Recent Progress in Fluorescent Probes for the Detection and Research of Hydrogen Sulfide in Cells

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Abstract: Hydrogen sulfide (H2S) is a gaseous signaling molecule that plays an important role in regulating various physiological activities in biological systems. As the fundamental structural and functional unit of organisms, cells are closely related to the homeostasis of their internal environment. The levels of H2S in different organelles maintain a certain balance, and any disruption of this balance will lead to various functional abnormalities that affect the health of organisms. Fluorescent imaging technology provides unique merits, such as simplicity, non-invasiveness, and real-time monitoring, and has become a powerful approach for the detection of molecules in biological systems. Based on the special physicochemical properties of H2S, numerous H2S-specific fluorogenic probes have been designed with different recognition mechanisms that enable rapid and accurate detection of H2S in cells. Therefore, this review briefly illustrates the design strategies, response principles, and biological applications of H2S-specific fluorescent probes and aims to provide relevant researchers with insight for future research.

Keywords: hydrogen sulfide (H2S); fluorescent probe; cell imaging

1. Introduction

Hydrogen sulfide (H2S) is a gaseous signaling molecule that regulates various physiological activities in biological systems [1]. It is naturally produced by enzymes such as cystathionine γ-lyase (CSE), cystathionine β-synthetase (CBS), and 3-mercaptopyruvate sulfurtransferase (3-MST) that catalyze sulfur-containing substances such as cysteine (Cys) and homocysteine (Hcy) in living systems [2,3]. H2S content varies in different tissues and organs of organisms [4–6], which reflects the corresponding health status. As a mediator of the inflammatory response [7–9], H2S affects physiological activities and pathological processes in vivo. The disruption of H2S balance in internal circulation is an important predisposing factor for many diseases, such as Alzheimer’s disease [10], Down’s syndrome [11], Parkinson’s disease [12], asthma [13], and so on [14,15]. Therefore, the detection of endogenous H2S is critical for the study of biological and pathological processes in vivo.

A cell is the basic structure and functional unit of an organism. The biological activity of organisms is closely related to cellular homeostasis [16]. Notably, the vital movement of cells requires the joint operation of various organelles [17]. Studies have found that H2S not only plays an important role in various organelles [18–20], but also plays a key role in various cell signaling pathways [21,22]. Therefore, the development of simple, effective, and precise methods to detect H2S in biological systems is critical for better understanding its subcellular distribution and functions in various physiological and pathological processes.

The traditional methods for the detection of H2S mainly include gas chromatography [23,24], colorimetry [25], electrochemical methods [26], etc., but these methods inevitably show some shortcomings. The pre-treatment process of the above methods is too complicated, which hinders the application of real-time H2S monitoring in vivo.
In addition, the low selectivity and sensitivity to H$_2$S also limit their further application in vivo [27,28]. Therefore, in order to overcome the above problems, it is necessary to develop fast, efficient, and real-time H$_2$S monitoring technology to realize sensitive monitoring of H$_2$S in cells. In recent years, through the unremitting efforts of researchers, the development of fluorescence imaging technology has brought hope for the detection of molecules in biological systems. Its non-invasiveness and quick detection of biomarkers have been pursued by researchers [29]. Selective activatable probes can be designed to specifically detect target biomarkers, providing effective tools for the visualization of H$_2$S in cells and in vivo [30–32].

In recent years, the rapid development of fluorescence imaging technology has led to the emergence of a series of H$_2$S fluorescence probes with excellent properties, high efficiency, and accuracy, enabling real-time monitoring of H$_2$S [33]. Furthermore, many fluorescent probes have been developed for H$_2$S detection in different organelles, achieving systematic research on physiological and pathological processes at the subcellular level [34,35]. Although an abundance of literature briefly summarizes the design principles of H$_2$S probes, their applications at the cellular level have not been elaborated in detail [36,37]. Therefore, this short review aims to summarize novel H$_2$S probes with various design strategies and their applications in biological research at the cellular level, so as to provide readers with a deeper understanding of H$_2$S probe design strategies and to develop novel H$_2$S probes for further clinical applications (Scheme 1).

