **Tz**-quenched RhoVR derivative and its application in confocal imaging, adapted with permission from ref. 163. Copyright 2022, American Chemical Society; (E) the structure of the **Tz** tag and its application in wash-free *S. aureus* bacteria imaging, adapted with permission from ref. 164. Copyright 2020, American Chemical Society; and (F) the structure of the **Tz**-caged probe and its application in bioimaging in live HSF cells adapted with permission from ref. 165. Copyright 2022, The Royal Society of Chemistry.

to 1133.7 fold. The probe was then used for the luminescent labelling, imaging and flow cytometric analyses of CHO-K1 cells.

On the other hand, the Huang group¹⁶⁰ reported a highly sensitive and specific “labeling after recognition” sensing strategy where luminophore labeling and then biological recognition were proposed (Fig. 25A). Based on the FRET from the iridium(III) complex (donor) to rhodamine (acceptor) a bioorthogonal probe was developed for luminescence and lifetime detection and imaging of caspase-3. The donor and acceptor FRET pair was linked using a caspase-3 sensitive linker, the DEVD tetrapeptide. From the luminescence lifetime analysis, it was found that the iridium(III) complex's emission lifetime was quenched by FRET to the rhodamine and then restored after caspase-3 promoted DEVD cleavage. Imaging of caspase-3 catalytic activity in apoptotic cells was achieved *via* photoluminescence lifetime imaging microscopy, confirming intracellular bioorthogonal bis-labeling and catalytic cleavage dynamics. This study presents a typical example of the “labeling after recognition” sensing strategy and a luminescence lifetime based bioorthogonal probe.

In comparison with metal complexes with prolonged emission lifetime at hundreds of nanoseconds, rare earth lanthanide chelates generally have a longer emission lifetime at microsecond levels. The prolonged emission lifetime makes the developed bioorthogonal probes suitable for background-

free time-resolved luminescence bioassay and imaging as the lifetime of autofluorescence is generally less than 10 ns. In this context, Wong *et al.*¹⁶¹ introduced a stable, biocompatible terbium complex, Tb-1, enabling bioorthogonal ligation with engineered cell-surface glycans, yielding responsive luminescence (Fig. 25B). They strategically incorporated a bioorthogonal luminescence resonance energy transfer (LRET) quencher into a tripodal terbium complex with low toxicity. Upon the IEDDA reaction, the electronic structure of the quencher changed, resulting in enhanced terbium emission intensity. The prolonged lifetime (approximately 1 ms) and bright emission ($\Phi \sim 10\%$) allowed for background-free luminescence imaging in A549 cancer cells and zebrafish.

Metal ions, such as sodium (Na^+), potassium (K^+), magnesium (Mg^{2+}), iron (Fe^{3+}) and calcium (Ca^{2+}), are essential for biological functions, thus monitoring levels of essential metal ions is crucial for disease diagnosis. Bioorthogonally activated probes have been developed for visualizing localization of these metal ions, such as Mg^{2+} . The Buccella group¹⁶² devised a two-step method to localize a fluorescent Mg^{2+} sensor within specific organelles (Fig. 25C). This method involved a rapid reaction between a Tz-functionalized pro-sensor (**Mag-S-Tz**) and a genetically encoded HaloTag fusion protein. Protein conjugation preserved the sensor's metal-binding properties, with a suitable dissociation constant ($K_d = 3.1 \text{ mM}$) for detecting intracellular Mg^{2+} concentration. This bioorthogonally



activated probe could detect Mg^{2+} in target organelles of HEK 293T cells, offering improved spatial resolution without interference.

Electric potential differences across lipid bilayers are fundamental in cellular physiology. In contrast to plasma membranes, transmembrane potentials of intracellular organelle bilayers, including those of the mitochondria, lysosomes, nuclei, and the endoplasmic reticulum (ER), are difficult to investigate using traditional methods like patch-clamp electrophysiology. Recently, Miller's group¹⁶³ introduced LUNAR RhoVR (Ligation Unquenched for Activation and Redistribution Rhodamine-based Voltage Reporter) for optically monitoring endoplasmic reticulum (ER) membrane potential changes in living cells (Fig. 25D). By pairing Tz-quenched RhoVR with TCO-conjugated ceramide (Cer-TCO) for ER targeted labelling, specific and bright fluorescence was observed, enabling voltage-sensitive imaging and validation of plasma-ER membrane functional coupling through a K^+ diffusion mediated change of ER membrane potential.

Both **BCN** and **TCO** are essential linkers for the IEDDA reaction with **Tz** in bioorthogonal "click" ligation. To enhance the availability of **TCO** reagents, Zanda and colleagues¹⁶⁴ synthesized a series of **TCO** linkers derived from *trans,trans*-1,5-cyclooctadiene for biomolecule conjugation (Fig. 25E). These linkers facilitated bioorthogonal ligation with **Tz** tags, offering linker flexibility crucial for stability and ligation efficiency, as exemplified by BSA. Labelling of biomolecules in cellular lysates and live bacteria like *S. aureus* was used to demonstrate their application in bioorthogonal ligation, showing promise for live-cell imaging. Moreover, the short PEG linker **TCO** enabled surface decoration of monoclonal

antibodies like rituximab and obinutuzumab with functional handles and minimal mAb aggregation.

Wu and coworkers¹⁶⁵ devised a series of fluorescent probes by exploring isonitrile-**Tz** click-to-release mechanism-based bioorthogonal activation (Fig. 25F). Efficient quenching of canonical fluorophores was observed across the visible spectrum using the **tBuTz**-cage, with substantial fluorescence activation upon isonitrile-induced cleavage. DNA and RNA templated chemistry accelerated this bioorthogonal cleavage reaction at low concentrations, suggesting potential for miRNA detection. Orthogonality between the **tBuTz**-caged probe cleavage and **iPrTz**-caged probe and **TCO** species cleavage was demonstrated, facilitating their use in live cells. This research using two orthogonal cleavage reactions in live cells has provided options for bioorthogonal fluorogenic probes in multi-target detection.

The distance between **Tz** and the fluorophore is one of the key points that should be taking into consideration when developing bioorthogonal probes for fluorescence labelling and imaging.^{144,166–168} On the basis of this consideration, Wombarcher's group¹⁶⁹ synthesized a series of fluorescence bioorthogonal probes with a close proximity **Tz**-fluorophore conjugate for wash-free live cell super resolution imaging (Fig. 26A). These probes were designed using a series of red and far-red fluorogenic **Tz** dyes (**HDyes**) for bioorthogonal labelling. The short distance between the **Tz** and fluorophores allowed for effective fluorescence quenching through an energy transfer process. **HDyes** displayed high fluorescence and was suitable for wash-free live-cell imaging, which was demonstrated in two-color labeling experiments and STED super-resolution imaging of UAA-labeled target proteins. The modular synthetic approach

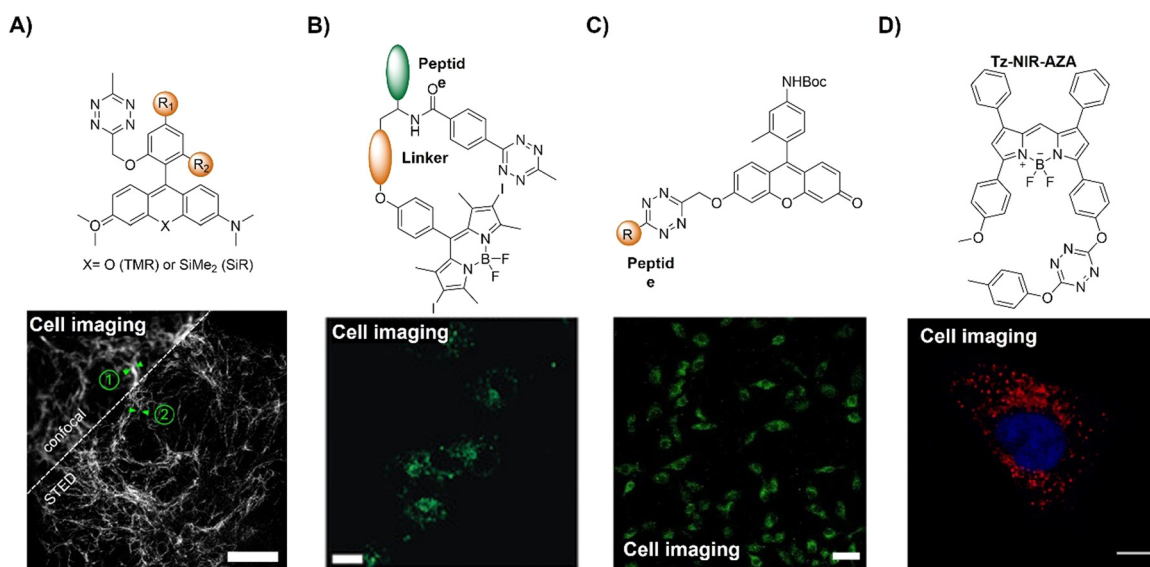


Fig. 26 Fluorescent probes based on the inverse-electron-demand Diels–Alder reaction using the FRET fluorescence response mechanism. (A) The structures of **HDyes** and live-cell confocal and STED imaging of COS-7 cells, adapted from ref. 169, under the license CC-BY-NC-ND 4.0; (B) the structure of **Tz**-bodipy and wash-free imaging in live HeLa cells, adapted with permission from ref. 170. Copyright 2020, Wiley-VCH GmbH; (C) the structure of the peptide–**Tz** conjugate and its application in cell imaging, adapted from ref. 171, under the license CC BY 4.0 DEED; and (D) the structure of **Tz**-substituted bodipy (**Tz**-NIR-AZA as an example) and its application in HeLa Kyoto cell imaging, adapted with permission from ref. 172. Copyright 2023, Elsevier Ltd.



yielded a self-blinking **HDye**, useful for long-term live-cell STED imaging and super-resolution optical fluctuation imaging (SOFI).

Typically, small molecules lack specificity in targeting cellular compartments, often distributing across multiple areas or randomly without achieving desired localization. Vázquez *et al.*¹⁷⁰ introduced a modular peptide-based platform separating the halogenated Bodipy photosensitizer (PS) and the quencher **Tz**, enhancing photochemical properties and enabling spatial activation within specific organelles upon peptide conjugation (Fig. 26B). Various dual-labeled peptides, differing only in peptidic vectors but sharing the same PS/quencher pair, controlled confined photodynamic effects *via* bioorthogonal *in situ* activation in specific subcellular organelles. For example, **Tz**-bodipy, activated specifically in the cytoplasmic membrane through bioorthogonal IEDDA, exhibited superior PDT performance, enhancing photodynamic effects at lower concentrations without dark toxicity. Compatible with common bioconjugation strategies, this versatile cysteine-based approach facilitated the exploration of diverse subcellular compartments and peptidic functionalities, advancing bioorthogonally activatable PS applicability.

In addition to the PDT,^{157,158} bioorthogonal chemistry has also enabled the delivery of a drug in the form of a prodrug into specific diseased tissues with minimal drug release in other tissues or side effects.^{125,171,173} For instance, Wu's group¹⁷¹ introduced 3-vinyl-6-oxymethyl-**Tz** (**voTz**) as a reagent for crafting **Tz**-caged prodrugs and selectively labeling peptides, generating bioorthogonally activatable peptide-prodrug conjugates (Fig. 26C). These stable prodrugs were bound specifically to

target cells, aiding cellular uptake. Subsequent bioorthogonal cleavage reactions activated the prodrug, significantly inhibiting tumor cell growth while ensuring remarkable biosafety for normal cells. *In vivo* trials using a tumor-bearing mice model confirmed the therapeutic efficacy and biosafety of this prodrug design strategy.

Bioorthogonal probes with NIR emission can also be obtained using bodipy derivatives as fluorophores. For example, Wu and coworkers¹⁷² explored the use of **Tz**-substituted BF_2 -azadipyrromethene probes (bodipy derivatives) and model **Tzs** for their reactivity with strained alkyne and alkene dienophiles, aiming to identify the optimal probe for cellular imaging applications (Fig. 26D). **BCN** exhibited superior reactivity, with reactions finishing within 15 min. The **Tz**-substituted BF_2 -azadipyrromethene displayed intense fluorescence at 715 nm as the cycloaddition progressed in an aqueous solution, with a remarkable 43-fold enhancement. Such good efficiency prompted its application in bioorthogonal cell imaging, allowing for dynamic imaging of the reaction in subcellular regions of live cells upon pre-incubation with a 2-deoxy-glucose-alkyne dienophile.

2.3.3. Other approaches. Apart from the TBET and FRET approaches, there are other mechanisms that have been applied to the design of IEDDA activated probes. Guo and coworkers¹⁷⁴ introduced a fluorogenic bioorthogonal reaction, employing the **IEDDA** reaction between styrene (an unstrained alkene) and a simple **Tz**-dipyridine structure (Fig. 27A). This reaction yielded a distinctive 4-phenyl-3,6-di(pyridin-2-yl)-1,4-dihydropyridazine (**PDHP**) fluorophore with solvatochromic emission. By genetically incorporating a styrene-derived

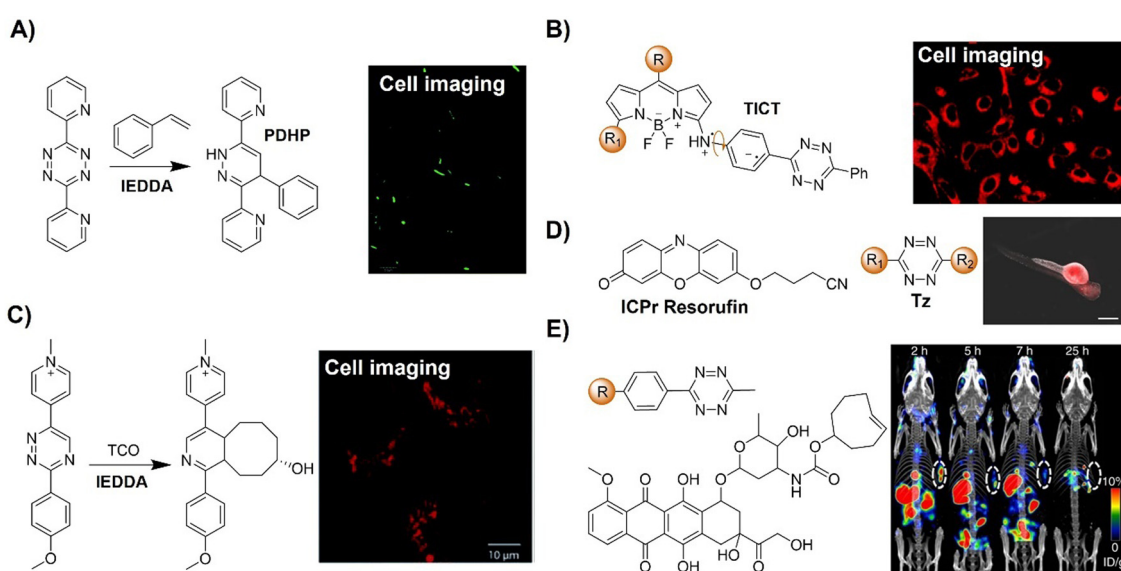


Fig. 27 Inverse-electron-demand Diels–Alder reaction-based bioorthogonal probes for fluorescence imaging. (A) The fluorogenic reaction between styrene/Tz and its application for selective labeling of *E. coli* cells, adapted from ref. 174, under the license CC BY 3.0 DEED; (B) the structure of **Tz**-bodipy, fluorescence quenching via the TICT process and its application in cell imaging, adapted with permission from ref. 175. Copyright 2022, The Royal Society of Chemistry; (C) fluorogenic reaction between **TCO** and 1,2,4-triazines and its application in cell imaging in live U2OS cells, adapted from ref. 176, under the license CC BY 3.0 DEED; (D) the structure of **ICPr-resorufin/Tz** and its application in zebrafish imaging, adapted with permission from ref. 177. Copyright 2018, American Chemical Society; and (E) the structures of the **TCO**-caged prodrug (**TCO-Dox**) and **Tz**-bearing **NapK(Tz)YF** and *in vivo* mice imaging, adapted from ref. 178 under the license CC BY 4.0 DEED.

unnatural amino acid (**KStyr**), site-specific and fluorogenic protein labeling in a wash-free manner was achieved in both cells and bacteria. He *et al.*¹⁷⁵ introduced highly efficient red-emitting fluorogenic probes with a twisted intramolecular charge transfer (TICT) mechanism for fluorescence quenching (Fig. 27B). TICT involves an electron transfer phenomenon upon photoexcitation, typically featuring an electron donor and acceptor linked by a single bond. Exploiting the correlation between the bodipy 5-position substituent's electron effect and fluorescence quantum yield, a stable, red-emitting fluorogenic probe was developed. Upon reaction with **TCO** and cyclopropane, the probe exhibited a remarkable 103-fold fluorescence enhancement. This probe enabled the wash-free imaging of live cells and exhibited suitability for long-term storage.

Recently, 1,2,4-triazines have emerged as versatile dienes in **IEDDA** reactions with strained dienophiles. In comparison with **Tz**, 1,2,4-triazines exhibit lower reactivity towards both **TCO** and **BCN** although they are highly stable under biological conditions. After a systematic investigation to understand the mechanism of reaction of various 1,2,4-triazines with **TCOs** in the **IEDDA** reaction, Vrabel's group discovered a strong dependence on the substitution patterns of both substrates (Fig. 27C).¹⁷⁶ This exploration led to the discovery of new *N*-alkyl pyridinium triazines with exceptional properties for bioconjugate chemistry. An efficient, modular synthetic strategy was then used to construct diverse heterobifunctional probes based on the pyridinium 1,2,4-triazine scaffold. The fluorogenic nature (up to 50-fold enhancement with 650 nm emission) of the reactions of these heterodienes with **TCOs** allowed for fluorogenic cell labeling.

Dissociative bioorthogonal reactions enable precise control over the release of bioactive agents and reporter probes. Franzini *et al.*¹⁷⁷ identified 3-isocyanopropyl (**ICPr**) substituents as removable masking groups within biological systems (Fig. 27D). The 3-isocyanopropyl derivatives are able to react with **Tzs** to yield 3-oxopropyl groups for bioorthogonal labeling. The study highlighted rapid reaction kinetics, achieving near-complete liberation of phenols and amines under physiological conditions. Compatibility with living organisms was confirmed by the release of the resorufin fluorophore and the mexiletine drug in zebrafish embryos implanted with **Tz**-modified beads. Due to their benefits including ease of synthesis, swift kinetics, diverse leaving groups, high release yields, and structural compactness, **ICPr** derivatives have emerged as appealing chemical caging moieties for applications in chemical biology and drug delivery.

A synergistic prodrug activation system was developed by Gao and coworkers¹⁷⁸ using a combination of **EISA** and **Tz/TCO** bioorthogonal decaging reactions (Fig. 27E). The system achieved spatiotemporal targeting and selective activation of prodrugs within cancer cells, crucial for enhancing chemotherapeutic selectivity. A **Tz** moiety was linked to the **EISA** motif to trigger inverse-electron-demand Diels-Alder (inv-DA) reaction-mediated chemical decaging of a **TCO**-caged effector molecule intracellularly. Cancer cells with elevated phosphatase levels underwent intracellular **EISA** via **Tz**-bearing **NapK(Tz)YF**,

leading to significant **Tz** accumulation and subsequent liberation of the **TCO**-caged prodrug (**TCO-Dox**), inducing cancer cell death while sparing normal cells. The increased emission of the bioorthogonal reaction allowed for imaging of cancer cells as well as the Dox release. This approach enhanced the selectivity of Dox delivery for cancer cells, offering potential for reducing adverse drug effects and demonstrating efficacy in a xeno-grafted cervical cancer mice model.

The **IEDDA** reaction between **Tz** and **TCO** offers precise control over bioactive agents and imaging probe release. For example, Bernardes *et al.*¹⁷⁹ employed a pre-targeted activation strategy using single-walled carbon nanotubes (SWCNTs) bearing **Tzs** (**Tz@SWCNTs**) and a **TCO**-caged molecule to deliver active effector molecules for tumor imaging (Fig. 28A). Optimized *in vivo* fluorescence imaging was achieved using a new fluorogenic NIR probe, **tHCA**, that could be activated *via* bioorthogonal chemistry, enabling tumor imaging in mice. This approach allowed for selective Dox prodrug activation for tumor chemotherapy and real-time fluorescence imaging in living cells. Combining **tHCA** with a pre-targeted bioorthogonal approach achieved non-invasive, high target-to-background ratio tumor visualization in a xenograft mice model, highlighting the potential of **Tz**-functionalized SWCNTs for targeted bioorthogonal decaging with minimal off-site activation.

In another example, a bioorthogonally activated drug delivery system (DDS) was obtained by Gao's research group,¹⁸⁰ and the bioorthogonal prodrug activation allowed for drug release in a precise imaging manner (Fig. 28B). This self-reporting bioorthogonal prodrug activation system was developed using fluorescence emission to interpret activation events. **Tz**, acting within ROS-responsive supramolecular assemblies, served both as a fluorescence quencher and as a prodrug activator. The subsequent **IEDDA** reaction with *trans*-cyclooctene-paclitaxel (**TCO-PTX**) facilitated the release of fluorescence and active drugs concurrently, enabling a linear relationship for quantifying **PTX** release. This fluorescence-based visualizable drug delivery system is promising for elucidating drug delivery processes and determining prodrug activation efficacy.

Senescent cell targeted bioorthogonal **Tz** ligation has contributed to the clearance of these cells in the body, slowing down the aging process as well as promoting anticancer treatment. For example, Wang's group¹⁸¹ presented a senolytic approach using senescent cell-sensitive bioorthogonal **Tz** ligation. In this system, a galactose (Gal) moiety was engineered into dihydrotetrazine, serving as both a recognition element for senescence-associated β -galactosidase (SA- β -gal) and a caging group for **Tz** (Fig. 28C). Gal-**Tz** efficiently released fluorescent hemicyanine (HCA) for the precise imaging and release of Dox from **TCO**-caged prodrugs for targeting and eliminating senescent HeLa and A549 cells over non-senescent cells with a 16.44 senolytic index. This approach was then used for the selective activation and delivery of proteolysis-targeting chimeras (PROTACs) as senolytics, potentially offering an efficient intervention for cell senescence by degrading targeted proteins catalytically. This research established bioorthogonal **Tz** ligation as a viable method for selectively removing senescent cells.