![Scheme 1. The mechanism and application of H$_2$S-activated fluorogenic probes.](image)

2. H$_2$S Probe

Organisms are complex systems where various molecules coexist and collaborate. Therefore, the detection of H$_2$S in vivo is susceptible to interference from other bioactive substances, such as persulfides (RSSH) [38] and reactive oxygen species (ROS) [39]. In view of the crucial role of H$_2$S in biological systems, a new type of H$_2$S-responsive fluorogenic probe with good biocompatibility, specificity, and optical properties is urgently needed to facilitate the detection of H$_2$S and further study of its physiological properties. H$_2$S has strong reduction ability and nucleophilicity, which are the main basis for designing H$_2$S response fluorogenic probes. Based on the activatable fluorogenic platform, a variety of fluorescent probes have been designed for detecting H$_2$S with different reaction strategies.

2.1. Probes Based on H$_2$S Reduction Reactions

2.1.1. Azide-Based H$_2$S Probes

As a bioorthogonal functional group, azide can be used to modify the chemical structure of various fluorescent dyes because of its convenient assembly. In addition, its efficient fluorescence quenching effect and good biocompatibility have been widely used.
Due to the reducibility of H$_2$S, the electron-withdrawing azide group will be converted to an electron-donating amino group by H$_2$S, restoring the D-π-A structure with high-emissive fluorescence and thereby achieving sensitive detection of H$_2$S. Notably, the selectivity of azide-based fluorogenic probes for H$_2$S is higher than that of reactive sulfur, oxygen, and nitrogen species. The chemical structures of azide-based H$_2$S probes introduced in this review are listed below (Figure 1).

In 2017, Wang et al. designed the two-photon Probe 1 [40] to visualize the intracellular and intercellular H$_2$S transfer processes. Probe 1 contains a H$_2$S-responsive azide group, a long-chain hydrophobic alkyl for anchoring the cell membrane, and a sulfonate to increase the hydrophilicity. Fluorescence imaging of endogenous H$_2$S in RAW 264.7 cells showed that fluorescence was mainly distributed in the plasma membrane rather than the cytoplasm, confirming its specific localization on the cytomembrane and the sensitive detection of intracellular H$_2$S release. In the same year, Lin et al. designed the azide-caged naphthalimide fluorescent Probe 2 [41] for the detection of mitochondrial H$_2$S (Figure 2A). When Probe 2 encounters high-level H$_2$S, the azide group transforms into an electron-donating amino group, which activates the probe with an intramolecular charge transfer (ICT) effect, emitting intense fluorescence at 540 nm (40 times). Due to its excellent responsiveness and the introduced triphenylphosphonium cations, Probe 2 shows mitochondrial distribution and is able to rapidly detect endogenous and exogenous H$_2$S, resulting in notable fluorescence imaging effects in cells (Figure 2B). In addition, Probe 2 can effectively detect H$_2$S in liver tissue using two-photon microscopy. Similarly, in order to improve the targeting of hepatocytes, Wang et al. introduced a galactose group into the quinoline fluorophore to obtain Probe 3 [42], which was effectively localized to hepatocytes through the specific recognition of the overexpressed asialoglycoprotein receptor (ASGPR). The low detection limit of fluorescent probes is crucial for the detection of trace substances and has significant impacts on cells and in vivo imaging. Probe 3 is an excellent probe for H$_2$S detection, with a reaction rate of only 1 min and a low detection limit of 126 nM in aqueous solution. Moreover, confocal imaging on different cell lines confirmed that Probe 3 can accurately and specifically detect H$_2$S in HepG-2 cells, making it a promising tool for detecting H$_2$S in cells and further investigating H$_2$S-related biological functions.
Wang et al. reported an anchored fluorescent Probe 4 [43] for selective imaging of H_2S with excellent properties. Based on this approach, Kim et al. reported two ratiometric therefore, the strategy of adding a self-immolative group between the fluorophore and probe was then further applied to investigate the role of CBS in the intracellular synthesis of H_2S by a gradual change in fluorescence from blue to yellow. Notably, Probe 7 showed two-photon Probes 6 and 7 [45]. After reacting with H_2S, the azide group was reduced to an amino group, and then the two-photon fluorophore was released, accompanied by a gradual change in fluorescence from blue to yellow. Notably, Probe 7 showed two-photon fluorescence and mitochondrial targeting properties, and can monitor the level of mitochondrial H_2S in living cells through changes in ratio fluorescence (F_{yellow} / F_{blue}). The probe was then further applied to investigate the role of CBS in the intracellular synthesis of H_2S (Figure 3), as well as pathological studies of Parkinson’s disease. Based on the same strategy, Liu et al. designed a ratiometric two-photon Probe 8 [46] to detect lysosomal H_2S. Due to its good biocompatibility and excellent two-photon properties, the probe enabled the visualization of both endogenous and exogenous H_2S in the lysosomes of living cells.

Protein labeling technology enables subcellular imaging of H_2S in different organelles. Wang et al. reported an anchored fluorescent Probe 4 [43] for selective imaging of H_2S in mitochondria and nuclei, respectively. Probe 4 includes the H_2S-activated fluorophore 7-azido coumarin (CouN_3) and additional O^2-benzylcytosine (BC) as a CLIP-label substrate. To verify the biological applicability of the technology, two plasmids, pCLIP-H2B and pCLIP-COX8A, that can instantaneously express CLIP fusion proteins were used to locate probes in the nucleus and mitochondria, respectively. Confocal fluorescence results indicated that the probe can effectively target the mitochondria and nucleus, respectively, and accurately detect changes in subcellular H_2S levels. By modifying the chemical structure of the probe, the optical properties of the probe can be effectively improved, which is conducive to the applications of probes in complex cellular environments. For this purpose, Lin et al. developed a series of H_2S probes based on rhodamine 110 dyes. In particular, Probe 5 [44] exhibits excellent optical performance and can visually detect H_2S generated by vascular endothelial growth factor (VEGF)-stimulated human umbilical vein endothelial cells (HUVECs). This finding confirms that H_2S production is dependent on VEGF receptor 2 (VEGFR2) and CSE.

In some cases, the direct addition of an azide group to a fluorophore is challenging. Therefore, the strategy of adding a self-immolative group between the fluorophore and the azide group has been proposed to facilitate the design and preparation of probes with excellent properties. Based on this approach, Kim et al. reported two ratiometric two-photon Probes 6 and 7 [45]. After reacting with H_2S, the azide group was reduced to an amino group, and then the two-photon fluorophore was released, accompanied by a gradual change in fluorescence from blue to yellow. Notably, Probe 7 showed two-photon fluorescence and mitochondrial targeting properties, and can monitor the level of mitochondrial H_2S in living cells through changes in ratio fluorescence (F_{yellow} / F_{blue}). The probe was then further applied to investigate the role of CBS in the intracellular synthesis of H_2S (Figure 3), as well as pathological studies of Parkinson’s disease. Based on the same strategy, Liu et al. designed a ratiometric two-photon Probe 8 [46] to detect lysosomal H_2S. Due to its good biocompatibility and excellent two-photon properties, the probe enabled the visualization of both endogenous and exogenous H_2S in the lysosomes of living cells.
parts: the introduced 4-azidobenzyl as the H\textsubscript{2}S complex probe photon the of excellent (VEGF)-stimulated probes mitochondria.fluorescent Probe 11 \cite{49} for the detection of H\textsubscript{2}S. The to an electron-donating amino group, the reductive product of the ICT process releases fluorescent Probe 11 \cite{49} for the detection of H\textsubscript{2}S. The combination of ratio fluorescence and NIR emission not only reduces background infection, but also reduces false results caused by photobleaching, enabling accurate detection of endogenous and exogenous H\textsubscript{2}S in cells. The chemical structures of nitro-/nitroso-based H\textsubscript{2}S probes introduced in this review are listed below (Figure 4).
Similarly, Hua et al. designed a H₂S-activated Probe 12 [50] employing N-annulated perylene and then introduced triphenylphosphonium salt or morpholine to design two organelle-targeting Probes 13 and 14 [50] (Figure 5A). All three probes can quantitatively detect low concentrations of H₂S in cells. Probes 13 and 14 exhibit excellent ratiometric fluorescence and biocompatibility, making them suitable tools for visual imaging of H₂S in mitochondria and lysosomes (Figure 5B). Aromatic nitroso compounds are reactive intermediates in biological reactions and are widely used in chemical and biological research. A flavylum derivative fluorescent Probe 15 [51] with nitroso as a recognition group was designed to detect H₂S. When the probe encounters H₂S, the sensitive fluorescence can be activated rapidly. With its good biocompatibility and optical properties, the probe can achieve fluorescence imaging of endogenous or exogenous H₂S in cells, making it an important tool for studying H₂S-related physiological processes.