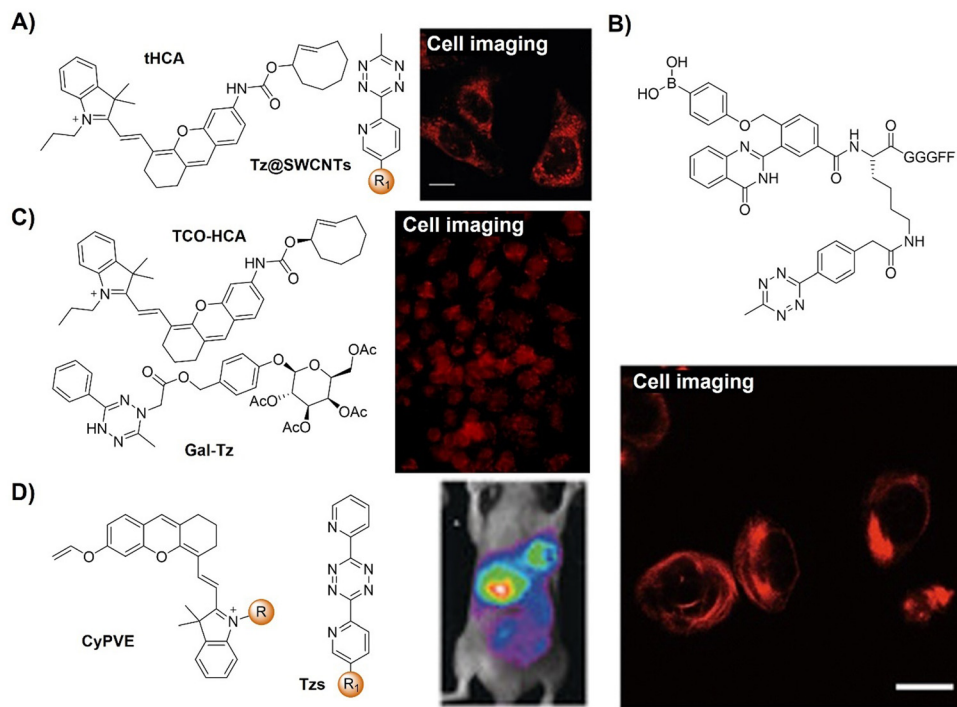


Fig. 28 Bioorthogonal probes for fluorescence imaging based on the inverse-electron-demand Diels–Alder reaction. (A) The structure and cell imaging of tHCA/Tz@SWCNTs, adapted from ref. 179, under the license CC BY 4.0 DEED; (B) the structure of the Tz-prodrug and its application in cell imaging, adapted with permission from ref. 180. Copyright 2022, The Royal Society of Chemistry; (C) the structure of TCO-HCA/Gal-Tz and its application in cell imaging, adapted with permission from ref. 181. Copyright 2024, Wiley-VCH GmbH; and (D) the structures of the CyPVE/Tzs and *in vivo* fluorescence images of tumor-bearing nude mice, adapted with permission from ref. 182. Copyright 2020, Wiley-VCH GmbH.

Bioorthogonal prodrug activation using exogenous click reactions also offers a solution for precise drug release with fluorescence turn-on response for efficient monitoring, overcoming challenges of current approaches based on endogenous receptors or overexpressed enzymes. For example, prodrug activation was developed by Wang and colleagues,¹⁸² in which the drug release was achieved synergistically by using a bioorthogonal reaction and the acidic tumor microenvironment (Fig. 28D). This approach uses a Tz-functionalized pH-sensitive polymer micelle, which disassembles in an acidic environment, triggering a bioorthogonal click reaction to activate PDT. The ROS generation could be specifically activated by Tzs through a bioorthogonal click reaction to trigger NIR fluorescence signals for precise tumor imaging and phototoxicity for cancer cell ablation, allowing for imaging-guided PDT. This design ensures Tz groups remain inert in circulating micelles but become active upon micellar disassembly in tumors, enabling specific prodrug activation. This method, exploiting the dysregulated pH common in cancers, offers a universal approach for bioorthogonal prodrug activation in acidic tumor microenvironments.

In another research, Ng and coworkers¹⁸³ developed a versatile bioorthogonal strategy enabling targeted delivery and site-specific activation of photosensitizers for precise tumor imaging and antitumoral PDT (Fig. 29A). This approach featured an isonitrile-caged distyryl boron dipyrromethene-based photosensitizer, NC-DSBDP, activated

through converting the *meso* ester substituent to carboxylate *via* the [4+1] cycloaddition with a Tz derivative. Using galactose or GE11 peptide-conjugated Tzs (gal-Tz and GE11-Tz), cancer cells expressing the asialoglycoprotein receptor or epidermal growth factor receptor were selectively labeled. Internalized NC-DSBDP underwent ester-to-carboxylate transformation triggered by these Tzs, activating fluorescence for imaging and ROS generation within the target cells for cancer cell ablation. *In vivo*, bioorthogonal activation led to effective tumor eradication in nude mice.

Despite the successful translation of PDT in cancer therapy, precisely controlling the administration of photosensitizers into a body is extremely hard, thus leading to side effects as the excess photosensitizers in the body could be activated for ROS generation under sunlight irradiation. The patients are thus required to stay in the dark for a few days after treatment. Interesting research led by the Ng group¹⁸⁴ reported a bioorthogonal strategy to mitigate photosensitivity post-PDT (Fig. 29B). In this protocol, BCN-Q, consisting of a BCN moiety that swiftly reacts with the Tz substituent of the photosensitizer PS-Tz, and the BHQ-3 quencher deactivate the PS-Tz photosensitizer through a bioorthogonal click reaction. As such the BHQ-3 quencher efficiently reduces the emission of PS-Tz and the generation of ROS *via* FRET in PBS and diverse cancer cell lines. In nude mice, BCN-Q injection effectively neutralizes the residual photosensitizer without compromising PDT efficacy, evidenced by minimal skin damage post-treatment. This approach



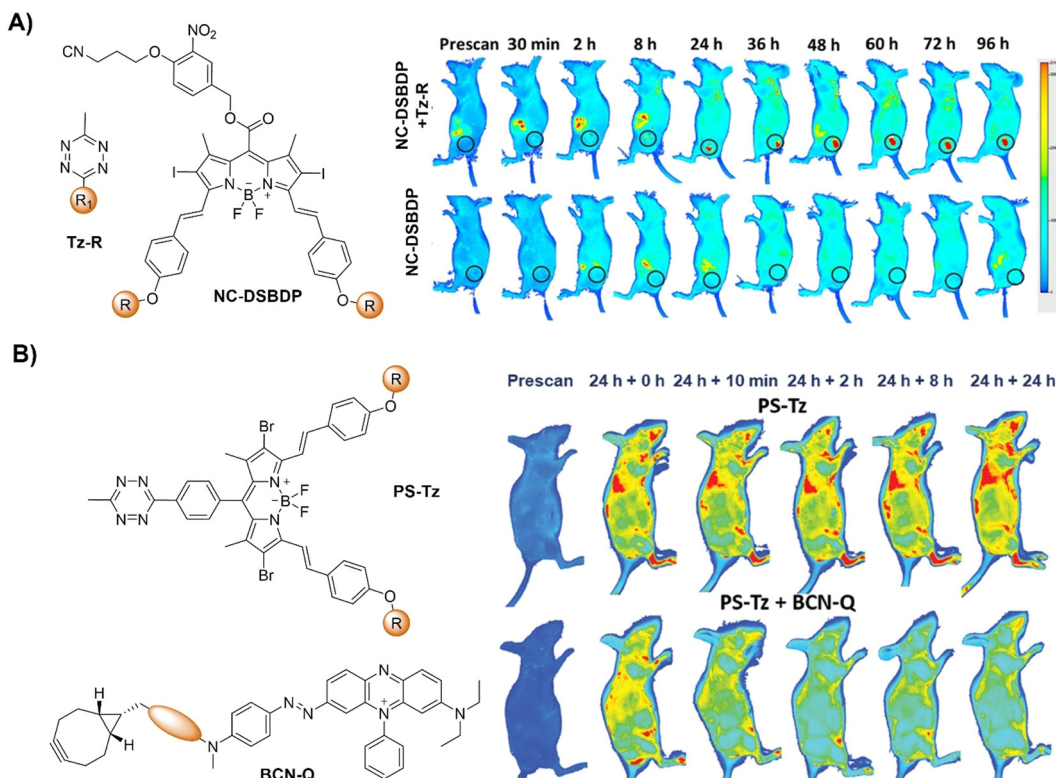


Fig. 29 Bioorthogonal probes for *in vivo* fluorescence imaging based on the inverse-electron-demand Diels–Alder reaction. (A) The structures of **Tzs** and **NC-DSBBDP** and *in vivo* fluorescence images of A431 tumor-bearing nude mice, adapted with permission from ref. 183. Copyright 2023, Elsevier B.V. and (B) the structures of **BCN-Q** and **PS-Tz** and NIR fluorescence images of the nude mice, adapted with permission from ref. 184. Copyright 2023, Wiley-VCH GmbH.

thus offered a direct “click-and-quench” mechanism, circumventing the need for tumor-specific photosensitizers and potentially enhancing the clinical viability of PDT by mitigating photosensitivity.

Spatiotemporal multiplexed labelling and imaging of multiple cellular targeted species, enabled by bioorthogonally activated probes, facilitates comprehensive profiling of the functions of these species in live cells. Carlson and co-workers¹⁸⁵ reported scission-accelerated fluorophore exchange (SAFE), a technique enabling spatiotemporal multiplexed immunofluorescence imaging of live cells (Fig. 30A). SAFE leveraged rapid IEDDA bioorthogonal click chemistry to swiftly remove immunofluorescence signals from labeled cells' surfaces. Strategies include multiparameter analysis ($n \geq 14$), non-disruptive imaging of murine peripheral blood mononuclear cells and bone marrow cells to study cellular differentiation, and longitudinal multiplexed imaging of bone marrow progenitor cells maturing into neutrophils over 6 days and real-time multiplexed cycling of living mouse hepatic tissues. SAFE is expected to have broad utility for investigating physiological dynamics in living systems.

Bioorthogonal reactions have been particularly useful for nucleic acid detection and imaging due to their specificity and adjustable reaction kinetics. For instance, Jiang and colleagues¹⁸⁶ reported a fluorogenic sensor for the precise RNA imaging in live cells by integrating a bioorthogonal

reaction with a DNA cascade reaction (Fig. 30B). In this probe system, two DNA hairpin probes were synthesized using Tzs or vinyl ether-caged fluorophores. Upon targeted mRNA-induced catalytic hairpin assembly, spatial proximity facilitated efficient bioorthogonal reactions, unmasking the vinyl ether group to activate fluorescence. This sensor exhibited high signal-to-noise ratios (~30 fold) and sensitive mRNA detection (4.6 pM detection limit). The “click to release” mechanism allowed for minimal background signals in biological environments for mRNA imaging in live cells and differentiated mRNA expression levels in tumor and normal cells. In another study led by Rentmeister,¹⁸⁹ a benzylic linker promoted methyltransferase catalyzed norbornene transfer reaction was employed for rapid bioorthogonal Tz ligation for the specific labelling of DNA and RNA in cell lysate.

A fluorescent probe for *in situ* lysosome labeling and lysosomal pH sensing was developed by Xu and coworkers¹⁸⁷ using fluorescence bioorthogonal reaction between 4-(6-methyl-1,2,4,5-tetrazin-3-yl) benzoic acid (TE) and norbornenes (NC). By pre-treatment of cells with morpholine-modified norbornene (**MP-NC**), lysosomes rapidly became decorated with functional substrates for bioorthogonal reactions (Fig. 30C). With the introduction of 4-(6-methyl-1,2,4,5-tetrazin-3-yl) benzoic acid (TE), lysosome-targeted probe **LP1** was constructed in live cells within 10 min, emitting bioorthogonal fluorescence at 515 nm.

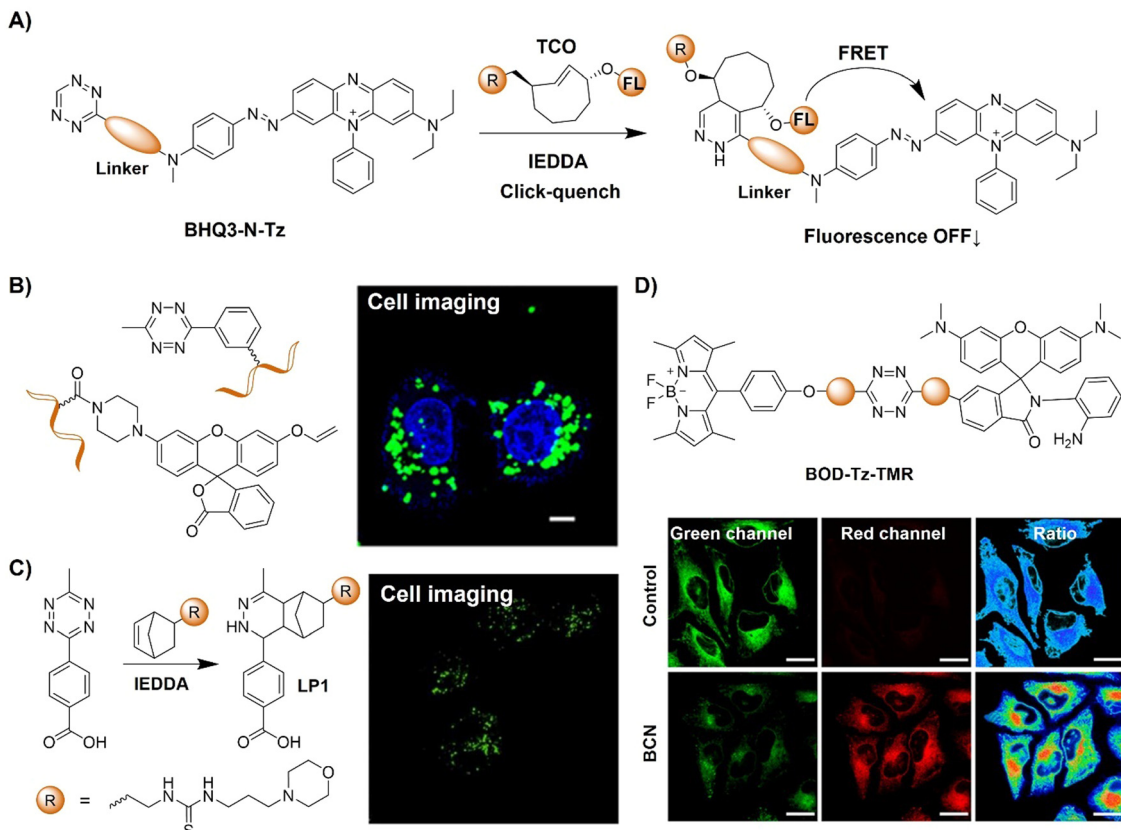


Fig. 30 Fluorescent probes designed for cell imaging based on the inverse-electron-demand Diels–Alder reaction. (A) Schematic illustration of the SAFE system; (B) the structures of reaction precursors (**Tz** and vinyl ether-caged fluorophore) and mRNA imaging in live cells, adapted with permission from ref. 186. Copyright 2022, American Chemical Society; (C) bioorthogonal reaction of TE-norbornenes and bioimaging of the MCF-7 cells, adapted with permission from ref. 187. Copyright 2022, Elsevier B.V; and (D) the structure of **BOD-Tz-TMR** and ratio imaging of HeLa cells, adapted with permission from ref. 188. Copyright 2023, American Chemical Society.

A **Tz**-based bioorthogonal probe for **NO** detection was developed by Xiao and coworkers¹⁸⁸ and the application of this probe for the detection of **NO** in atherosclerosis (AS) was demonstrated (Fig. 30D). AS is characterized by plaque formation in blood vessels, leading to severe cardiovascular conditions with varied levels of **NO**. In this probe system (**BOD-Tz-TMR**), bodipy and rhodamine with a well-documented **NO** sensing unit were linked using a **Tz** moiety, resulting in emission quenching of both fluorophores. Click **BCN** reaction allowed for switching on of the fluorescence of Bodipy, and sensing of **NO** led to the restoration of the rhodamine's emission. The bioorthogonal reactions with effective FRET from bodipy to rhodamine enabled the ratiometric fluorescence detection and imaging of **NO**. This sensor selectively targeted AS plaques, visualizing endogenous **NO** at two lesion stages in AS mouse models. Ratiometric signals confirmed **NO** involvement during AS, confirming increased **NO** generation with lesion progression. This bioorthogonally activated probe thus offered a useful approach for targeted delivery of small molecular probes to AS plaques for reactive biomarker identification and detection.

The rapid kinetics inherent in **Tz** ligation make it suitable for applications in the precise fluorescence imaging of live

cells. However, the stability of **Tzs** in aqueous solutions or in the presence of redox species varies (e.g. thiols and ROS), with the most reactive **Tzs** often being the least stable. For example, Li's research has identified that **Tz** could respond to superoxide ($O_2^{\bullet-}$), the primary ROS in mammalian cells.⁹⁹ Using 1,2,4,5-**Tz** as a superoxide-responsive trigger, a series of fluorescent probes for $O_2^{\bullet-}$ detection using various fluorophores with emission color modulation were developed. These probes exhibit exceptional specificity and fluorescence amplification for $O_2^{\bullet-}$ detection, enabling organelle-resolved multiplexed imaging. Abnormal $O_2^{\bullet-}$ generation in myocardial ischemia/reperfusion injury and high-content screening for superoxide homeostasis mediators has been investigated using these probes.

To tackle challenges associated with the poor stability of **Tz**, various strategies have been proposed.¹⁹⁰ For instance, the Fox group introduced a photochemical variant to this reaction, resulting in a new type of photoclick reaction named photo-induced **Tz** linkage (**photo-IEDDA**).^{191–194} Dihydrotetrazine (**DHTz**) was used as a precursor for this strategy, along with photosensitizers (such as methylene blue) that generate 1O_2 under light irradiation (>600 nm). These species can oxidize **DHTz** *in situ* to form **Tz**, which can then react with **TCO** to form



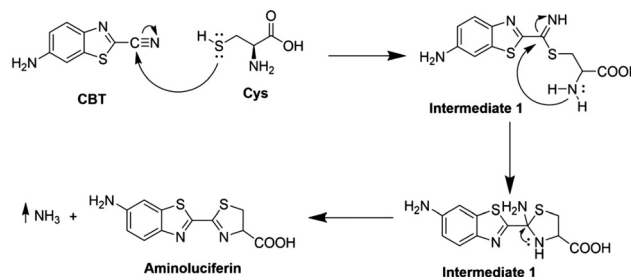
a cycloadduct. Additionally, disubstituted **Tzs**, particularly monomethyl **Tzs**, exhibit improved stability without significantly compromising reactivity. Alternatively, Prescher *et al.* have used 1,2,4-triazines in **IEDDA** reactions, which exhibit enhanced stability compared to the corresponding **Tzs** while maintaining satisfactory kinetics amongst various dienophiles.¹⁹⁵ On the other hand, Vrabel *et al.*¹⁹⁶ introduced triazinium ligation, a bioorthogonal conjugation based on *N*1-alkyl 1,2,4-triazinium salt reaction with strained alkynes. Optimized alkylation of 1,2,4-triazines yielded key triazinium reagents, facilitating tunable reaction rates. Alkylation boosted reactivity by three orders of magnitude, which was validated by DFT calculations. *tert*-Butylation has emerged as the optimal modification, offering highly reactive and stable salts. Triazinium ligation, orthogonal to strain-promoted azide–alkyne cycloaddition, efficiently modifies biomolecules, exhibiting enhanced cell permeability and intracellular labeling efficiency.

Optimization of dienophile structures strikes a balance between improved stability and reactivity. **TCO**, which has been extensively studied as a dienophile, can isomerize to less reactive *cis*-cyclooctenes, posing a challenge for bioorthogonal reactions.¹⁹⁷ To address this, the Fox group devised dioxolane-fused *trans*-cyclooctenes with increased stability and reduced propensity for isomerization. Other dienophiles employed in **IEDDA**-mediated ligation include norbornenes, cyclooctynes, and methylcyclopropenes. The Bonger research group introduced vinylboronic acids as strained dienophiles, notable for their water solubility and hydrolytic stability as well as high reactivity.^{198,199} These compounds engaged in rapid *in vivo* **Tz** ligation with stable precursors by coordination of the boronic acids to the phenol group, followed by swift intramolecular **IEDDA** reactions. With the continued effort of researchers, we are confident that the **IEDDA** reaction will find even more applications in a broader range of scenarios.

2.4. 1,2-Aminothiol-CBT click reaction

The biocompatible click reaction between *D*-cysteine and 2-cyanobenzothiazole (**2-CBT**), inspired by fireflies, has received considerable interest in the development of bioluminescent probes for bioassays and imaging. In nature, the reaction between *D*-cysteine and **2-CBT** is the final step in the synthesis of firefly luciferin (*D*-luciferin, *D*-Luc). In the presence of luciferase, Mg^{2+} , and ATP, *D*-Luc undergoes oxidation, emitting bioluminescence. In 2009, Rao and colleagues introduced a method to label proteins with terminal cysteines using **CBT** probes.⁴¹ This reaction,^{41,200–202} exhibiting click-like properties, proved to be simple and modular and required no catalysts or chromatographic purification. Its natural origin ensured excellent biocompatibility, with both reactants displaying good water solubility and selectivity.

Furthermore, the second-order rate constant, measured as $9.1\text{ M}^{-1}\text{ s}^{-1}$, surpassed that of the Huisgen click reaction by two orders of magnitude.⁴² Subsequent studies elucidated the mechanism of this click reaction. As illustrated in Scheme 4, the reaction between *D*-cysteine and 6-amino-2-cyanobenzothiazole (**NCBT**) at pH 7.4 shows the sulphydryl



Scheme 4 The proposed mechanism of the 1,2-aminothiol-CBT click reaction.

group's higher nucleophilicity than the amino group. This leads to the formation of intermediate 1, followed by the formation of a thiazole ring structure and the release of ammonia gas, resulting in the formation of the final product amino luciferin. During the past few decades, the **CBT-Cys** click reaction has found diverse applications in protein labeling, molecular imaging (encompassing optical imaging, magnetic resonance imaging, nuclear imaging, and photoacoustic imaging), tumor therapy, and various other fields. This section will focus on the latest advancements in this click reaction for *in situ* bioimaging, aiming to provide profound and comprehensive insights for its further implementation.

The exploration of *in vivo* molecular imaging is fundamental to unraveling the complexities of health, injury, aging, and disease by identifying specific biochemical processes like enzymatic activity, small-molecule fluxes, and post-translational modifications. As such probes that can be used for dual analyte detection *in situ* in living organisms are more desirable for understanding the multiple biochemical processes in a complicated environment. To this end, the Chang group²⁰³ developed a strategy for dual-analyte detection in live animals by leveraging the real-time generation of firefly luciferin from two complementary caged precursors sensitive to distinct biochemical targets (Fig. 31A). In this strategy, Peroxy Caged Luciferin-2 (**PCL-2**), a boronic acid probe is responsive to hydrogen peroxide (H_2O_2), and a peptide-based probe, *z*-Ile-Glu-ThrAsp-*D*-Cys (**IETDC**) can be activated by active caspase 8. Upon encountering the corresponding stimuli, **PCL-2** released 6-hydroxy-2-cyanobenzothiazole (**HCBT**), and **IETDC** liberated *D*-cysteine, resulting in the formation of firefly luciferin *in situ* and emitting bioluminescence only when both triggers are concurrently active. This system operates akin to AND-type molecular logic gates, providing insights into the simultaneous presence of H_2O_2 and caspase 8. Selective imaging of either H_2O_2 or caspase 8 activity was successfully demonstrated in both laboratory settings and live subjects. Furthermore, application of **PCL-2** and **IETDC** *in vivo* unveiled a simultaneous surge in H_2O_2 and caspase 8 activity during acute inflammation in live mice. In another study, the Dubikovskaya group²⁰⁴ introduced a biocompatible *in vivo* ligation reaction based on a selective reaction between *D*-cysteine and **CBT** *in vivo*, producing a luciferin substrate for bioluminescence imaging of firefly luciferase in mice (Fig. 31B). This “split luciferin” ligation reaction was used

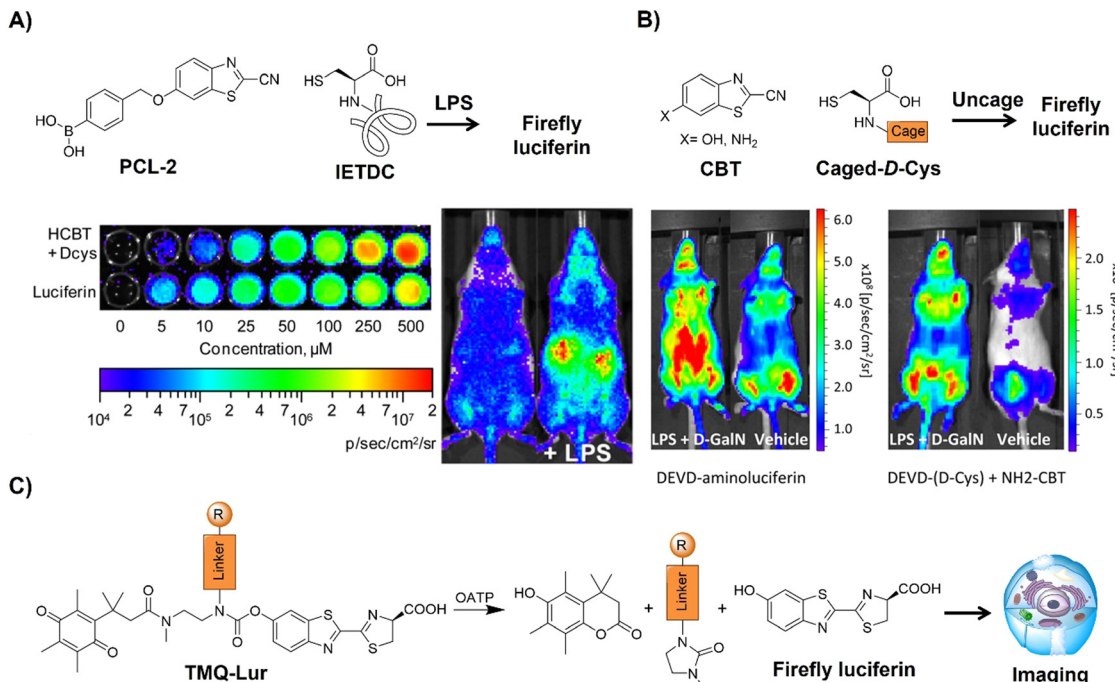


Fig. 31 Bioluminescent probes designed for imaging based on the 1,2-aminothiol-CBT click reaction. (A) Illustration of simultaneous detection of H_2O_2 and caspase 8 activity through release of **HCBT** and D -cysteine and *in situ* formation of firefly luciferin and representative image of PC3M-luc cells/mice. Adapted with permission from ref. 203. Copyright 2013, American Chemical Society. (B) Schematic illustration of the "split luciferin" ligation reaction and representative image of mice under different conditions. Adapted with permission from ref. 204. Copyright 2017, American Chemical Society. (C) Schematic illustration of the wash-free luminogenic probes for detecting **OATP** activity. Adapted with permission from ref. 205. Copyright 2016, American Chemical Society.

to investigate the click reactions of 6-amino/hydroxyl-CBT with D -cysteine and L -cysteine and revealed the high reactivity of CBTs towards D -cysteine. Utilizing bioluminescence imaging, visualizing the production of the luciferin substrates in live mice was achieved. Application of this reaction for real-time noninvasive imaging of apoptosis associated with caspase -3/-7 activity was also demonstrated.

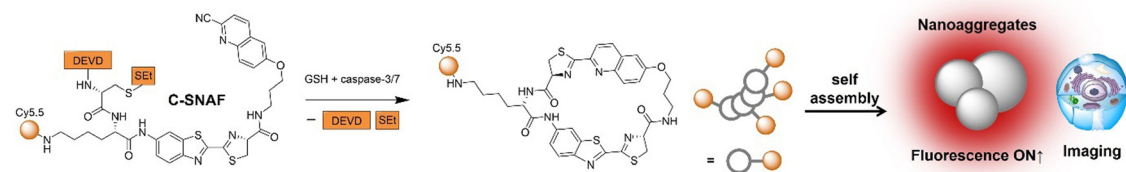
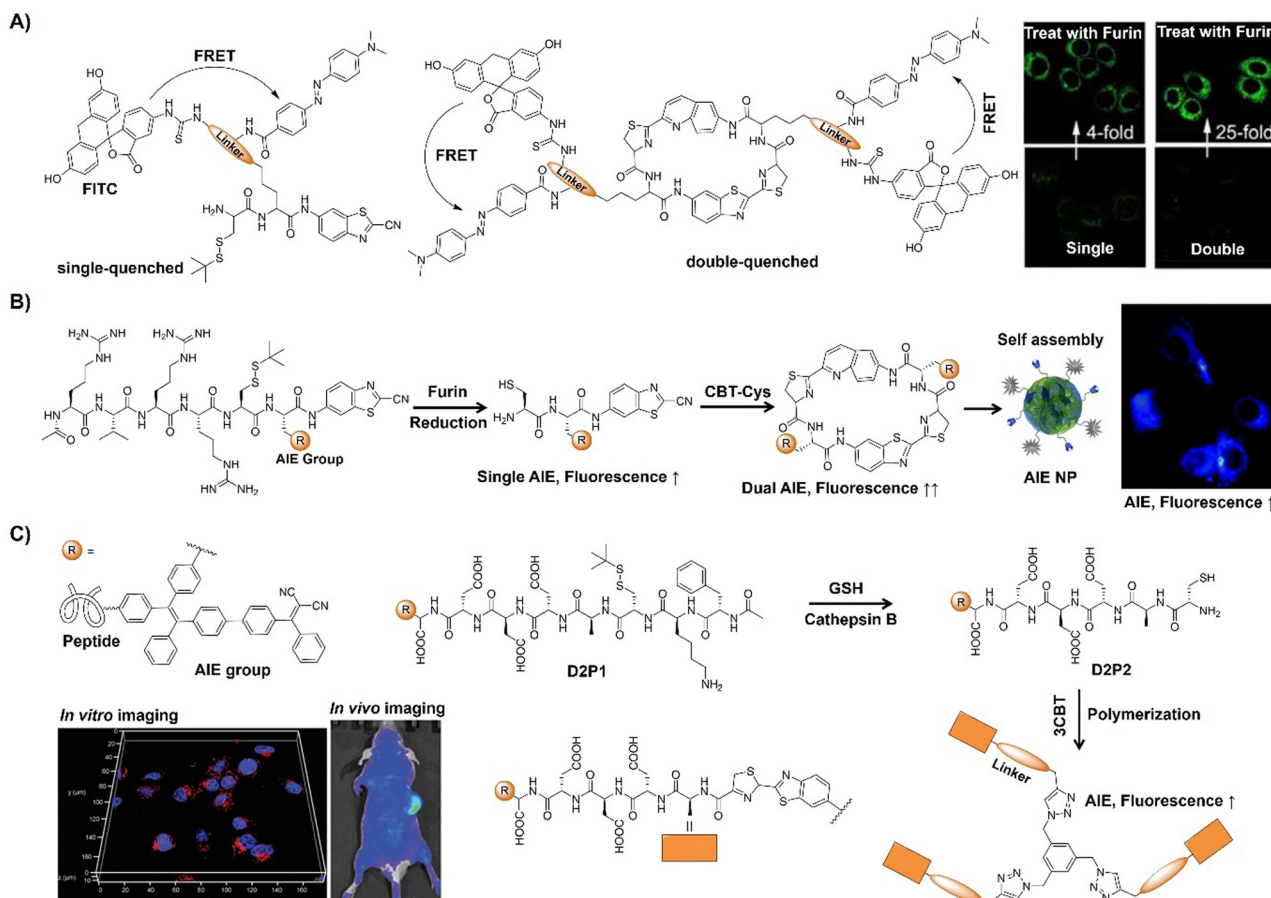
Through click reaction of CBT with D -cysteine, Cali and coworkers²⁰⁵ developed wash-free luminogenic probes for detecting the activity of organic anion-transporting polypeptides (OATP), pivotal in drug absorption, distribution, metabolism, and excretion (ADME) processes (Fig. 31C). This approach employed self-cleavage of trimethyl lock quinone-luciferin (TMQ-Lur) tethered to fluorescein as a recognized transporter substrate. Cellular uptake triggered quinone reduction and released luciferin *via* intramolecular lactonization, yielding detectable bioluminescence signals upon introduction of luciferase. The signal intensity corresponds to the activity of the transporter, enabling wash-free imaging, as demonstrated using HEK293 cells overexpressing OATP1B1*1a and OATP1B3. These results underscore the viability of measuring transporter activity *via* bioluminescence, with the potential for developing diverse transporter assays by simply altering the substrate.

Controlled self-assembly of small molecules within living systems offers extensive potential for applications in biology and medicine, such as in synthesizing supramolecules and nano/microstructures, both in solution and within living cells.

Bioorthogonal reactions based on CBTs and D -cysteine facilitate the controllable self-assembly of fluorescent small molecules to nanoparticles in the body. For example Rao's research²⁰⁰ employed an optimized first-order 1,2-aminothiol-CBT click reaction to control the self-assembly of fluorescent small molecules (Fig. 32). Imaging caspase-3/-7 activity in human tumor xenograft mouse models undergoing chemotherapy was demonstrated as one of the practical applications of this self-assembly system. Validation of the proposed mechanism was achieved using a series of control probes, and the self-assembled nanoaggregates were directly observed both *in vitro* and *in vivo* using super-resolution fluorescence microscopy. This bioorthogonal macrocyclization and nanoaggregation, triggered by caspase-3/-7 activity, facilitated effective monitoring of tumor therapy response *in vivo* and exhibited promise for imaging caspase-3/-7 activity in various diseases, including neurodegenerative diseases and organ-transplant rejection.

In addition to the probes for the analysis of caspase-3/7 activity, CBT- D -cysteine click condensation reactions have been used for the development of probes for detecting intracellular furin activity by the Liang group (Fig. 33A).²⁰⁶ The probe was developed using a dual quenching approach by combining a click condensation reaction to form the single quenched probe DABCYL-FITC-CBT. This strategy enabled enhanced sensitivity in detecting intracellular furin activity. Under physiological conditions, DABCYL-FITC-CBT underwent



Fig. 32 Schematic illustration of the mechanism of probe **C-SNAF** for caspase-3/7 activity analysis and imaging.Fig. 33 Fluorescent probes designed for bioimaging based on the 1,2-aminothiol-CBT click reaction. (A) Schematic illustration and confocal fluorescence and overlay images of MDA-MB-468 cells of the single/dual quenched nanoparticles. Adapted with permission from ref. 206. Copyright 2018, American Chemical Society. (B) Schematic illustration of a furin-controlled dual AIEgen for enhanced fluorescence sensing of furin activity. Adapted with permission from ref. 207. Copyright 2017, The Royal Society of Chemistry. (C) Schematic illustration of the formation of AIEgen-based nanostructures by enzyme-mediated intracellular reduction and condensation of **D2P1** and **3CBT** and their applications in *in vitro* and *in vivo* imaging. Adapted with permission from ref. 208. Copyright 2021, Wiley-VCH GmbH.

reduction-controlled condensation to form DABCYL-FITC-CBT-NPs, with the fluorescence intensity reducing by 1/2.8 of its original intensity. Upon furin cleavage, the dual-quenched DABCYL-FITC-CBT-NPs exhibited an 11-fold fluorescence “turn-on” compared to the single-quenched control probe FITC-DABCYL. Live cell imaging exhibited a 6.3-fold fluorescence “turn-on” for detecting intracellular furin activity. This versatile dual-quenching strategy is interesting and has the potential for enhanced sensitivity in detecting various

intracellular enzyme activities. Subsequently, Liang and coworkers²⁰⁷ developed a probe using AIEgens and a furin-controlled CBT click reaction for furin activity assays in live cells (Fig. 33B). Upon reduction and furin hydrolysis, dimer formation occurred, self-assembling into nanoparticles and intensifying the fluorescence through enhanced fluorogen aggregation. Compared to single AIE molecules lacking CBT groups, the fluorescence signals were boosted 1.7 and 3.4-fold *in vitro* and in living cells, respectively.

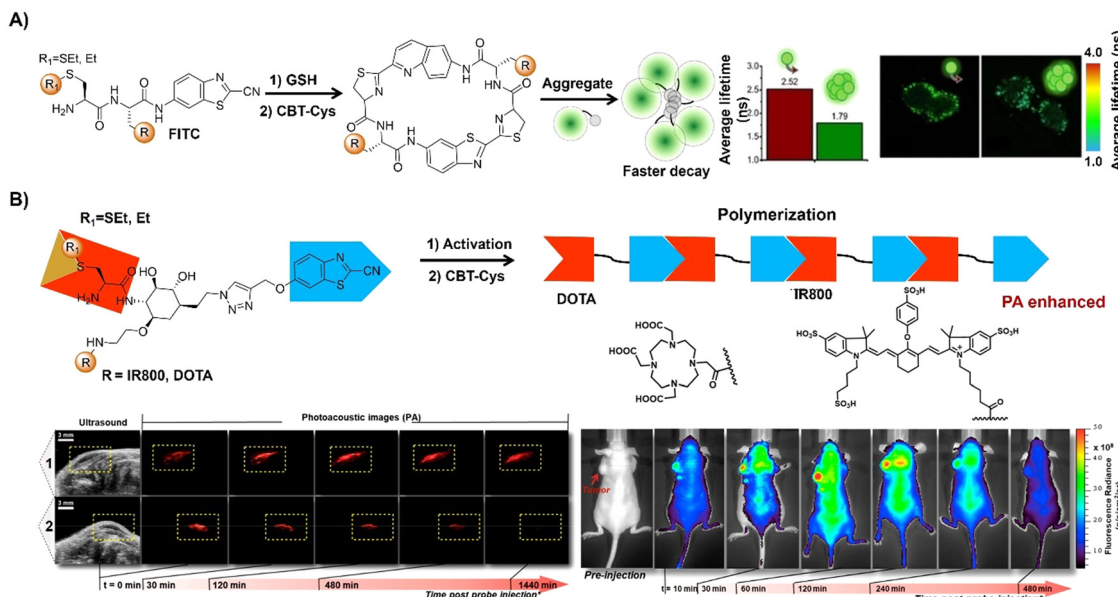


Fig. 34 Bioluminescent probes developed by exploring the 1,2-aminothiol-CBT click reaction. (A) Schematic illustration and cell imaging of a reduction-sensitive fluorescent probe obtained via **CBT-Cys** reaction. Adapted with permission from ref. 209. Copyright 2020, Wiley-VCH GmbH. (B) Schematic illustration of an *in situ* polymerization activated probe via **CBT-Cys** reaction and ultrasound and photoacoustic images of a subcutaneously injected probe in living mice and longitudinal fluorescence imaging of tumor-bearing mice. Adapted with permission from ref. 210. Copyright 2020, American Chemical Society.

Using a similar TPE-based AIEgen, Liu's group²⁰⁸ recently reported an enzyme-mediated AIEgen intracellular polymerization strategy to enhance drug localization within tumors (Fig. 33C). This approach used an AIEgen-peptide conjugate (**D2P1**) and cyanobenzothiazole-cysteine (**3CBT**) to form nanoaggregates in tumor cells *via* a rapid condensation reaction. Tumor-specific cathepsin protease cleavage triggered polymerization, promoting **D2P1** accumulation in tumors for fluorescence activation. This process disrupted cellular motility and inhibited cancer cell proliferation. Moreover, **D2P1** is able to generate ROS, boosting tumor PDT efficacy *in vivo*.

In addition to the analysis based on fluorescence intensity, fluorophore aggregation offers another approach for bioanalysis and imaging, *i.e.*, fluorescence lifetime imaging. For instance, Cui and colleagues²⁰⁹ introduced a reduction-sensitive fluorescent probe, exhibiting altered emission lifetime under reducing conditions as a result of the fluorophore aggregation of the CBT scaffold (Fig. 34A). Remarkable changes in lifetime occur in FITC emission when the compound interacts with glutathione (**GSH**), indicating the necessity of both a reduction-sensitive disulfide bond and reducing agent for lifetime shift. During FLIM imaging a lifetime decrease was detected after activation by GSH, suggesting the potential of the probe for lifetime imaging of the fluorophore's self-aggregation in cells. In 2020, the Rao group²¹⁰ introduced *in situ* polymerization of activatable bioorthogonal small molecules in response to changes in the reducing environment *in vivo* (Fig. 34B). A carbohydrate linker and **CBT-Cys** condensation reaction-based small molecule scaffold were developed for rapid condensation upon physicochemical changes, forming

pseudopolysaccharides. Evaluation of the fluorescence and photoacoustic properties before and after transformation confirmed *in situ* polymerization post-local and systemic administration in living mice, highlighting the potential of this probe for dual-modality optical imaging.

Apart from the probes developed based on **CBT-d**-cysteine initialized assembly, disassembly of self-quenched nanoparticles has also been adopted for the development of probes by Liang's group. For instance, in 2015, Liang and coworkers²¹¹ developed an enzyme-driven disassembly of self-quenched NIR fluorescent nanoparticles, enabling targeted detection of enzyme activity *in vitro* and *in vivo* (Fig. 35A). Through strategic design, the NIR probe's fluorescence was initially suppressed *via* reduction-controlled condensation and nanoparticle assembly, only to be activated upon NP disassembly by furin. Leveraging this enzymatic mechanism, precise NIR detection of furin was achieved *in vitro* and for live cell imaging. Additionally, the disassembly approach-based probe facilitated discriminative NIR imaging of MDA-MB-468 tumors in nude mice, suggesting the potential for tumor-targeted imaging in clinical applications. Expanding on this research, the same group then developed a NIR probe, Cys(StBu)-Lys(Biotin)-Lys-Lys(Cy5.5)-**CBT**, to generate self-quenched nanoparticles (**1-NPs**) for *in vitro* and *in vivo* tumor-targeted imaging (Fig. 35B).²¹² Biotinylated **1-NPs** exhibited active uptake by biotin receptor-overexpressing tumor cells *via* receptor-mediated endocytosis. Intracellular proteolytic cleavage dismantled **1-NPs**, yielding the fluorescent "turn-on" molecular probe Lys(Cy5.5)-Luciferin-Lys(Biotin)-Lys-OH (**1-d**-cleaved). This nanoprobe with NIR fluorescence facilitated tumor-targeted imaging in a nude mouse model.



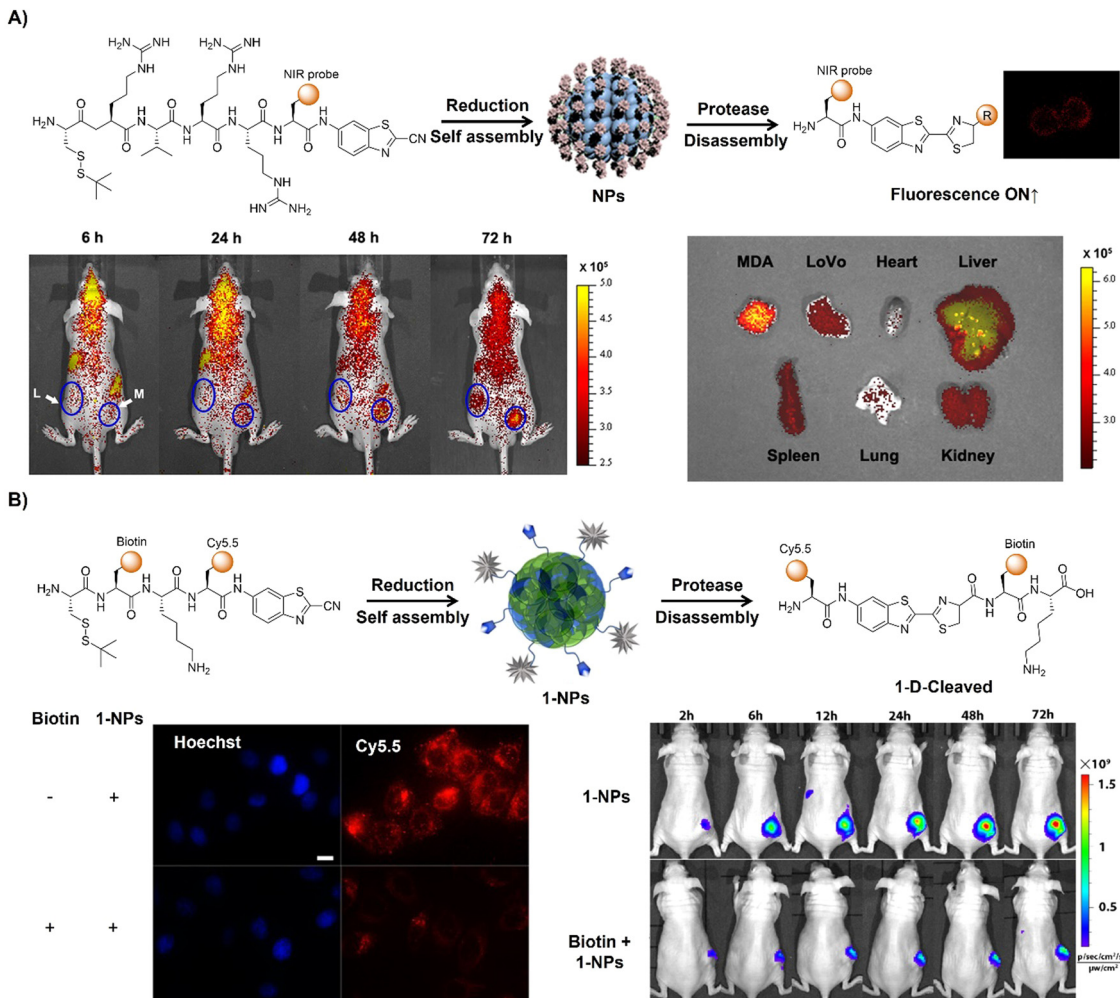


Fig. 35 Fluorescent probes developed for *in vitro* and *in vivo* imaging based on the 1,2-aminothiol-CBT click reaction. (A) Illustration of the enzyme-driven disassembly of self-quenched NIR fluorescent nanoparticles; *in vivo* active furin-overexpression-tumor-targeted imaging and *ex vivo* fluorescence images of different organs from tumor-bearing nude mice. Adapted with permission from ref. 211. Copyright 2015, American Chemical Society. (B) Schematic illustration of reduction-controlled self-assembly of **1-NPs** to turn the NIR signal "Off" and subsequent protease-controlled disassembly of **1-NPs** to turn the NIR signal "On" with fluorescence and overlay images of biotin receptor-positive HeLa cells and time-course fluorescence imaging of nude mice xenografted with HeLa tumor cells. Adapted with permission from ref. 212. Copyright 2017, American Chemical Society.