Figure 4. Chemical structures of nitro-/nitroso-based Probes 11–15.

Figure 5. (A) Fluorescent Probes 12–14 for H₂S detection. (B) Confocal fluorescence imaging of subcellular H₂S in living cells with Probes 13 and 14. Reproduced with permission from Ref. [50]. Copyright 2017 The Royal Society of Chemistry.
2.2. Probes Based on H₂S Nucleophilic Reactions

2.2.1. 2,4-Dinitrophenyl (DNP)-Based H₂S Probe

In addition to reducibility, the nucleophilicity of H₂S is also an important property in the design of fluorescent probes. DNP is a potent electron-withdrawing group that can efficiently quench fluorescence by covalently linking to the modifiable phenolic group in the fluorophore. When the nucleophilic H₂S is added, the resulting mercaptan undergoes intramolecular nucleophilic substitution, and the probe releases free fluorophores with potent fluorescence. Therefore, DNP-based fluorescent probes are easy to prepare and have excellent quenching effects, so DNP is widely used in the design of various fluorescent probes. Yuan et al. designed two dual-excitation ratiometric fluorescent Probes 16 and 17 [52] based on coumarin and rhodamine. Those probes can eliminate the interference of microenvironments and improve the accuracy of fluorescence measurement. Fluorescence imaging of H₂S in living cells under different excitations showed excellent dual-excitation ratiometric imaging in mitochondria, which is expected to be applied to fluorescence imaging of intracellular H₂S in biological systems. The chemical structures of DNP-based H₂S probes introduced in this review are listed below (Figures 6 and 7).

![Chemical structures of DNP-based Probes 16–18.](image)

NIR fluorescent probes can overcome the interference of scattering and auto-fluorescence and are more conducive to sensitive imaging in cells. Therefore, Jiang et al. designed a NIR fluorescent Probe 18 [53] with a selective mitochondrial localization function for the detection of H₂S in cells. After reacting with H₂S, the probe emits significant fluorescence at 720 nm, which can quickly respond to H₂S in mitochondria. Then, Mito-Tracker rhodamine 123 was used for the co-localization experiment, and the results showed that Probe 18 had good mito-targeting. Cell imaging demonstrated that Probe 18 was suitable for the detection of endogenous and exogenous H₂S with significant fluorescence effects. In addition, the level of H₂S within the endoplasmic reticulum plays a crucial role in various biological and physiological processes. Li et al. developed three pyrimidine-based fluorescent probes for targeted detection of H₂S in the endoplasmic reticulum. The most suitable fluorophore L1 was selected by optical analysis, and Probe 19 [54] with good lipophilicity and a fast response to H₂S was obtained (Figure 7A). Lipophilicity enables the probe to penetrate cells and improve the detection efficacy, which facilitates the detection of H₂S in the endoplasmic reticulum (Figure 7B). Probe 19 can be utilized to detect H₂S in the cellular environment and also facilitate research on the physiological activities of H₂S within the endoplasmic reticulum (Figure 7C).
2,4-Dinitrobenzenesulfonyl (DNPS) is a H$_2$S-responsive quencher that can be connected to fluorescent dyes through DNP and sulfonic acid groups. The chemical structures of DNPS-based H$_2$S probes introduced in this review are listed below (Figures 8 and 9). Chen et al. designed three fluorescent Probes 20–22 [55] based on the fluorescein fluorophore and promoted the H$_2$S nucleophilic reaction by utilizing the positive influence of ortho aldehyde groups, so that DNPS could leave quickly, and the fluorescence of the probe could be restored. In addition, Probe 22 with two aldehyde groups can form thiazolidine enantiomers with biothiols, which can inhibit the fluorescence of fluorescein and improve the selectivity for H$_2$S (Figure 8A). Similarly, using DNPS groups, chemical probes with different performance can be designed to meet the monitoring needs of biological systems.