Scheme 5 Proposed mechanism of the SPSAC reaction.

In brief, we have summarized the progress made in using CBT-Cys click reactions for *in situ* bioimaging in this section. The CBT-Cys click reaction relies on the condensation between CBT and *d*-Cys, as observed in fireflies. The mechanism has been proposed and verified experimentally, showcasing numerous advantages over conventional click reactions. This click reaction has exceptional biocompatibility because the reaction has natural origins. This reaction could be easily modulated by changes in pH, reduction, or enzymes under physiological

conditions. More importantly, its second-order rate constant is high at $9.19 \text{ M}^{-1} \text{ s}^{-1}$, surpassing that of CuAAC by two orders of magnitude. Furthermore, the resulting amphiphilic oligomeric products tend to self-assemble into nanostructures, a unique trait not seen in other click reactions. The versatility of the CBT-Cys click reaction facilitates a wide range of biomedical applications through modification of the chemical structures of the reactants. Under the influence of pH, reduction, or enzymes, the CBT-Cys click condensation reaction has successfully been applied in various fields, including protein labeling, molecular imaging (covering optical, magnetic resonance, nuclear, and photoacoustic imaging) of tumors, preparation of oligomeric nanostructures, and cancer therapy. The development of "smart" small molecular precursors enables precise control over intracellular spatiotemporal assembly. It is expected that this reaction will facilitate the development of novel dendrimer-based probes to enhance imaging contrast

(e.g., MRI and fluorescence imaging) by covalently attaching the probes at the periphery. Moreover, as molecular imaging advances towards multimodality, integrating multiple imaging modes into a single platform using the CBT-Cys click reaction holds promise for more accurate disease diagnosis through multimodal molecular imaging.

2.5. Strain promoted sydnone alkyne cycloaddition

Sydnones, a family of mesoionic compounds, have recently emerged as valuable components for click chemistry and bioorthogonal reactions. Reactions of sydnones with cyclooctynes, known as strain promoted sydnone–alkyne cycloaddition (SPSAC, Scheme 5),^{213,214} enable the synthesis of pyrazoles with intriguing properties. Sydnones offer the advantage of functionalizing mesoionic scaffolds at multiple positions, allowing for fine-tuning of their reactivity. This section will focus on the latest advancements in *in situ* activated bioimaging utilizing the sydnone–cyclooctyne reaction over the past few years, aiming to provide comprehensive and profound insights for its further implementation.

In 2014, Taran and colleagues synthesized a series of sydnone derivatives and investigated their reactivities towards **BCN** through a Cu-free cycloaddition reaction.²¹³ It was found that 4-chloro-sydnones could react with **BCN** more than 10 times faster than azide-based click reactions. The high reactivity of chlorosydnones was then applied to fluorescence labelling of

proteins using rhodamine-linked **BCN** (**BCN-TAMRA**). In 2016, the same group²¹⁴ reported the synthesis and dynamic reactivity of 4-fluorosydnones, an exceptional category of mesoionic dipoles exhibiting remarkable responsiveness towards both copper-catalyzed and strain-promoted cycloaddition reactions with alkynes (Fig. 36A). These mesoionic compounds were synthesized through electrophilic fluorination of σ -sydnone $\text{Pd}(\text{n})$ precursors in the presence of Selectfluor. Their interactions with both terminal and cyclic alkynes involve rapid and highly selective transformations, yielding 5-fluoro-1,4-pyrazoles with unprecedented bimolecular rate constants exceeding $10^4 \text{ M}^{-1} \text{ s}^{-1}$. Detailed kinetic investigations were conducted to elucidate the underlying reaction mechanism, further underlining the significance of 4-fluorosydnones, exemplified by their successful radiolabeling with $[^{18}\text{F}]$ Selectfluor. On the basis of these studies, the same group then synthesized a range of styryl-pyridinium-based sydnones tailored for wash-free protein labeling and in-cell imaging (Fig. 36B).²¹⁵ These “turn-on” sydnone probes exhibited high reactivity towards cyclooctynes, facilitating clean reactions with rate constants ranging from 10 to 100 $\text{M}^{-1} \text{ s}^{-1}$. Using these probes, highly effective protein labeling and cellular imaging applications were successfully demonstrated.

The Lo group²¹⁶ introduced a series of ruthenium(II) poly-pyridine complexes featuring a nitronate group as phosphogenetic bioorthogonal probes (Fig. 36C). Initially weakly emissive

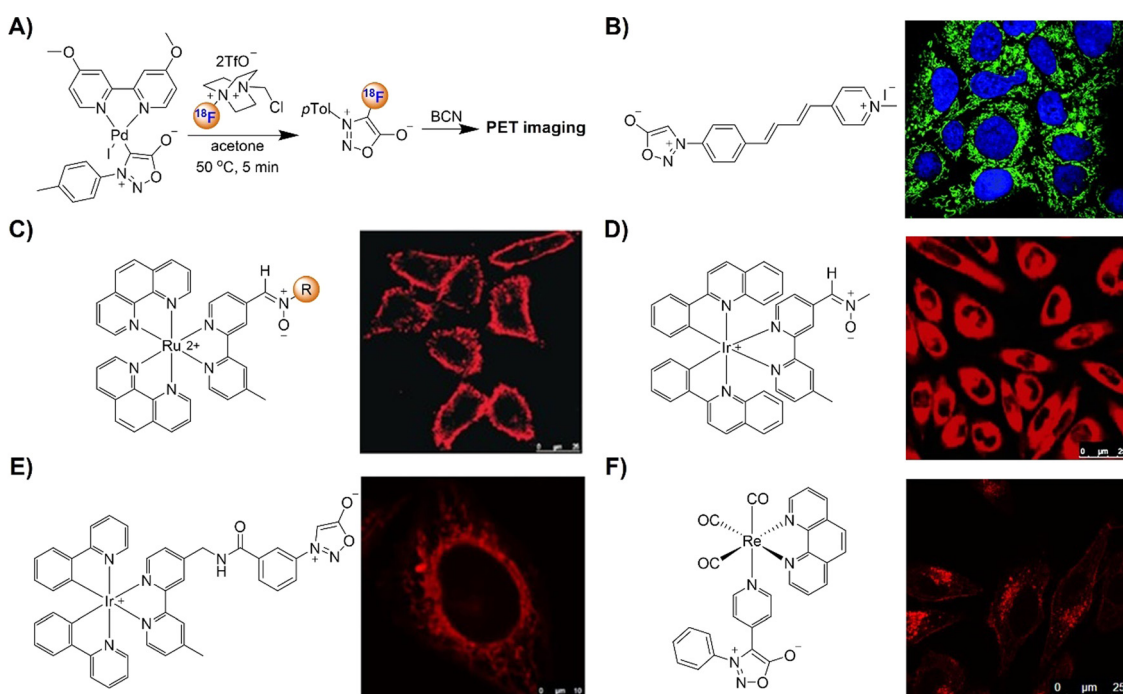


Fig. 36 Metal complex-based bioorthogonal probes for cell imaging based on the strain promoted sydnone alkyne cycloaddition. (A) Preparation and use of $[^{18}\text{F}]$ sydnone; (B) the structure of 6-sydnone-profluorophores and confocal cell imaging, adapted with permission from ref. 215. Copyright 2019, The Royal Society of Chemistry; (C) the structure of the Ru nitronate complex and LSCM images of the HeLa cells, adapted with permission from ref. 216. Copyright 2016, Wiley-VCH GmbH; (D) the structure of the Ir nitronate complex and LSCM images of the CHO-K1 cells, adapted with permission from ref. 217. Copyright 2016, Wiley-VCH GmbH; (E) the structure of the Ir sydnone complex and LSCM images of the HeLa cells, adapted with permission from ref. 218. Copyright 2018, Wiley-VCH GmbH; and (F) the structure of the Re polypyridine sydnone complex and LSCM images of the HeLa cells, adapted with permission from ref. 219. Copyright 2020, Wiley-VCH GmbH.

due to rapid C=N isomerization of the nitrone moiety, these complexes exhibited significant emission enhancement upon strain-promoted alkyne-nitronate cycloaddition (**SPANC**) with **BCN**-modified substrates. By incorporating a dicationic ruthenium(II) polypyridine unit at the α -C-position and a phenyl ring at the N-position of the nitronate, reaction kinetics were remarkably accelerated, up to 278-fold higher than other acyclic nitronate-**BCN** reactions. Remarkably, these complexes facilitated specific cell membrane/cytosol staining upon labeling with an exogenous substrate, **BCN**-modified decane, in live cells. Using a similar **SPANC** approach, the same research group designed three iridium(III) complexes incorporating a nitronate moiety as phosphorogenic bioorthogonal probes (Fig. 36D).²¹⁷ Similar to the luminescence quenching of the ruthenium(II) polypyridine complex,²¹⁶ isomerization of the C=N group quenched the emissions of these iridium(III) complex-based probes, while emission enhancement was obtained upon cycloaddition with strained cyclooctynes. Likewise, the cationic iridium(III) center promoted the reaction kinetics of nitronate with **BCN**. These nitronate complexes also served as phosphorogenic bioorthogonal labels and imaging agents for cyclooctyne-modified proteins, offering promising prospects for advancing phosphorogenic bioorthogonal probes and imaging reagents. Subsequently, the same research group employed the transformation of the highly polar mesoionic sydnone ring into less polar pyrazoles to create luminescent probes for the **SPSAC** reaction in bioorthogonal labeling and live cell imaging (Fig. 36E).²¹⁸ The developed iridium(III) sydnone complexes readily reacted with strained alkyne derivatives, resulting in significant emission enhancement (up to 24.9 fold) and extended lifetime. These complex-based probes were successfully utilized to label alkyne-modified proteins and ceramide molecules within live cells.

Following the success of ruthenium(II) and iridium(III) complex-based **SPSAC** bioorthogonal labelling, Lo's group then synthesized phosphorogenic rhodium(I) polypyridine sydnone complexes to expand the bioorthogonal toolkit (Fig. 36F).²¹⁹ Upon **SPSAC** with bicyclo[6.1.0]non-4-yn-9-ylmethanol (**BCN-OH**), these complexes displayed 8.8 to 17.3 fold enhancement in emission. Upon conjugation with **BCN**-modified bovine serum albumin (**BCN-BSA**), the emission enhancement of the rhodium(I) complex increased dramatically, reaching up to 38.9 times with lifetimes extended from 1.80 to 4.71 μ s. Due to their

low cytotoxicity and remarkable emission enhancement capabilities, these complexes were applied for bioimaging of lysosomes in live HeLa cells.

In 2018, the Frisicourt group²²⁰ reported the development of sydnone-modified coumarins as fluorogenic click reagents (Fig. 37A). The sydnone functionality quenched the fluorescence of the methoxycoumarin dye due to a nonemissive TICT state, with a low quantum yield of 0.004. Upon rapid strain-promoted cycloaddition with cyclooctyne **BCN**, these reagents produced highly fluorescent pyrazole cycloadducts with emission characteristics similar to 7-methoxycoumarin. Fluorescence quantum yields increased significantly (up to 0.66 and up to 132-fold enhancement), allowing for highly specific wash-free protein labeling.

The Liang group introduced a fluorogenic probe, Naph-Syd, featuring an off-on fluorescence response mechanism for cell imaging. The probe was developed by incorporating a sydnone functionality into a 1,8-naphthlimide fluorophore (Fig. 37B).²²¹ Sydnone played pivotal roles in (i) quenching the fluorescence of 1,8-naphthlimide through a PeT mechanism and (ii) click reaction with **BCN** derivatives. Upon rapid bioorthogonal cycloadditions with strained alkynes, pyrazole products were formed, leading to a remarkable 300-fold enhancement in 1,8-naphthlimide fluorescence, enabling fluorescence imaging in live cells under metal-free, wash-free conditions. Moreover, the probe exhibited simultaneous turn-on labeling of multiple targets by combining fluorogenic **Tz** cycloadditions with strained alkenes.

In another study of **SPANC** reaction-based fluorescence cell labelling, Tian and coworkers²²² developed a bioorthogonal turn-on probe utilizing nitronate-modified 1,8-naphthalimide (Fig. 37C). The rearrangement products resulting from nitronate-alkyne cycloadducts exhibited 252-fold fluorescence enhancement, along with a substantial Stokes shift and high Φ_F , up to 0.35. The high specificity of this **SPANC** reaction enabled effective protein labeling *in vitro* and mitochondrial labeling in living cells with a high signal-to-noise ratio under catalyst-free, wash-free conditions.

In addition to the labelling of proteins and live cells, **SPSAC** based bioorthogonal tags for labelling and tracking nucleosides and oligonucleotides in live cells have also been investigated by Wagenknecht and colleagues (Fig. 38A).²²³ Investigation of **SPSAC** kinetics with model substrates revealed rapid reactions

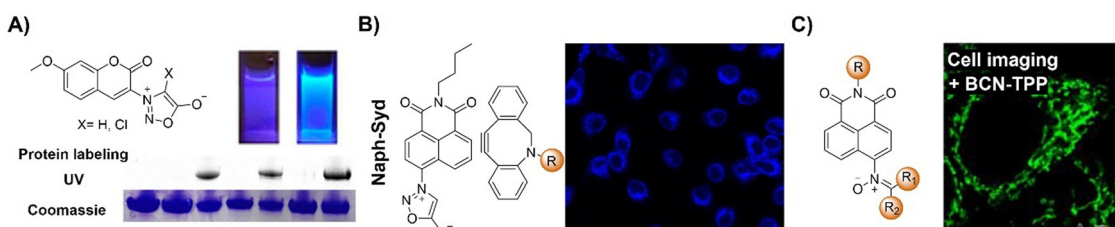


Fig. 37 Fluorescent probes developed for bioimaging based on the strain promoted sydnone alkyne cycloaddition. (A) The structures and protein labeling of sydnone-modified coumarins, adapted with permission from ref. 220. Copyright 2018, American Chemical Society; (B) the structure of Naph-Syd and fluorescence imaging of ManDIBAC, adapted with permission from ref. 221. Copyright 2019, Elsevier Ltd; and (C) the structure of the nitronate-modified 1,8-naphthalimide and mitochondrial labeling in live cells, adapted with permission from ref. 222. Copyright 2022, Wiley-VCH GmbH.

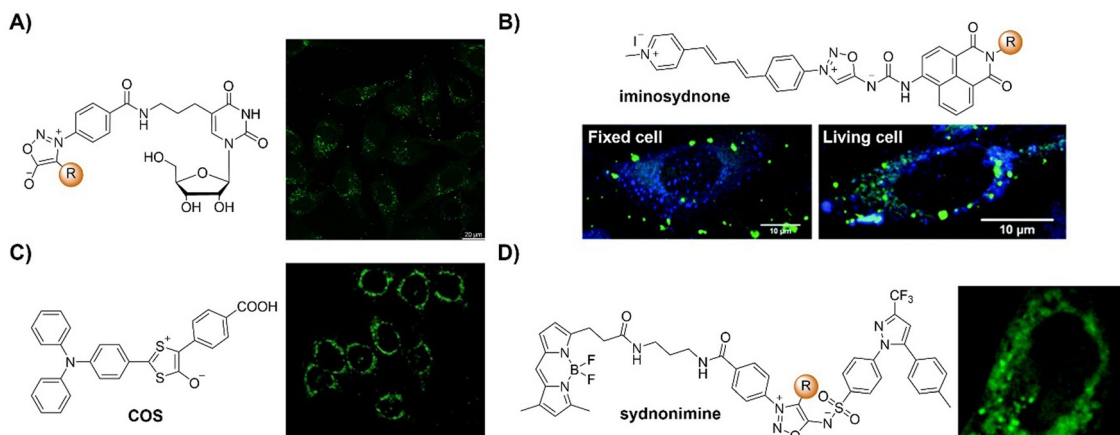


Fig. 38 Strain promoted sydnone alkyne cycloaddition-based bioorthogonal probes for fluorescence imaging. (A) The structures of the sydnone-modified 2'-deoxyuridines and confocal imaging of the HeLa cells, adapted with permission from ref. 223. Copyright 2021, Wiley-VCH GmbH; (B) the structure of the iminosydnone probe and comparison of the distribution of the two released products in fixed vs. living cells after reaction, adapted with permission from ref. 224. Copyright 2020, The Royal Society of Chemistry; (C) the structure of the **COS** probe and fluorescence images of reactions, adapted with permission from ref. 225. Copyright 2021, Wiley-VCH GmbH; and (D) the structures of the sydnonimines and fluorescence images of reactions in HeLa cells, adapted with permission from ref. 226. Copyright 2023, Elsevier Ltd.

with cyclooctyne probes (up to $k = 0.59 \text{ M}^{-1} \text{ s}^{-1}$), yielding a 70-fold fluorescence enhancement. Sydnones were integrated into both 2'-deoxyuridines at position 5 and 7-deaza-2'-deoxyadenosines at position 7, and the modified nucleosides were then synthetically incorporated into single-stranded DNAs, allowing for postsynthetic labeling with cyclooctyne probes both in solution and in live HeLa cells.

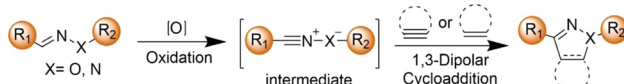
Probes based on fluorescence turn on click-and-release reactions enable the multicolor labelling of samples under biological conditions. For example, the Taran group²²⁴ synthesized a series of iminosydnone probes that can efficiently cleave and unquench two distinct fluorophores upon bioorthogonal reaction with cycloalkynes, resulting in remarkable fluorescence enhancements (Fig. 38B). These versatile iminosydnones served as fluorogenic cleavable linkers, enabling the cellular tracking of the two products generated from the click and release reaction. Examination of cell distributions revealed distinct cytoplasmic localization of the two released products, confirming their separation and movement within cells prior to precipitation. The compatibility of this reaction was validated in fluorescence labelling of fixed or living cells through monitoring the release of the double turn-on fluorescence signals. In another research conducted by the same group, release of isocyanates in live cells was also achieved using bioorthogonal click-and-release reactions, where the reactivity constants reached up to $1000 \text{ M}^{-1} \text{ s}^{-1}$.²²⁷

Inspired by the strain-promoted cycloaddition reaction involving 1,3-dithiolium-4-olates (**DTOs**) and cyclooctyne or benzyne derivatives, Liang and colleagues devised a carbonyl sulfide (**COS**)-based system for controlled release of H_2S (Fig. 38C).²²⁵ This system was activated by a bioorthogonal click-and-release reaction, enabling precise control over H_2S release in both PBS and live cells. Incorporating a diphenylamino substituent onto one phenyl ring of **DTO** facilitated simultaneous activation of a new thiophene fluorophore,

enabling spatiotemporal monitoring of intracellular H_2S release using green fluorescence signals. Beyond its bioorthogonal and visualization capabilities, this versatile platform allowed for targeted delivery of H_2S to mitochondria. The Liang group²²⁶ also explored the mitochondrial targeting capabilities of sydnones and sydnonimines (Fig. 38D). Experimental findings confirmed that both sydnones and sydnonimines exhibited significant mitochondrial distribution, while incorporation of a phenyl group into the C_4 position of sydnones markedly diminished their affinity for mitochondria. Leveraging the mitochondrial targeting properties and click-and-release reaction of sydnonimine, the anticancer efficacy of celecoxib delivered into mitochondria was evaluated in HeLa and HepG2 cells.

In summary, bioorthogonal chemistry based on mesoionic sydnones has experienced significant advances since their discovery in 1935. Initially acknowledged for their biological significance, their reactivity was only fully understood following Huisgen's work in 1962. However, harsh reaction conditions and inadequate control over regioselectivity impeded further exploration of their potential for cycloaddition reactions. Nonetheless, recent studies have achieved notable breakthroughs, generating increased interest within the chemical community. Sydnones, especially when combined with strained cycloalkynes, have emerged as potent tools for bioorthogonal chemistry, facilitating *in situ* bioimaging applications. Despite being larger in size compared to azides, sydnones offer adjustable reactivity through strategic substitution, enabling the development of ultra-fast reactions crucial for *in vivo* studies. For instance, Taran's research²¹³ conducted using 4-chlorosydnones resulted in a more than tenfold increase in reactivity towards **BCN** compared to conventional azides, proving invaluable for fluorescence protein labeling, particularly for proteins present in low abundance. Furthermore, advancements such as iminosydnones²²⁷ and highly strained alkynes²¹⁴ hold promise





Scheme 6 Proposed mechanism of nitrile imine–alkyne cycloaddition.

for further acceleration, with reaction rates exceeding $12\,000\text{ M}^{-1}\text{ s}^{-1}$, thereby enhancing their suitability for bioimaging. Despite notable advancements, challenges persist, notably, the lack of regiocontrol in the presence of unsymmetrical arynes. Addressing these challenges will undoubtedly propel future progress in this distinctive field of mesoionics.

2.6. Nitrile imine–alkyne cycloaddition

The 1,3-dipolar cycloaddition between nitrile imines and alkenes/alkynes is known to be a rapid bioconjugation reaction for labeling proteins or nucleic acids (Scheme 6).²²⁸ Since nitrile imines are inherently unstable, these dipoles are typically generated *in situ* through different methods like tetrazole photolysis, hydrazone oxidation, or chlorohydrazone hydrolysis. The remarkable reactivity and robustness of nitrile imines have recently sparked interest in their use for designing fluorogenic click probes. In this section, we will evaluate the applications of nitrile imine–alkene/alkyne 1,3-dipolar cycloaddition for *in situ* activated bioimaging.