Therefore, Zhu et al. designed Probe 23 based on a curcumin core with a BF$_2$ group, which can distinguish between two DNPS groups as a regioselective fluorescence probe. The probe has two DNPS groups, and the one DNPS group is solely used to quench fluorescence, while the other DNPS reacts with H$_2$S to form a stable fluorescence intermediate. The selectivity of Probe 23 [56] for H$_2$S is more than 10,000 times higher than that of biothiols, and its excellent H$_2$S visualization ability is confirmed by cell fluorescence imaging. Xu et al. obtained NIR Probe 24 [57] by connecting hemicyanidin with benzothiazole groups, which can achieve the detection of H$_2$S content in cells (Figure 9). In another work, a H$_2$S-responsive NIR fluorescent Probe 25 [58] uses 5,15-di(4-chlorophenyl)-10-(4-hydroxylphenyl)-corrole as the fluorophore, which has excellent NIR fluorescence and low cytotoxicity and realizes the specific detection of H$_2$S in cells.
localization the quenching dual-channel cells. H₂S this levels.

Figure 8. (A) Schematic illustration and comparison of Probes 20–22 for H₂S detection. (B) Chemical structure of DNPS-based Probes 23–25.

Figure 9. (A) Fluorescent Probe 24 for H₂S detection. (B) Fluorescence imaging of endogenous and exogenous H₂S in cells. Reproduced with permission from Ref. [57]. Copyright 2021 Elsevier.

2.2.2. 7-Nitro-1,2,3-benzoxadiazole (NBD)-Based H₂S Probe

NBD is similar to H₂S-specific DNP, and its excellent characteristics can also be responsible for fluorescence quenching and H₂S-specific recognition. After incubating with H₂S, the nucleophilicity of H₂S causes the cleavage of NBD from the probe, releasing H₂S concentration-dependent fluorescence signals. NBD-based probes have the advantages of
fast reaction speed and good biocompatibility and are widely used in biological systems. The chemical structures of the NBD-based H$_2$S probes introduced in this review are listed below (Figure 10). Li et al. designed a coumarin-based Probe 26 [59] with an NBD group as the cage group and introduced triphenylphosphonium salt as a localization group for specific detection of mitochondrial H$_2$S. After reacting with H$_2$S, the fluorescence of the probe at 565 nm was decreased, while the fluorescence of coumarin at 415 nm was increased. Due to the self-calibration function of ratiometric fluorescence, the dual-channel $F_{415\text{ nm}}/F_{565\text{ nm}}$ signal enables more accurate and quantitative analysis of H$_2$S levels. Cell imaging confirmed that the probe could visualize H$_2$S in the mitochondria of cells. Similarly, Probe 27 [60] was designed using naphthalimide derivative as the fluorophore. Free Probe 27 did not show fluorescence due to the strong fluorescence quenching effect of NBD but showed fluorescence enhancement (68 times) at 528 nm after the addition of Na$_2$S. The probe has stable fluorescence characteristics and can be used for confocal microscopy imaging of intracellular H$_2$S, which proves that the probe can effectively detect exogenous H$_2$S in cells.

A NIR fluorescent Probe 28 [61] was designed using a long $\pi$-conjugated rhodamine dye and an NBD quenching group. When reacting with H$_2$S, the fluorescence intensity is increased 10 times at 660 nm, and the quantum yield of the redshift product reaches 0.29. The probe kinetic test showed that the reaction rate $k_2$ was 29.8 M$^{-1}$S$^{-1}$, and the detection limit was 0.27 $\mu$M, demonstrating that the probe had a rapid response to H$_2$S. The positive charge of the probe allows for its mitochondrial location, enabling fast and efficient NIR fluorescence imaging of H$_2$S at the subcellular level. Li et al. reported a fluorescent Probe 29 for simultaneous detection of H$^+$ and H$_2$S, and then introduced a morpholine to improve its lysosome location [62]. The fluorescence of the probe can only be turned on in the presence of H$^+$ and H$_2$S simultaneously. Probe 29 could detect H$_2$S within the pH range of the lysosome, thus realizing the accurate location of acidic lysosomes (Figure 11). Feng et al. improved the previous NBD-type probe (HBT-NBD) and obtained highly efficient and specific Probes 30 and 31 [63]. Probe 30 is not suitable for physiological applications due to its instability under alkaline conditions. In contrast, Probe 31 exhibits excellent stability at physiological pH. The incorporation of aldehyde and ortho-methyl groups introduces reversible reaction sites and steric hindrance effects, enhancing its selectivity for H$_2$S compared to other biothiols. Moreover, due to its low pKa value, H$_2$S is more likely to perform a nuclear reaction with a probe than other biothiols under physiological pH conditions. Consequently, Probe 31 is more suitable for monitoring H$_2$S levels in living cells.

The fluorescent probes based on DNP and NBD exhibit sensitive fluorescence changes upon H$_2$S nucleophilic attack, enabling effective detection and imaging of H$_2$S. However, the selectivity of these probes is hampered by interference from endogenous nucleophilic thiols in living cells. To address this challenge, the probes can be customized by implementing strategies such as spatial hindrance and auxiliary groups (such as ortho-benzaldehyde).

Figure 10. Chemical structures of NBD-based Probes 26–31.
2.2.3. Disulfide Bond-Based H2S Probe

H2S is nucleophilic and can also attack disulfide bonds. Therefore, disulfide bonds are introduced into the chemical structure of fluorescent dyes as H2S recognition sites. When H2S nucleophilic attacks and disulfide bonds break, mercaptan intermediates are formed, and then intramolecular cyclization occurs between the intermediates and esters, releasing fluorescence groups and restoring fluorescence signals. The chemical structures of disulfide bond-based H2S probes introduced in this review are listed below (Figure 12). Ye et al. designed two kinds of two-photon fluorescent probes, Probes 32 and 33 [64], based on naphthalimide fluorescent dyes and disulfide bond-responsive groups. They were linked to 2,2-disulfide benzoic acid by ester bonds and amides, respectively, and introduced morpholine groups into the target lysosomes. Probe 32 itself has only weak fluorescence. After the nucleophilic attack of H2S, the disulfide bonds break, and the fluorescent product is released, emitting intense fluorescence. However, Probe 33 is attached to disulfides by an amide bond, and after the disulfide bond breaks, the thiol intermediate cannot continue the cyclization process, which will quench the fluorescence of the probe. Both single photon and two-photon cell imaging experiments showed that Probe 32 could achieve H2S fluorescence imaging, which confirmed its application value in biological cells. Probe 34 [65] is a ratiometric fluorescent probe based on ICT and the excited intramolecular proton transfer (ESIPT) effect based on solvation effects. It has good photostability at physiological pH and can be used to achieve accurate and rapid quantitative detection of H2S by ratiometric fluorescence λ25 nm/λ495 nm. The probe can achieve significant fluorescence imaging ratios with different concentrations of H2S through different channels in cells. The importance of CBS and CSE for H2S production has been confirmed in endogenous H2S imaging experiments.

Figure 11. (A) Fluorescent Probe 29 for H2S detection under acidic condition. (B) fluorescence imaging of H2S in cells after different treatments. Reproduced with permission from Ref. [62]. Copyright 2018 Elsevier.

Figure 12. Chemical structures of disulfide bond-based Probes 32–38.
The nucleophilicity of H$_2$S allows it to activate disulfide-based fluorescent probes, but endogenous biothiols cannot be ignored as interfering species. The substitution of sulfur with selenium (S-Se-) can effectively enhance the selectivity of such probes for H$_2$S, enabling the convenient design of a wide range of probes suitable for H$_2$S detection. In 2017, Xian et al. designed a series of Washington Red (WR) NIR dyes with large Stokes shifts (>110 nm) and modified the hydroxyl group of the selected WR6 by disulfide bonding to obtain Probe 35 [66]. Compared with Probe 35, Probe 36 [66] uses Se to replace the sulfur atom in the disulfide bond, which improves anti-interference ability against biological mercaptans, enhances specificity against H$_2$S, and successfully realizes fluorescence imaging of H$_2$S in cells. Probe 36 overcomes the weak anti-interference ability of previous WSP and Sep series probes and allows for better adaptation to bioimaging (Figure 13). In the same year, Yin et al. prepared a water-soluble ratio fluorescence Probe 37 by introducing 2-(pyridin-2-yl disulfanyl) benzoic into 4-hydroxycoumarin [67]. In 2020, Probe 38 [68] was designed based on the structure of dicyanoisophorone. Both probes have outstanding fluorescence characteristics in vitro and good biocompatibility, enabling their application in the intracellular detection of H$_2$S.