The Tian group²²⁸ introduced 4-oxime-1,8-naphthalimide (NP-Oxm) as a bioorthogonal turn-on probe for the fluorogenic labelling of proteins (Fig. 39A). Using this probe, highly reactive naphthalimide-nitrile oxide was produced *in situ*, which served as the bioorthogonal handle for 1,3-dipolar cycloaddition with alkenes or alkynes. Upon cycloaddition with alkenes, various 4-isoxazoline-naphthalimides (NP-Dmf, NP-Str, and NP-Nob) exhibited a 5- to 13-fold increase in fluorescence. Notably, the reaction products, 4-isoxazolyl-naphthalimides (NP-BCN and NP-DIBO), resulting from strain-promoted alkyne-nitrile oxide

cycloaddition, demonstrated photoactivatable fluorogenic properties *via* isoxazole–oxazole photoisomerization, boasting over 250-fold fluorescence enhancement with a high fluorescence quantum yield ($\Phi_F > 0.4$). Furthermore, NP-Oxm facilitated highly specific protein labeling in a tunable emission manner. This photoactivatable bioorthogonal turn-on strategy provided a useful toolkit for chemical biology investigations, enabling precise visualization of biomolecules in living cells.

In 2020, the same research group devised and synthesized four 4-hydrazone-1,8-naphthalimides as fluorogenic click probes (Fig. 39B).²²⁹ These compounds produced nitrile imines *in situ* that can react with BCN *via* a highly fluorescent turn-on strain-promoted 1,3-dipolar cycloaddition, yielding 4-pyrazol-1,8-naphthalimides. These derivatives showed remarkable characteristics including large Stokes shift (up to 152 nm), high Φ_F (up to 0.85), and substantial fluorescence enhancement (up to 196.9-fold). These hydrazone probes were effectively employed for the fluorogenic labeling of alkyne modified BSA proteins, demonstrating their potential as valuable tools for protein visualization and detection.

Following this success, the same research group²³⁰ then synthesized a series of nitroso-modified naphthylene fluorophores for bioorthogonal nitroso–diene Diels–Alder reactions (Fig. 39C). Upon photoactivation, these probes exhibited remarkable properties, *i.e.*, formation of bridged cycloadducts between 4-nitroso-1,8-naphthlimide or 5-nitroso-naphthalene-1-sulfonamide and cyclic dienes led to aziridine-containing rearrangement products. These products displayed remarkable fluorescence enhancement, with an increase of fluorescence intensity up to 3604-fold, accompanied by large Stokes shifts of up to 188 nm, a high Φ_F of up to 0.92, superior photostability, and distinct solvatochromic effects. The cycloadducts formed between nitroso compounds and multiple methyl-substituted cyclopentadiene *via* [2+2] cycloaddition underwent spontaneous rearrangement to produce fluorescent hydroxylamine

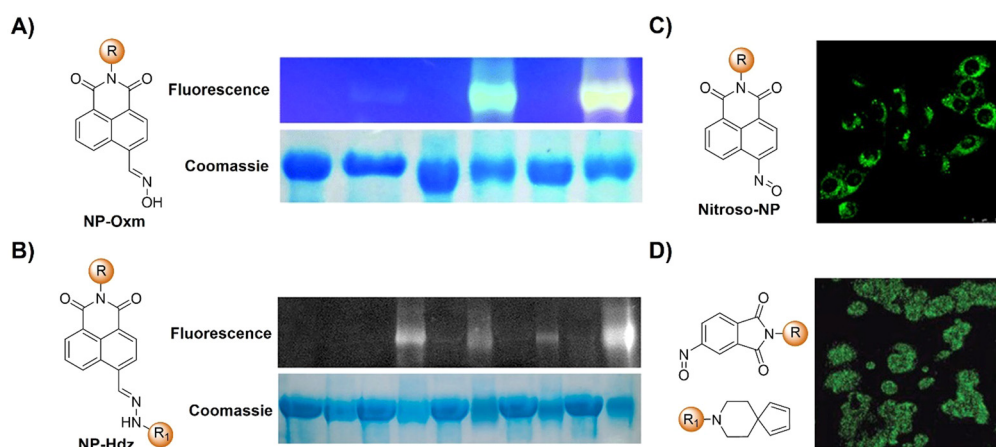


Fig. 39 Fluorescent probes based on the nitrile imine–alkyne cycloaddition. (A) The structure and protein labeling of NP-Oxm, adapted with permission from ref. 228. Copyright 2019, The Royal Society of Chemistry; (B) the structure and protein labeling of NP-Hdz, adapted with permission from ref. 229. Copyright 2020, Wiley-VCH GmbH; (C) the structure of Nitroso-NP and fluorescence images of reactions, adapted with permission from ref. 230. Copyright 2021, American Chemical Society; and (D) the structure of the nitroso-modified naphthylene and fluorescence images of reactions, adapted with permission from ref. 231. Copyright 2019, Wiley-VCH GmbH.



products. These nitroso–diene probe pairs were effectively utilized for selective fluorogenic labeling of diene-modified proteins and for wash-free organelle-specific mammalian live-cell imaging in real time, in a spatiotemporally controlled manner.

In another study, Jiang and coworkers²³¹ harnessed the fluorogenic reactivity of nitroso-based bioorthogonal Diels–Alder reactions to develop the fluorogenic nitroso probe (Fig. 39D). This reaction facilitated the generation of a twisted “donor–acceptor–donor” skeleton, crucial for effective separation of HOMO/LUMO electron densities, thereby enabling the development of probes with thermally activated delayed fluorescence (TADF) properties. This unique feature enabled selective activation of Point of Interest (POI) within specific cell compartments while leaving other labeled molecules intact. The precise spatial and temporal control afforded by this fluorogenic labeling system holds promise for monitoring POI dynamics in live cells across various applications. Expanding beyond model protocols, they then devised an efficient “click-like” fluorogenic nitroso–diene pair probe for general, controllable protein labeling in live cells. Proof-of-concept studies underscored the optimized probe’s potency for fluorogenic protein labeling with minimal background signals under UV irradiation. The utility of this bioorthogonal reaction in living mice was assessed for noninvasive *in vivo* fluorescence imaging.

2.7. Photoclick reaction

Photoclick reactions^{232–234} synergize the benefits of photochemistry²³⁵ and classical click chemistry.^{34–37} Unlike standard click reactions occurring between molecules in their ground state, in photoclick reactions, the partners remain chemically inert until exposure to specific light wavelengths. This illumination activates one of the partners, triggering the formation of new covalent bonds. This property enables the synthesis of diverse organic structures under mild, metal-free conditions, with precise control. Moreover, light irradiation offers non-invasive characteristics, affording superior spatial resolution and temporal control over reactive intermediate generation. These attributes are particularly advantageous for bioorthogonal transformations, making photoclick chemistry an integral part of the toolbox for surface functionalization,²³⁶ polymer conjugation,^{237,238} photocross-linking,²³⁹ and bioorthogonal labeling.^{92,240,241}

Over the past few decades, researchers have explored an expanding array of photoclick reactions. These encompass diverse reactions such as the photoinduced cycloaddition of tetrazoles and alkenes (NITEC),^{242–244} UV-light initiated thiolene and thiol–yne reactions,^{127,245,246} light-triggered azide–alkyne cycloadditions,^{247,248} diarylsydnone–alkene photoclick reaction (photo-DASAC),²⁴⁹ photo-induced azirine–alkene cycloaddition,²⁵⁰ light-triggered oxime ligation reactions,²⁵¹ and light-mediated coupling of acylsilanes with indoles,²⁵² along with visible-light-induced 9,10-phenanthrenequinone–electron rich alkene (PQ-ERA) photocycloadditions.^{253–255} Notably, certain photoclick reactions yield products with excellent

fluorescence properties, exemplified by NITEC, photo-DASAC, and PQ-ERA. Leveraging the fluorescence responses, these reactions have been extensively used in precise labeling and imaging of biological systems. In this section, we aim to evaluate the evolution of photoclick reactions specifically for *in situ* activated bioimaging.

2.7.1. Light-triggered nitrile imine-mediated tetrazole–ene cycloaddition (NITEC). Initially, highly reactive nitrile imines were generated through the thermal decomposition ($>150\text{ }^{\circ}\text{C}$) of tetrazoles, liberating N_2 in the process.²⁵⁶ This decomposition can also be achieved *via* light exposure (refer to Fig. 40A), utilizing a wavelength that corresponds to the attached chromophore.²⁵⁷ The resultant photochemically induced nitrile imine undergoes a 1,3-dipolar cycloaddition with alkenes, yielding a conjugated five-membered pyrazoline ring as a photoclick product. Importantly, this process, known as NITEC, yields pyrazolines that typically exhibit fluorescence, thus facilitating real-time monitoring of the reaction. Various alkenes can partake in this reaction, particularly if they are conjugated or attached to electron-withdrawing substituents, exhibiting enhanced dipolarophilicity. A notable advantage of nitrile imine formation is the exclusive production of inert N_2 as a byproduct. The NITEC reaction exhibits good compatibility with a range of solvents, including aqueous media. Since first reported in 2008,^{242,243} NITEC has emerged as a pivotal methodology for diverse applications such as surface functionalization, polymer conjugation, protein labeling, and cell imaging especially in the last decade, owing to its favorable kinetics and biocompatibility.²⁴⁴ The 1,3-dipolar cycloaddition reaction between an alkene and a tetrazole represents one elegant and rare example of fluorophore-forming bioorthogonal chemistry. As such, this is an attractive reaction for imaging applications in live cells that requires less intensive washing steps and/or need spatiotemporal resolution.

In 2013, the Zhang group²⁵⁸ introduced photodegradable hydrogels for three-dimensional cell encapsulation (Fig. 40B). These hydrogels, formed by short peptides with a biaryl-substituted tetrazole phototrigger, offer precise spatial and temporal control over cellular microenvironments. Upon light exposure, the phototrigger underwent rapid intramolecular photoclick ligation, yielding a highly fluorescent pyrazoline, facilitating precise imaging. Tet(i)-GFF and Tet(ii)-GFRGD gels, incorporating these triggers, exhibited robust mechanical strength and biocompatibility, supporting extended cell culture. Triggered activation disrupted hydrogel assembly, modulating microenvironments for cells both within and on the gel. This approach is promising for biomedical applications, offering targeted control over cellular environments critical for various research and therapeutic endeavors.

Most photoclick reactions, activated by UV light (300–365 nm), face limitations in biological systems due to phototoxicity. To address this, Lin’s group synthesized oligothiophene-based tetrazoles to minimize phototoxicity. For example, a terthiophene-tetrazole exhibited excellent photoreactivity under 405 nm laser (ring rupture quantum



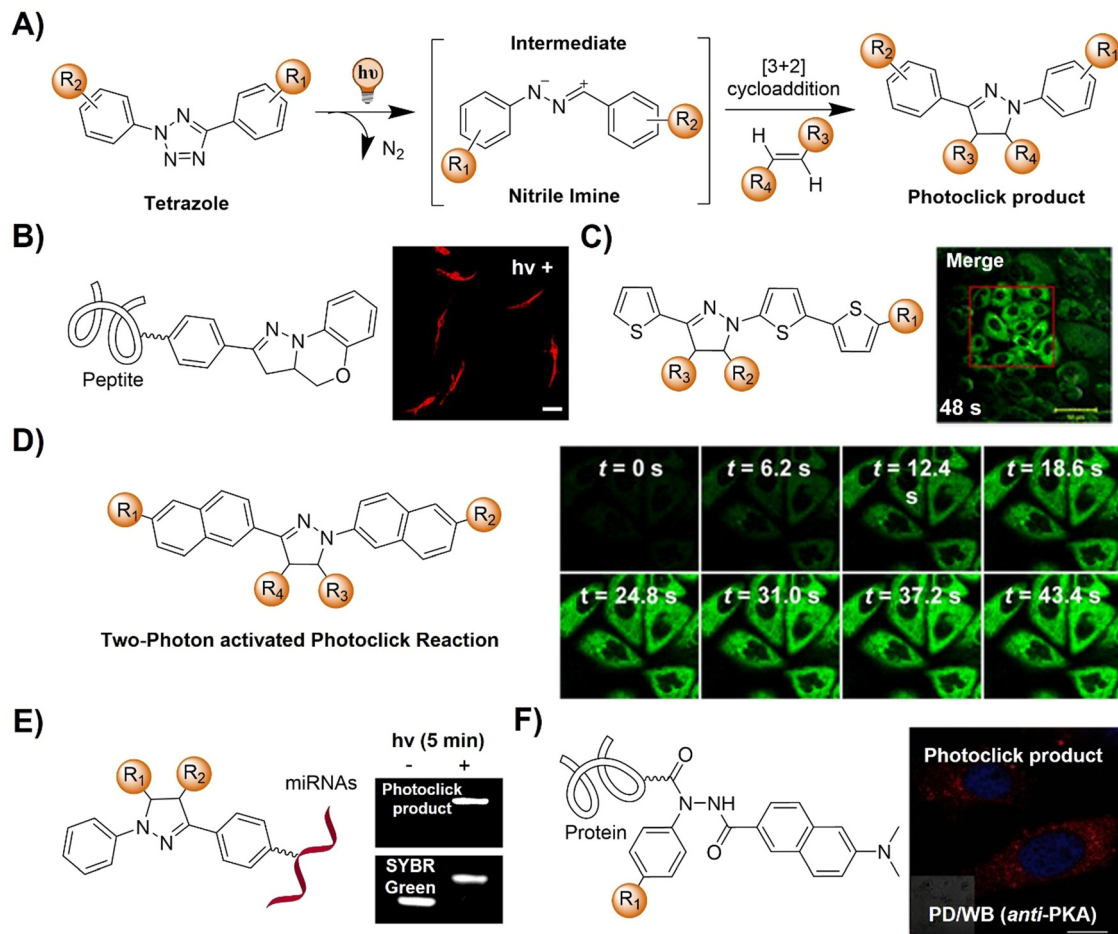


Fig. 40 Fluorescent probes based on the light-triggered nitrile imine-mediated tetrazole–ene cycloaddition. (A) Proposed mechanism of the NITEC photoclick reaction; (B) chemical structure of the Tet(II)-GFRGD photoclick product and morphology of hMSCs encapsulated in Tet(II)-GFRGD gels, adapted with permission from ref. 258. Copyright 2013, American Chemical Society; (C) the structure and *in situ* cell imaging of the oligothiophene-based tetrazole photoclick product, adapted with permission from ref. 259. Copyright 2013, The Royal Society of Chemistry; (D) the structure and *in situ* CHO cell imaging of the naphthalene-tetrazole photoclick product, adapted with permission from ref. 260. Copyright 2013, American Chemical Society; (E) the structure and HepG2 cell imaging of the miRNA-tetrazole photoclick product, adapted with permission from ref. 261. Copyright 2016, American Chemical Society; (F) the structures and live HepG2 cell imaging of the two-photon tetrazole conjugates, adapted with permission from ref. 240. Copyright 2016, Wiley-VCH GmbH.

yield: 0.16, Fig. 40C).²⁵⁹ On the other hand, a water-soluble terthiophene-tetrazole exhibited faster kinetics with a fumarate dipolarophile ($k_2 = 1299 \pm 110 \text{ M}^{-1} \text{ s}^{-1}$), enabling real-time, spatiotemporally controlled precise imaging of microtubules in live mammalian cells. Given the widespread availability of 405 nm lasers in fluorescence microscopes, this class of thiophene-tetrazoles activated by 405 nm lasers have potential for the broader adoption of photoclick chemistry in cell biology research.

To further advance the application of NITEC to living organisms with enhanced spatiotemporal precision, the Lin group devised naphthalene-based tetrazoles activated efficiently *via* two-photon excitation using a 700 nm femtosecond pulsed laser (Fig. 40D).²⁶⁰ In this two-photon bioorthogonal probe, a water-soluble, cell-permeable naphthalene-based tetrazole that exhibits a significant two-photon cross-section of 3.8 GM was identified for the cycloaddition reaction with

acrylamide. This two-photon-triggered, fluorogenic photoclick chemistry was successfully employed for site-selective labeling of alkene-encoded proteins *in vitro* and for visualizing ligand-bound microtubules in cultured cells. With genetic methods enabling the incorporation of unnatural amino acids containing reactive alkene dipolarophiles in both mammalian cells and *Drosophila melanogaster* fruit flies, this technique offers a new approach to dissect protein function in living organisms with exceptional spatiotemporal precision.

MicroRNAs (miRNAs) are pivotal in post-transcriptional gene regulation, yet identifying their interactions with target genes in cells remains challenging due to untagged miRNAs. Zhang *et al.*²⁶¹ introduced photoclickable miRNAs, pre-tagged with molecular handles, maintaining intracellular functionality while allowing attachment of probes *via* photoclick reactions (Fig. 40E). Upon transfection, these miRNAs formed functional complexes that can effectively suppress target gene expression.

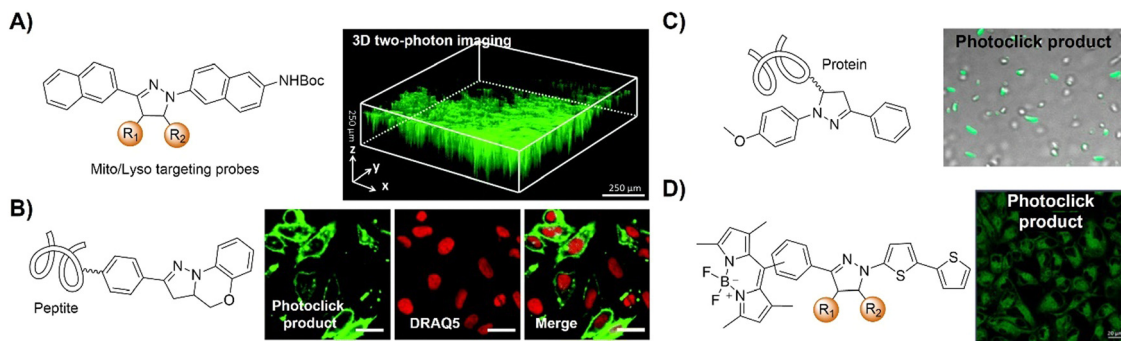


Fig. 41 Bioorthogonal probes for fluorescence cell imaging based on the light-triggered nitrile imine-mediated tetrazole–ene cycloaddition. (A) The structure and 3D two-photon imaging of the naphthalene-substituted tetrazole photocclick product, adapted with permission from ref. 262. Copyright 2016, The Royal Society of Chemistry; (B) the structure and HeLa cell imaging of the biaryl-tetrazole-modified peptide photocclick product, adapted with permission from ref. 263. Copyright 2017, The Royal Society of Chemistry; (C) the structure and *E. coli* cell labeling of the styrene-tetrazole photocclick product, adapted with permission from ref. 264. Copyright 2017, American Chemical Society; and (D) the structure and cell imaging of the bodipy-tetrazole photocclick product, adapted with permission from ref. 265. Copyright 2018, Wiley-VCH GmbH.

Target genes bound to photoclickable miRNAs were tagged for pull-down and identification. Using miR-106a, miR-27, and miR-122, their intracellular function matched intact miRNAs, surpassing biotinylated versions. Attachment of biotin handles to complexes facilitated pull-down and analysis, unveiling a regulatory pathway in miR-122-mediated cell apoptosis in HepG2 cells.

Harnessing the fluorescence quenching capability of tetrazoles, Yao and coworkers²⁴⁰ designed and synthesized a series of tetrazole-Bodipy conjugates (Fig. 40F). Emission of these compounds was quenched, while upon irradiation and protein labeling, their emission was switched on. In water, an 80-fold fluorescence enhancement was observed post-photoactivation due to a reaction with acetic acid, with similar effects observed in the presence of proteins like HSA and BSA. Staurosporine (STS)-labeled tetrazole-Bodipy enabled selective imaging of endogenous protein kinase A (PKA) in HepG2 cells. Intense fluorescence emission in the cell cytoplasm persisted after multiple wash cycles, indicating the formation of a robust covalent bond.

In another study, Zhang and colleagues²⁶² fabricated pyrazoline-based two-photon probes with diverse functionalities for two-photon fluorescence imaging of protease in apoptosis (Fig. 41A). Using this probe system, photoclick reaction of naphthalene-substituted tetrazoles with alkenes was employed to generate naphthalene-substituted pyrazoline, a highly fluorescent two-photon fluorophore, *in situ* under mild and biocompatible conditions. Various types of two-photon probes, such as those targeting the mitochondria and lysosomes, as well as a complex probe incorporating both a targeting ligand (cRGD) and a short peptide substrate for caspase-3 activation, were developed. Notably, these probes exhibited selective activation by caspase-3 in cells or tumor tissue, enabling 3D imaging of caspase-3 distribution in tumor tissue. The versatility of this method in incorporating multiple functionalities into two-photon probes through a gentle photoclick protocol holds promise for constructing diverse imaging probes for targeted tumor imaging.

Following this success, the same research group²⁶³ devised alkaline phosphatase (ALP)-guided self-assembly using a biaryl-tetrazole-modified peptide precursor, Tet-Gly-DPhe-DTyr(H₂PO₃). The peptide precursor could form the photo-responsive hydrogelator Tet-GDFDY upon ALP-mediated dephosphorylation (Fig. 41B). This enabled enzyme-instructed self-assembly-photo-induced disassembly (EISA-PIDA) processes on ALP-overexpressing cancer cell membranes. Tet-GDFDY exhibited rapid fluorescence activation under brief light exposure, facilitating real-time monitoring of EISA-PIDA on HeLa cells. Additionally, photo-triggered disassembly of Tet-GDFDY disrupted pericellular nanofibers, potentially offering a mechanism for combating EISA-induced cell death.

To enhance the fluorogenicity of the alkene-tetrazole reaction for protein labelling, the Gou group²⁶⁴ explored the use of styrene as a dipolarophile for alkene-tetrazole reaction. Remarkably, an over 30-fold increase was obtained in fluorescence quantum yield for the reaction product in aqueous environments (Fig. 41C). Mechanistic investigations suggested that this enhancement stems from inadequate protonation of the styrene-tetrazole reaction product. This discovery offered valuable insights for optimizing alkene-tetrazole reactions in biological research. The efficacy of the styrene-tetrazole reaction for fluorogenic protein labeling was demonstrated both *in vitro* and *in vivo* through genetic incorporation of a styrene-containing unnatural amino acid. The combination of styrene with various tetrazole derivatives promises the enhanced and diversified application of alkene-tetrazole chemistry in live-cell real-time imaging.