![Figure 13. (A) The structures of NIR fluorescent Probes 35, 36. (B) Response mechanisms of WSP and Sep probes to H$_2$S. (C) Fluorescence imaging of endogenous H$_2$S in cells using Probe 36. Reproduced with permission from Ref. [66]. Copyright 2017 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.](image)

2.2.4. Other Nucleophilic-Based of H$_2$S Probes

Due to the nucleophilic nature of H$_2$S, various nucleophilic reactions can be carried out, such as nucleophilic substitution, nucleophilic addition, etc. Based on this strategy, probes with different structures can also be designed to meet the needs of various types of cell imaging. The chemical structures of other nucleophilic-based H$_2$S probes introduced in this review are listed below (Figure 14). Han et al. developed a cyanine dye-based turn-off fluorescent Probe 39 [69]. The strong fluorescence of the probe itself is turned off under the nucleophilic attack of H$_2$S. Thus, using fluorescence or UV–visible spectroscopy, the probe
can even detect H$_2$S in serum. Cell imaging experiments showed that the fluorescence of Probe 39 could be effectively turned off against H$_2$S, and NIR fluorescence emission characteristics reduced the signal interference in biological samples. Similarly, Probe 40 [70] also uses cyanine dyes as fluorophore and selects 2-carboxybenzaldehyde as the H$_2$S reaction site. Based on the tautomerism of ketones and enols in the nucleophilic addition product of Probe 40, a ratiometric fluorescence probe is designed to reduce environmental and instrumental interference. In the imaging of exogenous H$_2$S, the pixel intensity was only 0.0768. With the extension of reaction time with H$_2$S, the pixel intensity increased to 0.7091. In vitro imaging confirmed that Probe 40 could locate mitochondria and detect cellular H$_2$S in the NIR region.

Figure 14. Chemical structures of other nucleophilic-based Probes 39–42.

In 2020, Zhang et al. designed the NIR Probe 41 [71] based on O-carboxybenzaldehyde to realize the fluorescence detection of endogenous/exogenous H$_2$S in the endoplasmic reticulum. The excellent organelle localization of Probe 41 confirmed the important physiological role of H$_2$S in endoplasmic reticulum stress. Probe 42 [72] selected N-methyl-2-methoxyaniline moiety as the NO response site, while 4-nitrobenzenethiol-substituted boron dipyrromethene could respond to H$_2$S. NO and H$_2$S cycles can be dynamically visualized by the 645 nm channel (NO) and the 936 nm NIR II channel (H$_2$S). Imaging experiments on colonic smooth muscle cells and HepG-2 cells showed that Probe 42 can indeed dynamically monitor NO and H$_2$S within the cells, facilitating the study of the signal transduction between NO and H$_2$S (Figure 15).

Figure 15. Cont.
In addition to the mentioned H$_2$S fluorescent probes, it is important to consider additional recognition sites during probe design, while taking into account the specific chemical properties of the fluorophore itself. This approach allows for the development of more versatile H$_2$S fluorescent probes with improved sensitivity and specificity.

### 2.3. Probes Based on Metal and H$_2$S Reactions

Due to its special affinity for Cu$^{2+}$, H$_2$S can effectively combine with Cu$^{2+}$ to form CuS precipitates with $K_{sp}$ about $10^{-45}$ (25 °C, water) [36]. When H$_2$S and Cu$^{2+}$ react rapidly, Cu$^{2+}$, which quenches the fluorescence, immediately leaves, and the probe resumes the potent fluorescence. Fluorescent dyes designed using this mechanism allow for rapid detection of H$_2$S. The chemical structures of metal-based H$_2$S probes introduced in this review are listed below (Figure 16). In 2011, Sasakura et al. designed four fluorescent probes, Probes 43–46 [73], by incorporating fluorescein and a heterocyclic copper. Azazamacrocycles can form a stable metal complex with Cu$^{2+}$, which has an obvious quenching effect on the fluorophore. Probe 44 has the best optical properties and strong anti-interference ability, enabling selective fluorescence imaging of endogenous/exogenous H$_2$S within cells.