The development of photoclick reaction initialized responsive probes for the detection of biomolecules, such as ROS, was reported by the Lin group (Fig. 41D).²⁶⁵ Three bodipy-tetrazoles featuring *para*-, *meta*-, or *ortho*-linkages between the bithiophene-tetrazole and the bodipy fluorophore were synthesized and their photoclick and sensing reactions were investigated. These compounds exhibited remarkable reactivity towards dimethyl fumarate *in vitro* in photoclick reactions. Intriguingly, the fluorescence of bodipy experienced



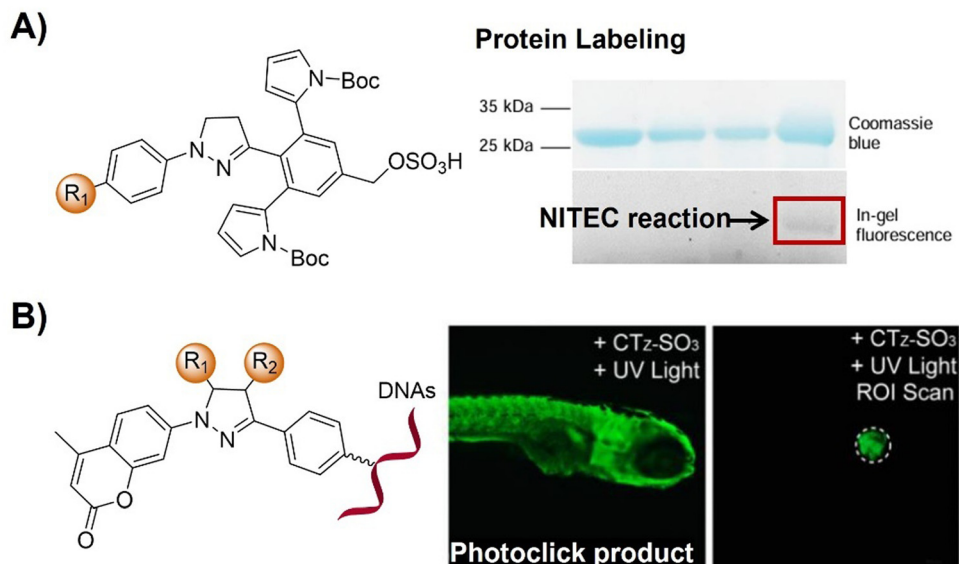


Fig. 42 Fluorescent probes for protein labelling and *in vivo* imaging based on the light-triggered nitrile imine-mediated tetrazole–ene cycloaddition. (A) The structure and protein labeling of the sterically shielded nitrile imine photoclick product, adapted with permission from ref. 266. Copyright 2018, American Chemical Society; (B) the structure and live zebrafish imaging of the coumarin-fused tetrazole photoclick product, adapted with permission from ref. 267. Copyright 2019, American Chemical Society.

a significant decrease post photoclick reaction; yet, this fluorescence loss could be reversed by treatment with chemical oxidants like H_2O_2 . A water-soluble bodipy-tetrazole derivative was also developed, serving as an “off-on” fluorescent probe for the detection of H_2O_2 . This probe facilitated the initial bioorthogonal *in situ* generation at microtubules and subsequent detection of H_2O_2 within HeLa cells.

For rapid bioorthogonal labelling of proteins in live cells, the same research group employed a bioinspired approach for designing sterically shielded tetrazoles for bioorthogonal photoinduced cycloaddition with alkene dipolarophiles (Fig. 42A).²⁶⁶ The resulting photogenerated nitrile imine exhibited significant enhancement of selectivity for cycloaddition over nucleophilic addition. Computational and structural analyses indicated that adjacent *N*-Boc-pyrrole groups provided a steric shield, raising the activation energy to prevent thiolate addition. *In situ* studies revealed a notably prolonged half-life of 102 s for the shielded nitrile imine compared to its unshielded counterpart. Leveraging the environmentally sensitive pyrazoline fluorophore produced by the cycloaddition, the shielded tetrazole enabled the creation of a fluorescent sensor for monitoring ligand-induced conformational transitions of glutamine binding protein (QBP). Additionally, due to its rapid kinetics and exceptional selectivity, the shielded nitrile imine efficiently labeled an alkene encoded GCGR in live mammalian cells at a concentration as low as 500 nM within 1 min, surpassing the efficiency of optimized Tz reagents. Apart from protein labelling, the “photoclick” method for nucleic acid labeling was explored by Zhang and coworkers in 2019,²⁶⁷ in which fluorescence “switch-on” labeling of synthesized DNA with coumarin-fused tetrazole probes was achieved (Fig. 42B). In this nucleic acid labeling system,

DNA and electron-rich alkenes could quench pyrazoline fluorescence while minimally affecting the emission of coumarin fluorophores. **CTz-SO₃**, a water-soluble coumarin-fused tetrazole probe designed in this research, could precisely label and visualize 5-vinyl-2'-deoxyuridine (VdU)-incorporated cellular DNA in real time without fixation or DNA denaturation.

In addition to protein and nucleic acid labelling, intramolecular photoclick reactions with fluorogenic probes also enabled the labelling of subcellular structures, like lysosomes and mitochondria. For example, Huang *et al.*²⁶⁸ devised a straightforward strategy for designing tetrazole-based photoactivatable fluorophores (Mt-Tet and Ly-Tet) that target cellular mitochondria and lysosomes guided by TPP and morpholine, respectively, and exhibit turn on fluorescence *via* intramolecular photoclick reaction (Fig. 43A). Upon mild photoirradiation, robust fluorescence activation occurred within ~1 min *in vitro* and ~3 min in live cells, allowing for real-time imaging of subcellular organelles. These probes exhibited minimal fluorescence prior to activation, resulting in a reduced background signal compared to standard fluorescence labeling. The spatiotemporally controllable nature of these probes enabled the observation of organelle dynamics by super-resolution Stochastic Optical Reconstruction Microscopy (STORM) imaging.

Sialic acid plays a crucial role in profiling cell activities across various biological and pathological contexts. To understand the biological functions of sialic acids under endogenous conditions, photoclick based bioorthogonal probes have been developed for their precise labelling by Zhang and colleagues.²⁶⁹ With this probe system, upconverting nanoparticles (UCNPs) that can be activated by NIR light while emitting UV-visible light were used to prepare the nanoprobe



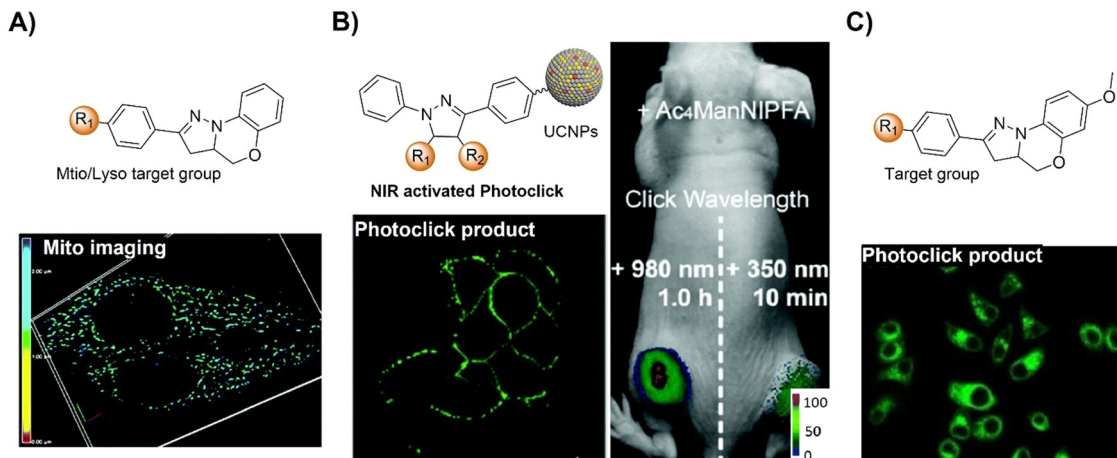


Fig. 43 Bioorthogonal probes for fluorescence bioimaging based on the light-triggered nitrile imine-mediated tetrazole–ene cycloaddition. (A) The structure and super-resolution STORM imaging of the intramolecular **NITEC** photoclick product, adapted with permission from ref. 268. Copyright 2021, The Royal Society of Chemistry; (B) the structure, *in vitro* and *in vivo* imaging of the NIR light-activated **NITEC** photoclick product, adapted with permission from ref. 269. Copyright 2020, The Royal Society of Chemistry; and (C) the structure and endoplasmic reticulum (**ER**) fluorescence labeling of the fluorescent pyrazoline, adapted with permission from ref. 270. Copyright 2022, Wiley-VCH GmbH.

(tetrazole-UCNP) for spatiotemporal labeling of sialic acids (Fig. 43B). Combined with stable *N*-alkene-*D*-mannosamine (Ac₄ManNIPFA), this strategy facilitated real-time imaging of metabolically synthesized alkene sialic acids on cell surfaces through fluorogenic cycloaddition. In addition, spatially selective sialic acid visualization was achieved in specific tumor tissues of mice under NIR light activation, enabling *in situ* controllable labeling in complex biological systems.

Effective labelling of decarboxylases enables direct monitoring of oxidative decarboxylation *in situ* in biological samples. In 2020, Schwaneberg and coworkers²⁷¹ devised a robust 96-well photoclick assay for quantifying styrene derivatives and other alkene products resulting from decarboxylation reactions in the micromolar range. This method features user-friendly handling and fluorescence read-out that facilitate high-throughput screening of decarboxylase variant libraries within hours. The assay requires minimal instrumentation, leveraging commonly available UV light irradiation. To demonstrate the application of this method, analysis of the specificity for different substrates of OleT-BM₃R fusion enzyme was evaluated, holding promise for enhancing substrate selectivity, H₂O₂ resistance, and optimizing electron transfer efficiency in OleT and other decarboxylases.

An *et al.*²⁷⁰ introduced a photoactivatable fluorogenic azide–alkyne click (**PFAAC**) reaction based on the azide-tetrazole probe platform. Upon light irradiation, selective photolysis of azide and tetrazole rendered these probes nonfluorescent, generating photoactivatable fluorescent reporters post-click reaction (Fig. 43C). This chemistry enabled specific detection of AAC reactions with a remarkable over 1000-fold “turn-on” ratio in a spatiotemporally controlled manner, validated in live cell imaging for organelle labeling. Incorporating a chemical linker into the azide-tetrazole probe expanded **PFAAC**’s utility, allowing for dual-labeling of biomolecules, as confirmed in endoplasmic reticulum (**ER**) fluorescence labeling with

pyrazoline and *c*y5 dyes. Given the widespread use of click chemistry, **PFAAC** holds broad potential for biomolecular imaging, tracking, and detection.

In summary, since 2008, the **NITEC** photoclick reaction has gained increasing attention in various biological applications. Unlike other click reactions, such as CuAAC and IEDDA, **NITEC** offers unique features like substrate stability in physiological environments, low toxicity, and fluorogenicity enabling efficient sensing and imaging of specific biomolecules. While primarily used for bioorthogonal protein labeling and wash-free *in situ* imaging, tetrazoles have also modified nucleic acid structures and have been involved in the development of biomaterials like hydrogels. Despite optimizations, drawbacks like UV-restricted activation wavelengths and collateral reactions with biological nucleophiles have hindered their biological utility. Strategies to shift activation wavelengths towards the NIR or explore photocatalysis with UCNPs have shown some promise. Addressing collateral reactions, especially with the formation of undesired pyrazoline, requires modification of the tetrazole scaffold. However, such reactions can also be harnessed for applications like studying protein–protein interactions. With increasing interest in photoclick chemistry and the versatility of light-generated intermediates, photoactivatable tetrazoles hold immense potential across various research fields, from biology to materials science, with scope for further optimization and exploration in novel applications.

2.7.2. PQ-ERA photocycloaddition. The phenanthrenequinone (**PQ**)-electron-rich alkene (**ERA**) [4+2] photocycloaddition reaction, first uncovered in the 1940s by Schönberg and coworkers,²⁵⁵ involves the interaction of photoexcited **PQ** with **ERAs** such as styrene, stilbene, and triphenyl ethylene. The reaction mechanism initiates with the photoexcitation of **PQ** using UV to blue light (Fig. 44A). Upon excitation, ground state **PQ** populates the S₂ (1 $\pi\pi^*$) state, undergoing internal conversion to form the lowest excited S₁ (1n $\pi\pi^*$) singlet state. A highly



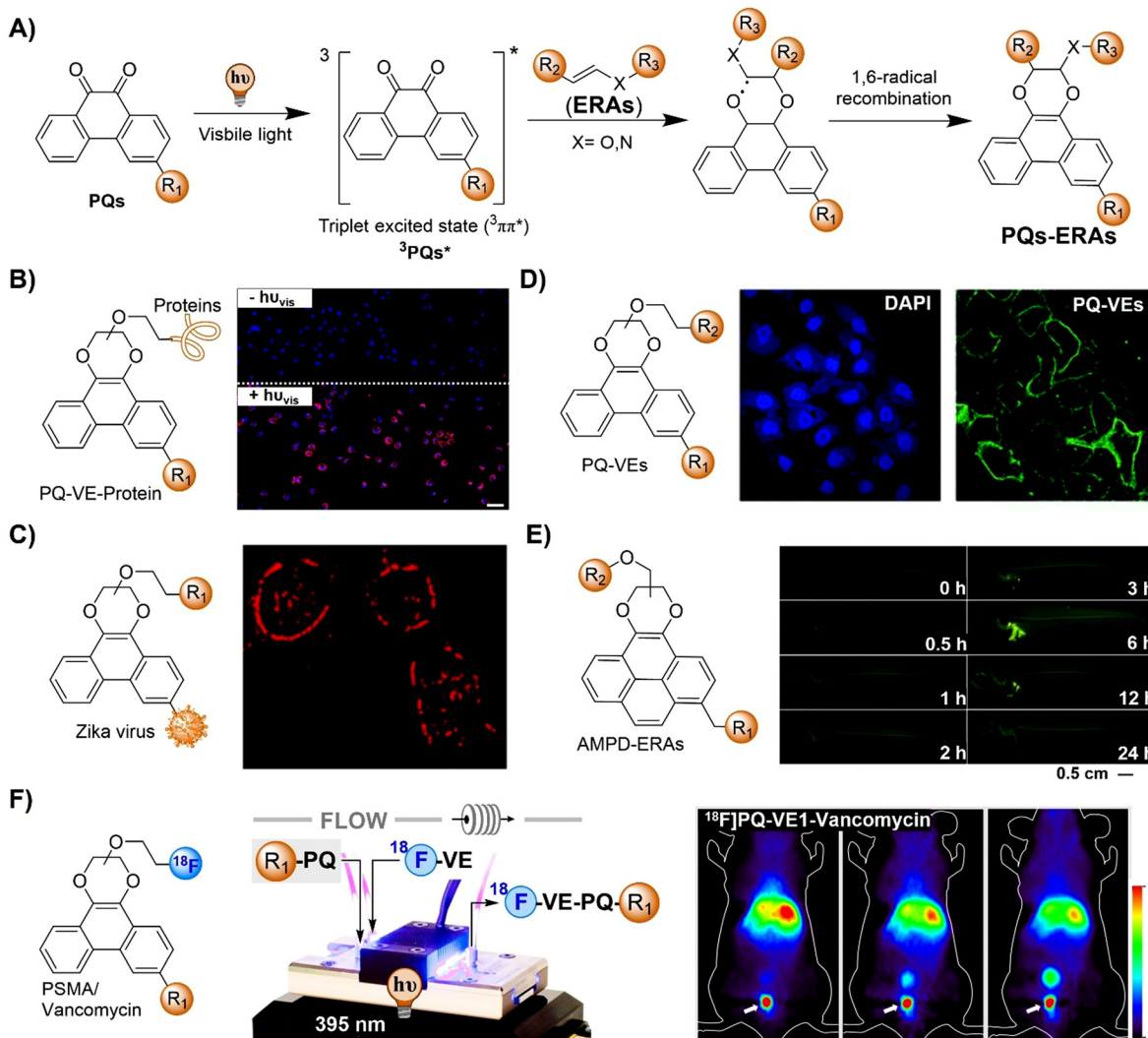


Fig. 44 PQ-ERA photocycloaddition reaction based bioorthogonal probes for fluorescence imaging. (A) Proposed mechanism of the **PQ-ERA** photocycloaddition reaction; (B) spatial labeling of cetuximab on the surface of A549 cells with pretagged VE that was further labeled through the **PQ-ERA** photocycloaddition reaction, adapted with permission from ref. 254. Copyright 2018, American Chemical Society; (C) **PQ-ERA** photocycloaddition for the labeling and visualization of the Zika virus, adapted with permission from ref. 275, licensed under CC-BY 4.0; (D) sequential photoirradiation-orthogonal reporter tagging (SPORT) for cell labelling via **PQ-ERA** photocycloaddition, adapted with permission from ref. 276, under the license CC BY-NC 3.0 DEED; (E) the structure and *in vivo* imaging of the **AMPD** probe on glass catfish; adapted with permission from ref. 277. Copyright 2024, Elsevier B.V; and (F) short-lived ^{18}F -PET tracers prepared via **PQ-ERA** photoclick reaction and their application in PET imaging, adapted from ref. 253, licensed under CC-BY-NC-ND 4.0.

efficient intersystem crossing (ISC) process ensues, populating the two lowest triplet states, including T_1 ($^3\pi\pi^*$) and T_2 ($^3\text{n}\pi^*$). Notably, the reactive state for the **PQ-ERA** reaction (Fig. 44A) is T_1 , with T_2 also present and closely aligned in energy. Subsequently, a [4+2] photocycloaddition with an **ERA** occurs, enabling the reaction to form the fluorescence product (Fig. 44A).^{272–274} Despite its efficacy, the reliance on a potent mercury lamp as a light source and the prolonged reaction time (days) have impeded its broader application in synthesis for over 80 years.

In 2018, Zhang and colleagues addressed the limitations of the mercury lamp light source by employing a suitable high-power LED light source.²⁵⁴ Using this hand-held LED lamp, the **PQ-ERA** photoclick reaction could be executed within a short

time frame of 5 min, achieving high yields (up to 93%) even in an aqueous environment (Fig. 44B). The application of the **PQ-ERA** photoclick reaction system was demonstrated for labeling cell membranes in living cells, underscoring its biocompatibility and potential for biological applications (Fig. 44B). Shortly after, Zheng *et al.* introduced the **PQ-ERA** bioorthogonal cycloaddition technique to construct a fluorogenic probe in 2022²⁷⁵ for labeling and visualization of the Zika virus through a straightforward process mediated by visible light (450 nm, Fig. 44C).

Based on the **PQ-ERA** photoclick reaction, Xie and coworkers²⁷⁶ introduced a method, namely sequential photoirradiation-orthogonal reporter tagging (SPORT), for cell labelling (Fig. 44D). This technique enabled the controlled labeling of surface

biomolecules in live cells using visible LED light. Introducing metabolic precursors into live cells triggered a bioorthogonal reaction to convert azide/alkyne functional groups into a photoactive functional group known as **PQ**. Subsequently, the **PQ** groups underwent visible light-induced photoclick reactions with vinyl ether-biotin using a white light LED. The effectiveness of this approach was demonstrated by selective labeling of a variety of biomacromolecules, including sialome, mucin-type *O*-glycome, phospholipids and membrane proteins, highlighting the **SPORT** strategy for precise investigations of biomolecules within complex biological systems.

Multimodal bioorthogonal small molecule probes also contribute significantly to tracking drugs in living organisms. For instance, Bai and colleagues²⁷⁷ introduced 1-(azidomethyl) pyrene-4,5-dione (**AMPD**), a versatile probe incorporating adjacent dione structures at the pyrene core for drug target detection *in vivo* (Fig. 44E). **AMPD** selectively interacts with **ERA**-labeled drugs under blue LED light, triggering specific *in situ* fluorescence for *in vivo* tracking and flow cytometric sorting. The methyl azide group enhances target protein enrichment *via* click chemistry. With exceptional biocompatibility, **AMPD** facilitated the visual tracking in live animals, tissues, cells, and gels, expediting drug discovery by improving comprehension of *in vivo* drug-target interactions. Beyond its utility in optical imaging, the **PQ-ERA** photoclick reaction has diversified its applications across a spectrum of molecular imaging techniques, including PET imaging. The Feringa group pioneered the development of an expedited photoclick method for synthesizing short-lived ¹⁸F-PET tracers, leveraging the **PQ-ERA** photoclick reaction (Fig. 44F).²⁵³ The corresponding precursors are readily accessible synthetically and can be tailored with various targeting moieties. Using a flow photo-microreactor, the photoclick reaction could be completed within 60 s, facilitating the rapid generation of clinically relevant tracers for prostate cancer, bacterial infection imaging and *in vivo* PET imaging,²⁷⁸ thus underscoring the practicality of this approach.

The **PQ-ERA** reaction, as an emerging photoclick system, is particularly favored due to its excellent biocompatibility, use of visible light, and the inherent fluorescence properties of the products. This has led to its widespread adoption for high spatiotemporal resolution light-activated bioimaging.²⁷⁹ Nevertheless, the light to be used for activation of the click reaction at about 400 nm or even visible light exhibits limited tissue penetration depth, impeding its application for *in vivo* deep tissue photoclick labelling and imaging. Moreover, the reaction rate of the **PQ-ERA** photoclick reaction is relatively low, approximately $1\text{ M}^{-1}\text{ s}^{-1}$, necessitating stronger light sources and longer illumination periods, potentially leading to phototoxicity concerns.

To overcome these challenges, Feringa's group recently introduced a series of modified **PQ-ERA** systems. Employing triplet optimization,^{272,273,280} triplet-triplet energy transfer (TTET),²⁷⁴ and upconversion nanoparticles (UCNPs),²⁸¹ they successfully engineered ultrafast **PQ-ERA** reactions with reaction rates (k_2) reaching up to $1970\text{ M}^{-1}\text{ s}^{-1}$. These reactions were induced by long-wavelength visible light/NIR light,

extending up to 980 nm. While the biocompatibility of these enhanced systems requires further refinement, these probes exhibit promise for the biological application of the **PQ-ERA** photoclick reaction.

2.7.3. Diarylsyndone-alkene photoclick reaction. Sydnone (**Syds**) are unique molecular structures with remarkable physical and chemical properties (di-polar with delocalized positive and negative charges across the ring) and a broad range of biological activities and have potential as precursors for synthesizing diverse heterocyclic systems.²⁸² Upon photoirradiation, sydnone can convert into the nitrile imine intermediates with elegant applications in bioorthogonal photoclick chemistry (Fig. 45A). The Yu group has made enormous efforts in the field of diarylsyndone-alkene photoclick reactions. For instance, inspired by the remarkable reactivity observed in a specific reaction, the Yu group²⁴⁹ introduced a highly efficient visible light-induced diarylsyndone (**DASyD**)-alkene cycloaddition (**photo-DASAC**) method with fluorogenic properties (Fig. 45A–C). This approach involved synthesizing a library of **DASyds** and evaluating them in light-induced 1,3-dipolar cycloaddition reactions. Among these compounds, **DASyD** exhibited exceptional reactivity in the fluorogenic photoclick reaction with various alkenes such as **TCO**, norbornene, methyl methacrylate, diethyl fumarate, and 5-vinyl-2'-deoxyuridine. The emission at 533 nm was increased significantly, allowing for the application of **DASyD** in labelling of **TCO**-appended proteins, *e.g.*, lysozyme (lyso-**TCO**).