Yang et al. also designed three triarylboron-based Probes 47–49 [74] with different numbers of diphenylamine and cyclen, which can chelate Cu$^{2+}$ according to the affinity of cyclen to Cu$^{2+}$, thus inhibiting the ICT process and effectively quenching the probe fluorescence. The finite aggregates induced by probe and nonchromophoric analyte can effectively inhibit the aggregation-caused quenching (ACQ) effect and enhance the photostability of the probe. The selected Probe 48 has good membrane permeability and excellent two-photon properties, which can accurately image endogenous H$_2$S in mitochondria. Using the fluorescence lifetime microscopy (FLIM) technique, it can be clearly shown that H$_2$S is uniformly distributed within mitochondria. Similarly, Huang et al. designed two Cu(II)-cyclen-based BODIPY fluorescent Probes 50 and 51 [75] with fluorescence emission wavelengths of 765 and 680 nm, respectively. The detection limit of H$_2$S by Probe 50 is as low as 80 nM in vitro. In cell imaging, the probe showed excellent fluorescence signals in response to cellular H$_2$S.

Yuan et al. designed Probe 52 [76] based on the time-gated luminescence (TGL) technique of luminescent lanthanide complexes. Upon the reaction with H$_2$S, the chelated Cu$^{2+}$ leaves, and the fluorescence of Eu$^{3+}$ at 610 nm is significantly enhanced (62 times), while the fluorescence of Tb$^{3+}$ at 540 nm does not change significantly. A proportional TGL cell fluorescence imaging experiment showed that the addition of Cu$^{2+}$ can effectively quench fluorescence. In addition, the probe can be used for rapid and quantitative detection of...
intracellular H$_2$S, showing that TGL technology is suitable for fluorescence imaging of biological systems rich in biological background autofluorescence interference.

![Chemical structures of metal-based Probes 43–52.](image)

Figure 16. Chemical structures of metal-based Probes 43–52.

The metal-based probes exhibit rapid responses, but their biocompatibility is a significant concern. To mitigate these issues, the design of efficient sites capable of strong chelation with Cu$^{2+}$ should be prioritized, thereby preventing the binding of Cu$^{2+}$ with various cellular proteins and minimizing the potential toxicity of Cu$^{2+}$ to living cells. In addition, the impediment posed by the metabolic mechanisms of CuS precipitation in cells should be given careful consideration.

3. Summary and Outlook

As an important biological signaling molecule in biological systems, H$_2$S plays a unique role in various physiological and pathological activities, and the level of intracellular H$_2$S is closely related to the homeostasis of the biological environment. Therefore, it is extremely important to develop techniques that can detect H$_2$S in organisms, especially in cells that play various physiological functions. Fluorescence imaging probes have good biocompatibility and chemical stability, which is suitable for the application of biological systems. This review carefully analyzes fluorescent probes based on the physicochemical properties of H$_2$S and their applications in intracellular H$_2$S detection and H$_2$S-related biological processes, providing ideas and strategies for designing functional fluorescent probes for H$_2$S. However, there are still some challenges that hinder the development of H$_2$S-activated fluorescent probes, such as the following: (1) the resolution of most current cell imaging approaches is more than 200 nm, which makes it difficult to achieve accurate detection of intracellular molecular processes. Therefore, there is an urgent need to develop high-resolution cell imaging probes and techniques to obtain high-resolution information on intracellular H$_2$S to facilitate further studies of physiological mechanisms. (2) Fluorescent probes are usually easy to photobleach, making it difficult to meet the demand for long-term dynamic observation of H$_2$S. Therefore, developing fluorescent dyes with excellent light stability and strong anti-bleaching ability is necessary to achieve continuous analysis of intracellular H$_2$S fluctuation and H$_2$S-related pathways. (3) Fluorescent probes for H$_2$S detection should also be combined with other techniques to study physiological processes and gain a deeper understanding of the physiological significance of H$_2$S signal transduction. So far, researchers are continuing their exploration of the detection of cellular H$_2$S, hoping to develop more fluorescent probes with excellent characteristics suitable for cellular H$_2$S research in the future.
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