Following this success, the Yu group²⁸³ introduced rapid photoclick reactions between **DASyD** and strained alkynes, specifically **BCN**, illustrated in Fig. 45D. The reaction between **DASyD** and **BCN** exhibited efficient and selective progress under 405 nm light irradiation. Although competing cycloaddition products emerged from the non-light-activated **DASyD**-alkyne pathway, their occurrence was noticeably slower compared to the **photo-DASAC** process (Fig. 45D). The slow thermal-mediated cycloaddition reaction was ascribed to steric repulsion induced by the adjacent *ortho*-diaryl units on the twisted **DASyD**, hindering the non-light-activated **DASAC** pathway. To highlight the biocompatibility of this reaction, selective labelling of proteins was achieved on the surfaces of A549 cells (Fig. 45D). Subsequently, the same group engineered a set of photoactivatable β -diarylsyndone-1-alanines (**DASAs**) known for their outstanding photoreactivity and heightened fluorescence turn-on when interacting with alkenes through a **photo-DASAC** reaction in a biocompatible setting (Fig. 45E).²⁸⁴ The resulting fluorescent pyrazoline-alanine exhibited impressive environmental sensing properties, enhancing its probe capabilities. Integrating the **DASA** residue onto the side chain of linear peptides led to the formation of macrocyclic peptides through the *in situ* photo-cyclization with the alkene residue. These peptides demonstrated fluorogenic translocation across live cell membranes.

2.7.4. Other photoclick approaches. Apart from the above three photoclick reactions, other photoclick reactions have also been utilized for developing bioorthogonally activated probes



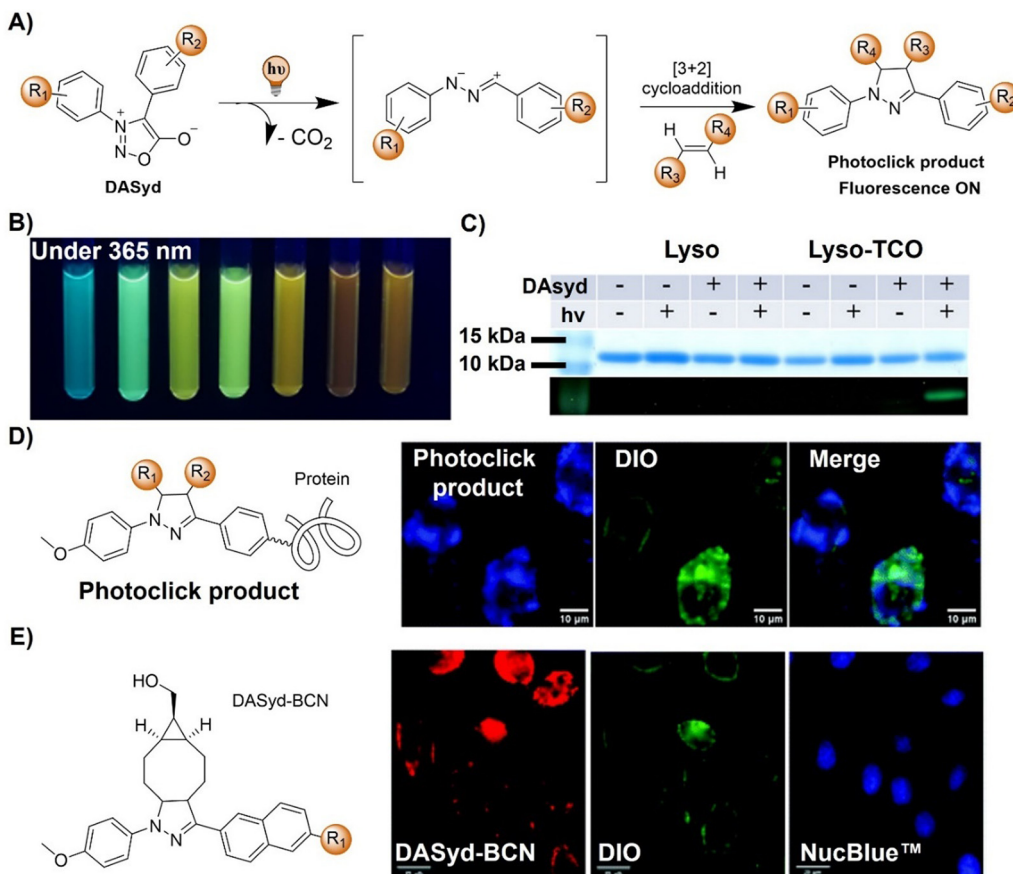


Fig. 45 Bioorthogonal probes for fluorescence imaging based on the diarylsyndone–alkene photoclick reaction. (A) Proposed mechanism of the **photo-DASAC** photoclick reaction; (B) fluorescence of different **photo-DASAC** photoclick products, under excitation at 365 nm; (C) selective photoinduced fluorogenic labeling of proteins; adapted with permission from ref. 249. Copyright 2018, American Chemical Society; (D) the structure of the **photo-DASAC** product and its precise labelling of living A549 cells, adapted with permission from ref. 283. Copyright 2019, The Royal Society of Chemistry; and (E) the structure of the **DASyd-BCN** and *in situ* fluorescence imaging of live A549 cells, adapted with permission from ref. 284. Copyright 2019, The Royal Society of Chemistry.

for *in situ* bioimaging. For instance, the Yu group²⁸⁵ reported a photo-induced defluorination acyl fluoride exchange (**photo-DAFEx**), utilizing acyl fluorides from *m*-trifluoromethylaniline's photo-defluorination to conjugate with primary/secondary amines and thiols in water (Fig. 46). Computational results and experiments revealed that water-mediated cleavage of the *m*-NH₂PhF₂C(sp₃)-F bond in the excited triplet state plays a crucial role in defluorination. The formed benzoyl amide linkages exhibited satisfactory fluorogenicity, enabling *in situ* visualization of this photoclick reaction. This photoclick strategy was used to label small molecules, cyclized peptides, and functionalized proteins *in vitro* and to develop photo-affinity probes targeting endogenous carbonic anhydrase II (hCA-II) in living cells.

In summary, photoclick reactions offer high spatial-temporal control and rapid reaction kinetics, making them valuable for precise optical imaging. However, several limitations need to be considered when designing photoclick reaction-based bioorthogonal probes. For instance, photoclick reactions are generally triggered by UV or short-wavelength-visible light (violet or blue light, ranging from 380 to

465 nm), which are potentially highly phototoxic in biological studies. The limited penetration ability of short-wavelength light hampers the application of photoclick reaction in deep biological tissue and for *in vivo* imaging. Although two-photon excitation and UCNPs NITEC reactions have been developed for cell imaging, the efficiency of photoclick reaction remains low requiring prolonged exposure to high-intensity laser illumination, diminishing the practical value. In terms of upconversion luminescence-based photoclick reactions, the toxicity of UCNPs presents another concern for biological studies *in vivo*. Future research in this domain will need to concentrate on developing photoclick systems with high biocompatibility and activity triggered by long-wavelength visible light and NIR light, to enhance their applicability for *in situ* precision imaging, both *in vitro* and *in vivo*.

2.8. Bioorthogonal nanozyme-catalyzed reactions

The incorporation of transition metal catalysts into nanoscaffolds enables the development of nanocatalysts that replicate crucial enzymatic behavior. These catalysts, known as bioorthogonal nanozymes,²⁸⁶ engage in bioorthogonal chemistry not

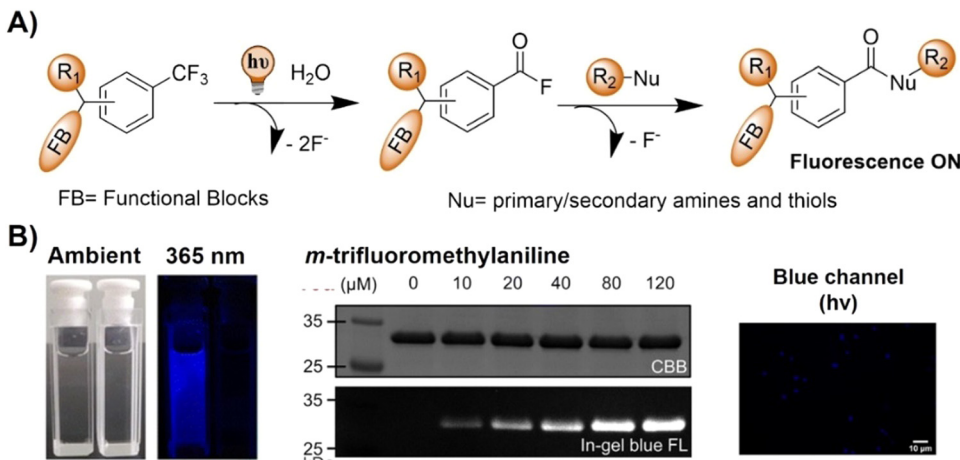


Fig. 46 Bioorthogonal probes developed using other photoclick reactions for fluorescent protein labelling and cell imaging. (A) Proposed mechanism and (B) fluorescence changes, protein labeling and cell imaging of the photo-DAFEx photoclick reaction, adapted with permission from ref. 285. Copyright 2023, The Royal Society of Chemistry.

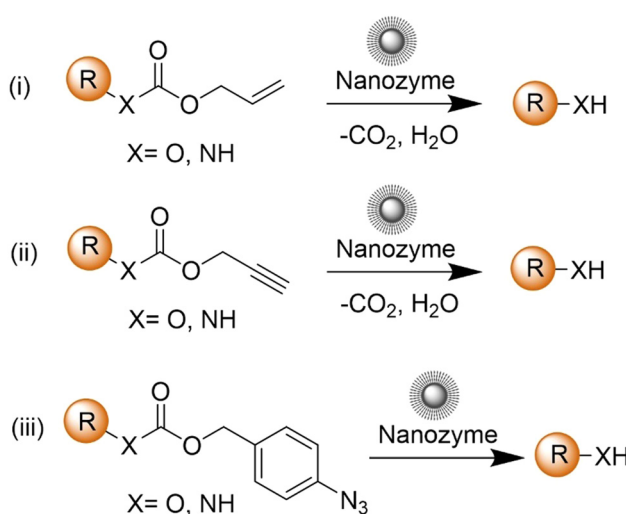
accessible to natural living systems (Scheme 7).^{287,288} This unique capability allows bioorthogonal nanozymes to function as on-site “factories” for producing bioactive molecules as needed. Co-engineering the transition metal catalysts and the nanometric scaffold is essential to create highly effective bioorthogonal nanozymes. In this section, we will provide an overview of recent advances in the field, with a specific focus on the application of bioorthogonal nanozymes for *in situ* imaging.

Artificial nanoscale catalysts with high turnover are excellent alternatives for initializing bioorthogonal activation for specific labelling and imaging of cellular molecules. In 2015, Meggers and coworkers²⁸⁹ engineered cytochrome P450 monooxygenase enzyme variants that are capable of efficiently removing protecting groups from propargylic and benzylic ethers for bioorthogonal activation (Fig. 47A). These modified cytochrome P450 monooxygenases deprotected compounds *in vitro* and within living *E. coli* cells, releasing fluorescent

alcohols that are suitable for *in situ* bioimaging. Results of this research demonstrated the potential of bioorthogonal enzyme/protecting group pairs for targeted release of imaging agents or catalytic activation of prodrugs at specific sites.

Bioorthogonal catalysis has been widely adopted in investigating intracellular chemistry, while delivering and controlling synthetic catalytic systems within cells remains challenging. To tackle this issue, Rotello and colleagues²⁹¹ developed a bioorthogonal nanozyme based on AuNPs, employing transition metal catalysis for *in situ* imaging and therapy (Fig. 47B). These nanozymes with protein size and surface features offered promise for diverse *in vitro* and *in vivo* applications. Controlling their activity *via* CB[7] host-guest interactions with AuNP ligands’ benzyl headgroups has enabled efficient and reversible catalysis regulation. This strategy could be further used to promote therapeutic applications like prodrug activation to treat non-cancerous chronic diseases. The gated platform ensures controllable catalyst loading, reducing interference and protecting against poisoning, release, distribution, or clearance. The application of this biomimetic and bioorthogonal system was then demonstrated in imaging and prodrug systems through triggered cleavage of allylcarbamates for activating fluorescence and propargyl groups for the release prodrug inside living cells, respectively.

Traditional methods involving Pd catalysts are limited by their poor cellular selectivity, requirement for stoichiometric or excessive Pd complex, and potential cellular toxicity. Therefore, targeted localization of Pd catalysts within cells would advance their applications in biological investigations. Bradley *et al.*²⁹⁰ expanded Pd-catalyzed chemistry in living systems by introducing a method for delivering and tracking an active organometallic complex within mammalian cells (Fig. 47C). These water-soluble and traceable catalysts, based on Pd(II)-carbene complexes linked to a fluorescently labeled homing peptide, enabled targeted cell delivery. This biocompatible homogeneous Pd catalyst, built on a previously reported carbene Pd



Scheme 7 Illustration of bioorthogonal nanozyme-catalyzed reactions.



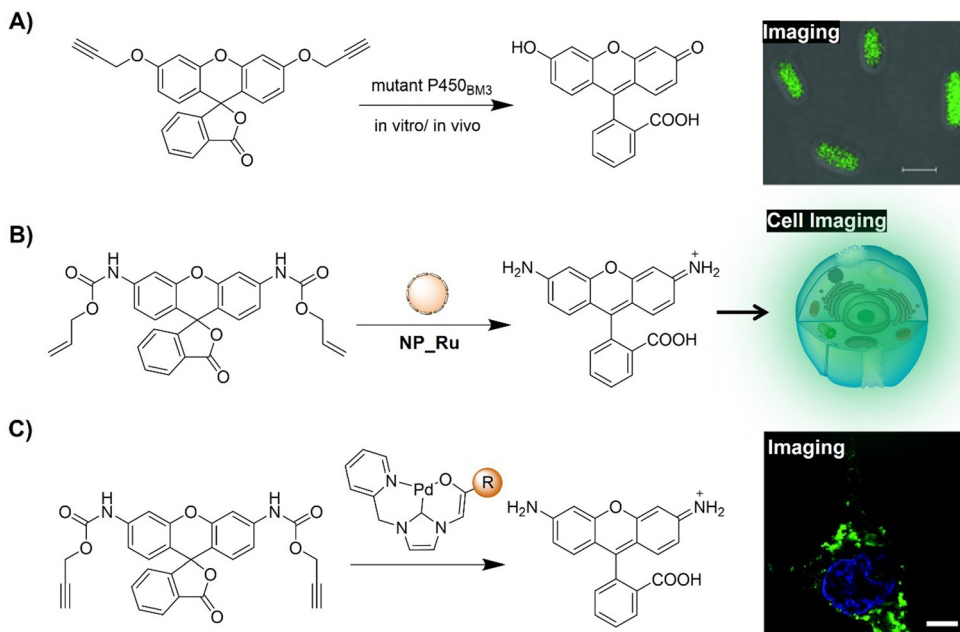


Fig. 47 Bioorthogonal nanozyme-catalyzed reaction-based probes for fluorescence cell imaging. (A) Mutant P450_{BM3} catalyzed decaging of Proc-FITC and bioimaging of *E. coli*, adapted with permission from ref. 289. Copyright 2015, Wiley-VCH GmbH; (B) schematic illustration of NP_Ru catalyzed decaging of Proc-rhodamine; and (C) Pd catalyzed decaging of Proc-rhodamine and confocal microscopy images of PC-3 cells, adapted with permission from ref. 290. Copyright 2017, The Royal Society of Chemistry.

ligand coupled to a peptide, exhibited low-cytotoxicity and significant catalytic activity within mammalian cells, allowing for application in bioorthogonal activation of a profluorophore for prostate cancer cell identification and imaging.

Bioorthogonal catalysts are also able to be employed to manipulate the fate of exo-/endo-genous molecules in complicated biological systems. For instance, Broceta and coworkers²⁹² engineered a heterogeneous catalytic system, unlocking previously inaccessible chemical traits of Au-NPs within biological contexts (Fig. 48A). In this catalytic system, bioorthogonal uncaging of various cytotoxic precursors was achieved *via* unexplored gold reactivity, offering a safe avenue for therapeutic activation through nonbiological routes. Moreover, this solid-supported catalyst could promote the localized release of fluorescein within the brain of a zebrafish, enabling precise *in situ* imaging. Results of this research enabled chemical modulation of the activity of bioorthogonal reagents in living organisms.

Manipulating biological processes by bioorthogonal bond cleavage reactions facilitates *in vivo* therapeutic applications. For example, Wu and colleagues²⁹³ devised a bioorthogonally activated prodrug for precise bioimaging and therapy, comprising a Pd-mediated cleavable propargyl, a coumarin fluorophore, and a potent anticancer agent (Fig. 48B). *In vitro* studies revealed that the Pd complex triggered propargyl cleavage, initiating a cascade of reactions to activate the coumarin fluorophore for imaging and releasing the anticancer drug for therapy. These components were encapsulated in phospholipid-liposomes, forming a two-component bioorthogonal nanosystem. Internalized by HeLa cells, the nanosystem exhibited

strong intracellular fluorescence and effectively inhibited tumor growth in a mouse model. The success of this research validated the potential of appropriately formulated bioorthogonal systems for *in vivo* applications, offering a new approach to develop imaging-guided prodrug delivery systems.

Remote control of the catalytic activity of nanoreactors by magnetic fields is promising for advancing bioorthogonal chemistry in biological studies. In research led by Lee,²⁹⁴ MAG-NERs, hollow-silica-based nanoreactors that are responsive to magnetic induction, were engineered for bioorthogonal organic synthesis in live cells (Fig. 48C). Catalytic metal (Pd) growth within the silica's confines, guided by magnetic fields, accumulated at the Fe₃O₄-core interface. Pd nanoclusters at this interface catalyzed carbocyclization to activate the fluorescence of nonfluorescent substrates under physiological conditions. MAG-NERs enabled bioorthogonal catalysis at ambient temperature *in vitro* under remote operation. Customization with diverse catalytic nanoclusters enhanced the activity with a biofriendly, remotely controllable magnetic field, advancing biomedical and biological precision imaging applications.

In addition to the remote magnetic field, integrating bioorthogonal catalysts into thermal responsive nanoparticle platforms yields bioorthogonal "nanozymes" that are controllable *via* endogenous or exogenous thermal stimuli. In research led by Rotello,²⁹⁵ thermoresponsive nanozymes were prepared by confining porphyrin (FETTP) supramolecular assemblies within gold nanoparticle monolayers (Fig. 48D). Retaining stimuli-responsive behavior in biological settings, these assemblies enabled localized therapy and *in situ* imaging. Upon heating, the confined assemblies within the scaffold reversibly



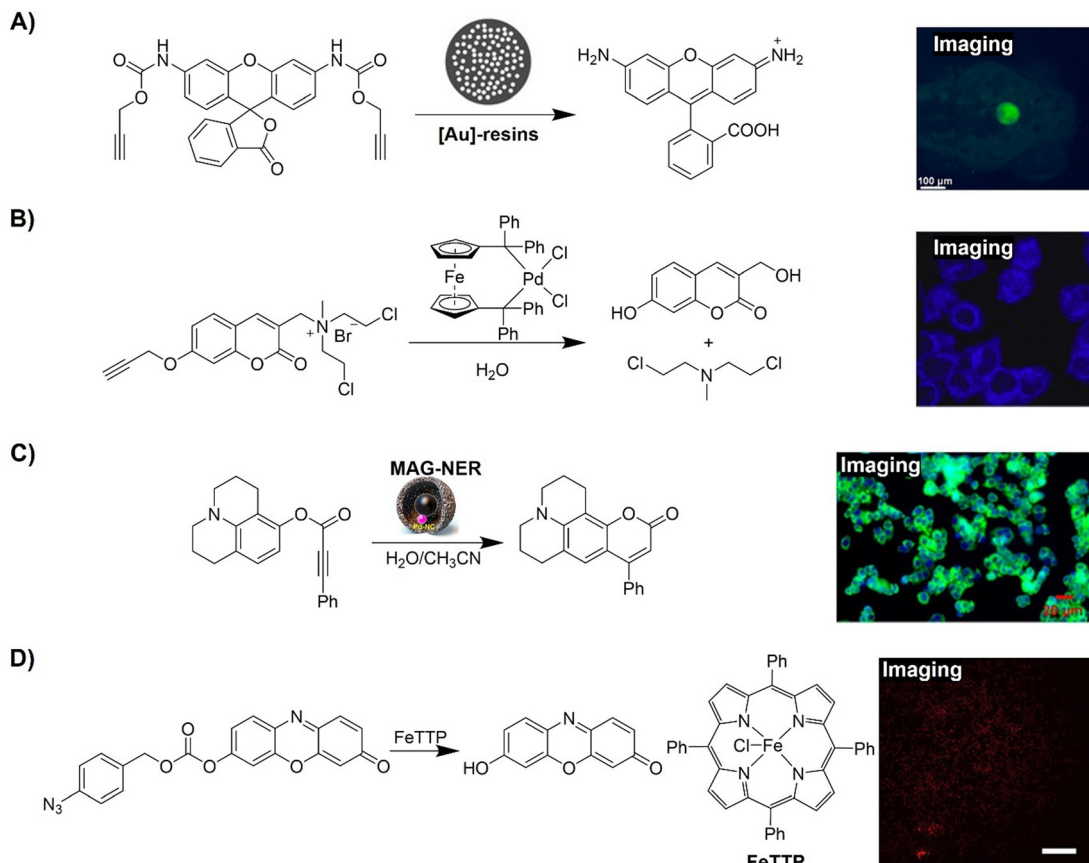


Fig. 48 Fluorescent probes for cell and *in vivo* imaging developed using bioorthogonal catalyzed reactions. (A) [Au]-resin catalyzed decaging of Procrhodamine and precise *in situ* imaging of zebrafish brain, adapted with permission from ref. 292. Copyright 2017, Wiley-VCH GmbH; (B) Pd complex catalyzed activation of the coumarin fluorophore for cell imaging and releasing the anticancer drug for therapy, adapted with permission from ref. 293. Copyright 2017, Elsevier Ltd; (C) MAG-NER catalyzed carbocyclization to activate the fluorescence of nonfluorescent substrates for precise imaging applications, adapted with permission from ref. 294. Copyright 2020, American Chemical Society; and (D) conversion of the profluorophore pro-Res through the **FeTTP** catalytic reduction of aryl azides, adapted with permission from ref. 295. Copyright 2020, Elsevier Inc.

dissociated, restoring catalytic activity. This gating mechanism, driven by confined assembly dynamics, resulted in nanozymes with thermoresponsive behavior in complex biological systems. This thermal responsive and reversibly activated system was then used for activating antibiotic based prodrugs for bacterial biofilm elimination.

In another example of bioorthogonal activation of prodrug therapeutics, Rotello and coworkers²⁹⁶ demonstrated selective intracellular activation of molecules using endogenously activated bioorthogonal nanozymes (Fig. 49A). In this research, AuNP ligand structures were able to facilitate protein corona formation, which in turn controlled the catalytic activity of the nanozyme. A hard “irreversible” corona deactivated nanozymes *via* aggregation and steric hindrance, while a soft “reversible” corona partially reduced the activity of nanozymes. Proteolytic degradation of the corona by endogenous proteases in endosomes and lysosomes restored catalytic activity. Engineering ligand monolayers on nanoparticles yielded a selective intracellular activation system (without tetra(ethylene glycol) (TEG) and an always-on system (with TEG).

Bioorthogonal activation of artificial metalloenzymes has also been adopted for recreating the reaction networks of signal processing between cells and organisms in protocells. For example, Walther and colleagues²⁹⁸ engineered all-DNA protocells hosting an artificial metalloenzyme catalyzing olefin metathesis, triggering downstream morphogenetic responses of varying complexity (Fig. 49B). The enzyme uncaged a pro-fluorescence signal, generating a self-reporting metabolite, weakening DNA duplex interactions and fostering growth, intraparticle adaptation, and interparticle fusion. These processes that mimic cellular adaptation and adhesion offered insight into life-like behavior *via* abiotic bioorthogonal chemical and mechanical transformations in synthetic protocells.

Late stage bioorthogonal diversification of amino acids and peptides has been employed for drug discovery and molecular bioimaging. In a research led by Ackermann,²⁹⁷ a modular linchpin strategy was engineered for stereo-divergent manganese(i)-catalyzed C–H labeling of complex peptides with bodipy probes (Fig. 49C). Validating manganese(i) catalysis,



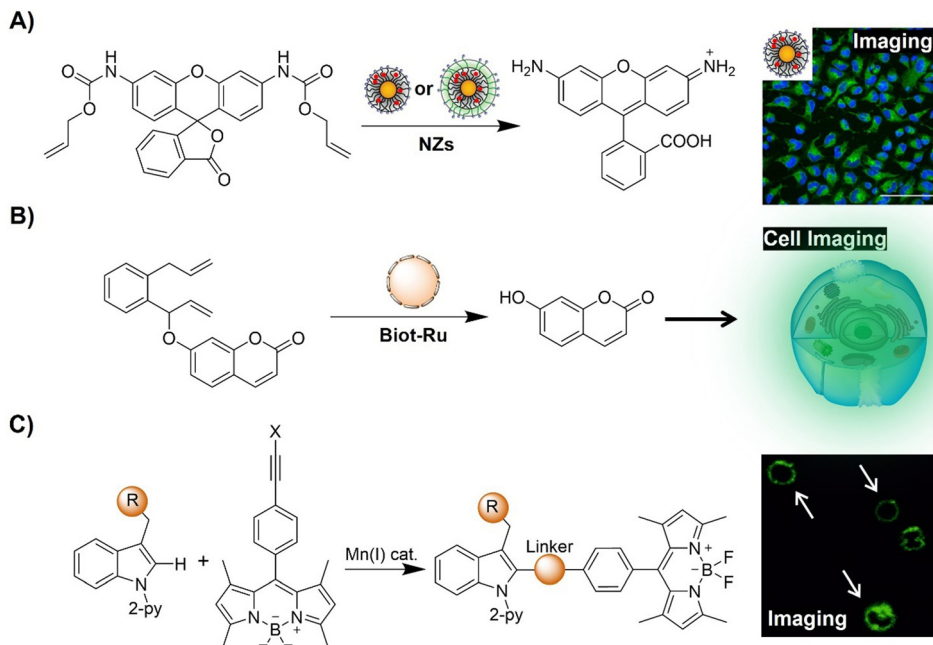


Fig. 49 Fluorescent probes for cell imaging developed using bioorthogonal catalyzed reactions. (A) AuNP nanozyme-catalyzed decaging of Pro-rhodamine and precise *in situ* HeLa cell imaging, adapted with permission from ref. 296. Copyright 2020, American Chemical Society; (B) schematic illustration of the Biot-Ru catalyzed activation of the coumarin fluorophore for precise *in situ* cell imaging; and (C) Mn(I) catalyzed C–H labeling of complex peptides with bodipy probes for immune cell membrane imaging, adapted from ref. 297, licensed under CC BY 4.0.

sensitive molecular rotors for monitoring immune cell membrane changes were obtained. A cholesterol-responsive probe emitting bright fluorescence was also developed, facilitating rapid screening for $CD8^+$ T cell modulators. This divergent C–H activation strategy promises efficient development of activatable fluorophores for real-time, physiological imaging of cell function.

Bioorthogonal transformation of imaging and therapeutic substrates *via* transition metal catalysts (TMCs) offers versatile biomedical applications. In research involving bioorthogonal transformations in complex biological environments, Sasmal and coworkers²⁹⁹ introduced an organoiridium complex to uncage allyloxycarbonyl-protected amines under biologically relevant conditions and within living cells (Fig. 50A). This uncaging chemistry demonstrated potential applications in diagnosis and treatment by controlled activation of the pro-fluorophore (Rho-alloc) and the prodrug (caged Dox) in HeLa cells. The fluorescence intensity increased up to 56-fold over 8 h, offering a valuable tool for diverse biological and therapeutic applications. In another work, Rotello *et al.*³⁰⁰ employed a polymeric scaffold to encapsulate transition metal catalysts (TMCs), yielding bioorthogonal “polyzymes” for activating intracellular anticancer therapeutics (Fig. 50B). In this “polyzyme” system, the hydrophobic interior of these polymers protected TMCs from deactivation in biological environments, without the clearance issues seen with inorganic nanoparticle analogues. Polyzymes³⁰¹ exhibited efficient intracellular activation of pro-dyes and prodrugs *in vitro*, indicative of their potential for disease diagnosis and treatment through on-site generation of imaging and therapeutic agents (Fig. 50C).

Precise localization of bioorthogonal catalysis is crucial to enhance therapeutic efficacy and minimize off-target effects. Towards this end, Rotello *et al.*³⁰² integrated biomimetic nanozymes with red blood cells (RBCs) as nanozymes for the selective targeting of bacterial infections (Fig. 51A). Utilizing non-immunogenic RBC-hitchhiked nanozymes, this approach selectively eliminated pathogenic bacterial infections. Toxins from bacteria induced hemolysis, leading to nanozyme accumulation and activation at sites of infection. These accumulated nanozymes subsequently released antibiotics, acting as *in situ* “dye and drug factories” for imaging and eradicating preformed biofilms without harming non-virulent bacteria. This approach exhibited potential for the treatment of recurrent bacterial infections, including chronic wounds and medical device-associated infections, offering targeted therapy without microbe-specific probes. By leveraging cellular interactions, this strategy encouraged diverse biomedical applications, providing a modular platform for imaging and therapeutics across various diseases.

Building on this research, Rotello *et al.*³⁰³ then devised a strategy utilizing surface-functionalized ZnS nanoparticles to achieve both enhanced bioorthogonal catalysis and biodegradability (Fig. 51B). Ligand-mediated acceleration of catalysis facilitated the uncaging of allyl-caged molecules by ruthenium catalysts, with released thiols acting as nucleophiles to accelerate the rate-determining step. Unlike enzyme-mimicking nanozymes such as FeS, ZnS-supported nanozymes underwent chemical transformations inaccessible to natural enzymes. This study highlighted the active involvement of nanoscaffolds as cofactors in bioorthogonal catalysis, surpassing non-degradable

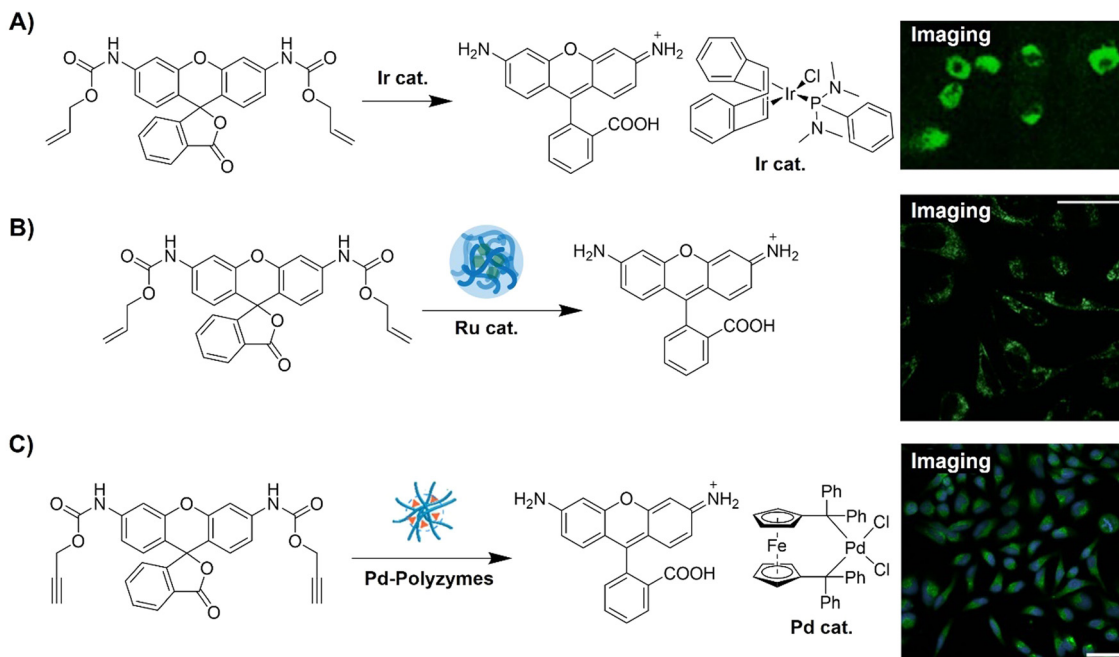


Fig. 50 Fluorescent probes for cell imaging developed using metal-based catalysts. (A) Ir complex catalyzed decaging of a profluorophore (Rho-alloc) and a prodrug (caged Dox) in HeLa cells, adapted with permission from ref. 299. Copyright 2021, American Chemical Society; (B) the Ru-polyzyme-catalyzed activation of Proc-rhodamine for precise *in situ* cell imaging, adapted with permission from ref. 300. Copyright 2020, Wiley-VCH GmbH; (C) the Pd-polyzyme-catalyzed activation of Proc-rhodamine for precise *in situ* HeLa cell imaging, adapted with permission from ref. 301. Copyright 2022, American Chemical Society.

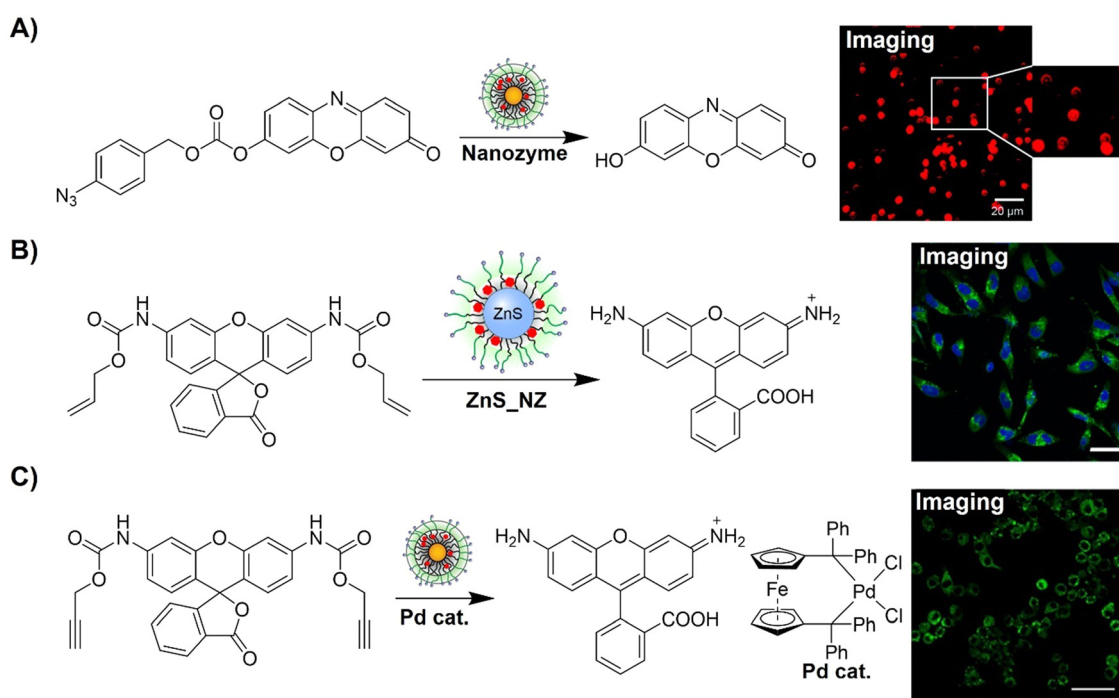


Fig. 51 Fluorescent probes for cell imaging developed using nanozyme-catalyzed activation. (A) Conversion of the profluorophore pro-Res through the nanozyme catalytic reduction of aryl azides for cell imaging, adapted with permission from ref. 302. Copyright 2021, The Royal Society of Chemistry; (B) the ZnS_NZ catalyzed activation of Proc-rhodamine for precise *in situ* cell imaging, adapted with permission from ref. 303. Copyright 2022, American Chemical Society; and (C) the Pd-nanozyme catalyzed activation of Proc-rhodamine for precise *in situ* cell imaging, adapted with permission from ref. 304, licensed under CC-BY-NC-ND 4.0.

gold nanoparticle nanozymes for fluorophore and drug release *in vitro*. ZnS nanozymes provided a biocompatible platform for *in situ* production of imaging and therapeutic agents, facilitating clinical translation of bioorthogonal catalysis and enabling localized therapies with minimal side effects and avoidance of particle accumulation.

Macrophage migration to tumor sites *via* chemoattractant gradients offers an active targeting strategy for cancer therapy. Taking this fact into consideration, Rotello *et al.*³⁰⁴ introduced bioorthogonal “nanozymes” with TMCs to transform profluorophores and prodrugs into *in situ* imaging agents and chemotherapeutics within macrophages, promoting macrophage-based drug delivery (Fig. 51C). These nanozymes, embedded in self-assembled monolayer-coated gold nanoparticles, solubilize and stabilize TMCs. Intracellularly localized nanozyme-loaded macrophages retained activity after prolonged incubation, maintaining migratory ability toward tumor chemoattractants and efficiently killing cancer cells. This macrophage-based delivery approach offered tunable dosages and delivery rates, minimizing off-target toxicity while activating prodrugs at tumor sites.

Understanding cellular functions requires detailed analysis of subcellular proteomes in precise spatial and temporal contexts. Chen *et al.*³⁰⁵ introduced CAT-Prox, a bioorthogonal and photocatalytic decaging-enabled proximity labelling strategy, for spatiotemporally resolved mitochondrial proteome profiling in living cells (Fig. 52A). In this bioorthogonal and photocatalytic system, Ir(ppy)₂bpy served as a bioorthogonal and mitochondria-targeting catalyst to control the release of azidobenzyl-caged quinone methide for protein labelling. CAT-Prox exhibited mitochondria specificity in live HeLa and RAW264.7 cells, enriching approximately 300 proteins with

70% specificity. For applications using lipopolysaccharide (LPS)-stimulated RAW264.7 macrophages, CAT-Prox dynamically dissected mitochondrial proteomes. By integrating photocatalytic decaging with proximity-based protein labelling, CAT-Prox exhibited excellent performance as a catalytic and non-genetic alternative for live-cell proteome analysis.

Tian and colleagues³⁰⁶ developed a versatile method for the *in situ* assembly of ICT-based light-up fluorophores using bioorthogonal Suzuki–Miyaura cross-coupling (Fig. 52B). This approach was applied to five fluorophore scaffolds, including naphthalimide, coumarin, naphthalene sulfonate, nitrobenzoxadiazole, and acetonaphthone, for the development of bioorthogonally activated probes for fluorescence imaging. The iodine group served as the bioorthogonal handle, quenching fluorescence by blocking ICT when strategically positioned. Upon Suzuki–Miyaura reaction, multicolor fluorophores were generated *in situ*, exhibiting significant fluorescence enhancement (up to 5277-fold) and high Φ_F (up to 0.82). Structure–property relationships of 1,8-naphthalimide probes were systematically explored, enabling fine-tuning of the photophysical properties. This bioorthogonal Suzuki–Miyaura cross-coupling was then applied for peptide modification, fluorogenic protein labeling, and live-cell mitochondria imaging with a high signal-to-noise ratio, representing a valuable expansion of the chemical biology toolbox for diverse biomedical applications.

In summary, bioorthogonal nanozymes represent a groundbreaking approach for biological investigations, enabling the localized synthesis of fluorescent probes and drugs at disease sites. This technique is particularly beneficial for precise *in situ* imaging and targeted drug delivery under challenging conditions such as tumors, bacterial infections, and during inflammation, where conventional treatments pose risks including

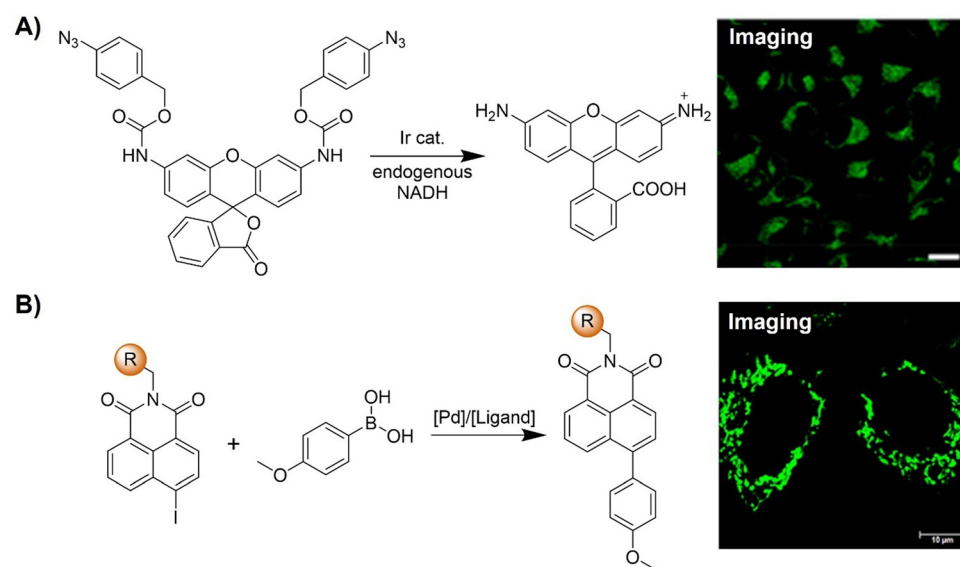


Fig. 52 Fluorescent probes developed for cell imaging using metal complex-based bioorthogonal activation. (A) Evaluation of Ir catalytic activity by PAB-rhodamine decaging in mitochondria in living cells through the CAT-Prox strategy, adapted with permission from ref. 305. Copyright 2021, American Chemical Society; (B) the Pd catalyzed fluorophore activation via Suzuki–Miyaura reaction for precise *in situ* cell imaging, adapted with permission from ref. 306. Copyright 2022, Elsevier B.V.



off-target effects. Furthermore, nanozymes offer extensive therapeutic potential due to their adaptability and versatile design possibilities. Manipulating various components like nanoparticles, ligands, and TMCs enhances adaptability and enables fine-tuning for future clinical applications.

Ensuring the long-term stability and efficacy of nanozymes is paramount, enabling high drug yields with minimal doses and adaptability to biological changes. Addressing potential cyto- and genotoxicity concerns arising from nanoscaffolds and TMCs is crucial for clinical translation, emphasizing the development of biodegradable scaffolds and non-toxic metal catalysts. Advances in organocatalysis may provide alternatives as well as low toxicity bioorthogonal TMCs based on metals such as silver and manganese. Enhancing the controllability of nanozymes can mitigate safety concerns, although reliable on/off switchable nanozymes have yet to be validated *in vivo*. Comprehensive studies on the distribution and metabolic pathways of nanozyme components in complex physiological environments are essential for clinical translation. Research efforts should concentrate on comprehensively understanding the complete lifecycle of nanozymes *in vivo* to effectively address safety concerns.

In brief, bioorthogonal nanozymes facilitate the creation of *in situ* probes and drug carriers, heralding a new era of enzyme-mimicking therapeutics. Their modular design holds promise for significant advancements in disease detection, deep tissue imaging, and precision therapy for biomedical applications.

3. Conclusion and outlook

Over the past two decades, we have witnessed a remarkable surge in the advancement of bioorthogonal chemistry, signifying a dynamic evolution that challenges longstanding assumptions in both biology and medicine. Particularly notable is the profound impact on fluorescence labeling, which has achieved a significant milestone through the integration of bioorthogonal ligation reactions with sophisticated molecular biological techniques. This symbiotic relationship allows for selective and biocompatible transformations, elevating our ability to scrutinize biological processes with unparalleled precision.

This review offers a comprehensive examination of diverse types of bioorthogonally activated *in situ* fluorescent labels. These probes, endowed with both bioorthogonal and fluorogenic properties, exhibit exceptional efficacy and precision in tagging specific biological targets with complementary chemical reporters. Despite the rapid expansion of the repertoire of bioorthogonal reactions, designing optimal bioorthogonal chemical probes tailored for specific biological systems remains a formidable challenge.

Primarily, there is an urgent need to devise strategies enabling diverse tagging, crucial for facilitating multicolor imaging. While certain orthogonal reactions, such as SPAAC-CuAAC, and specific combinations like the photoclick reaction with SPAAC, permit dual labeling and have achieved intricate multicolor imaging, further advancements in additional

orthogonal reactions are still required. Additionally, while current bioorthogonal reactions enable selective labeling of intracellular structures, their predominant application typically involves tagging extracellular targets such as receptors or glycans. To extend the utility of bioorthogonal tagging to intracellular and *in vivo* scenarios, there is a pressing need for the development of non-toxic, membrane-permeable labels applicable within bioorthogonal contexts. Furthermore, in addition to the continuous optimization of the performance of the existing systems, it is noteworthy that there are ample opportunities for developing novel types of bioorthogonal click reactions based on innovative reactant structures that allow for wash-free *in situ* precise imaging applications. We are confident that there are more unexplored avenues that need investigation. Through discovering new families of clickable molecules that are capable of meeting the increasingly diverse needs of *in vivo* applications, it is important to promote current bioorthogonal click chemistry for further advancements in this field.

In conclusion, bioorthogonal chemistry has revolutionized biological labeling and imaging, significantly broadening our horizons within complex physiological environments. The ongoing development of both existing and emerging bioorthogonally activated fluorescent probes holds immense promise for deepening our understanding of living systems. This heightened understanding facilitates improved comprehension of biomolecular processes and aids in the design of more effective drugs, particularly for targeted treatments such as those for tumors. With the relentless pace of innovations and developments, we firmly believe that the future of bioorthogonally activated probes for precise biolabeling and bioimaging is exceptionally promising.

Data availability

Data sharing is not applicable to this article as no new data were created or analyzed in this study.

Conflicts of interest

There are no conflicts to declare.

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