# Chemical Science

## EDGE ARTICLE



Cite this: Chem. Sci., 2022, 13, 2992

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 27th October 2021 Accepted 31st January 2022

DOI: 10.1039/d1sc05930k

rsc.li/chemical-science

### Introduction

With strong tissue penetration ability, minor cell and tissue damage, and low interference from autofluorescence, nearinfrared (NIR) fluorescence imaging is gradually replacing traditional fluorescence imaging as one of the most popular tools for real-time, *in situ*, visual tracking of biomolecules.<sup>1-5</sup> The recent success of *in situ* tracking of ferroptosis (an irondependent oxidative stress) based on NIR fluorescence sensing has provided a basis for the treatment and drug design for neurodegenerative diseases, acute kidney injuries, and



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Ferroptosis is closely associated with cancer, neurodegenerative diseases and ischemia-reperfusion injury and the detection of its pathological process is very important for early disease diagnosis. Fluorescence based sensing technologies have become excellent tools due to the real-time detection of cellular physiological or pathological processes. However, to date the detection of ferroptosis using reducing substances as markers has not been achieved since the reducing substances are not only present at extremely low concentrations during ferroptosis but also play a key role in the further development of ferroptosis. Significantly, sensors for reducing substances usually consume reducing substances, instigating a redox imbalance, which further aggravates the progression of ferroptosis. In this work, a  $H_2S$ triggered and H<sub>2</sub>S releasing near-infrared fluorescent probe (HL-H<sub>2</sub>S) was developed for the high-fidelity in situ imaging of ferroptosis. In the imaging process, HL-H<sub>2</sub>S consumes H<sub>2</sub>S and releases carbonyl sulfide, which is then catalyzed by carbonic anhydrase to produce H<sub>2</sub>S. Importantly, this strategy does not intensify ferroptosis since it avoids disruption of the redox homeostasis. Furthermore, using erastin as an inducer for ferroptosis, the observed trends for Fe<sup>2+</sup>, MDA, and GSH, indicate that the introduction of the HL-H<sub>2</sub>S probe does not exacerbate ferroptosis. In contrast, ferroptosis progression was significantly promoted when the release of H<sub>2</sub>S from HL-H<sub>2</sub>S was inhibited using AZ. These results indicate that the H<sub>2</sub>S triggered and H<sub>2</sub>S releasing fluorescent probe did not interfere with the progression of ferroptosis, thus enabling high-fidelity in situ imaging of ferroptosis.

> malignant tumors.6,7 However, the current NIR fluorescent probes used to monitor ferroptosis progression detect oxidizing substances such as lipid ROS and free ferrous ions, but the detection of reducing substances is rarely reported.6,8,9 Compared with oxidizing substances, reducing substances such as GSH and GPX4 are considered to be the key biomarkers to directly monitor ferroptosis, because ferroptosis can be regulated by the X<sub>c</sub><sup>-/</sup>GSH/GPX4 system by controlling the generation of phospholipid hydroperoxides.10 Current methods for the detection of antioxidants GSH or GPX4, for example western blotting make it impossible to achieve high sensitivity, in situ imaging and real-time tracking.11,12 As such the development of probes able to understand cell redox are required in order to understand cell ferroptosis, since the molecular regulation mechanisms of ferroptosis are complicated, and remain to be fully elucidated. In biology, the level of ferroptosis is determined by the simultaneous characterization of multiple targets.13-15

> As a gaseous signaling molecule, H<sub>2</sub>S plays an important role in human physiological and pathological processes.<sup>16,17</sup> H<sub>2</sub>S has been defined as an important endogenous neuroprotective agent that exerts protective effects through antioxidant, antiinflammatory, and anti-apoptotic mechanisms.<sup>18,19</sup> A



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 $<sup>\</sup>dagger$  Electronic supplementary information (ESI) available: Experimental details and characterization probe  $HL\text{-}H_2S$  supplementary spectral and cells experiments results. See DOI: 10.1039/d1sc05930k

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significant amount of evidence suggests that endogenous H<sub>2</sub>S is produced from cysteine desulfurization catalyzed by cystathionine  $\gamma$  lyase (CSE) or cystathionine  $\beta$  synthase (CBS).<sup>20,21</sup> Since ferroptosis inhibits the cystine/glutamate transporter (system x<sub>c</sub><sup>-</sup>), cysteine uptake is decreased, which leads to H<sub>2</sub>S depletion.<sup>22</sup> Thus, H<sub>2</sub>S can be considered as a representative reducing substance and as such enables monitoring of the ferroptosis process. However, it is difficult for current NIR fluorescent probes to facilitate *in situ* H<sub>2</sub>S tracking because extremely low concentrations of H<sub>2</sub>S will enhance the progression of ferroptosis.<sup>21,23</sup> Significantly, traditional sensors for reducing substances require the consumption of the reducing substance, leading to a redox imbalance, which further aggravates the progression of ferroptosis.<sup>24-26</sup>

Aiming to address these challenges, we have developed a benzyl thiocarbamate with an azide as the recognition site for H<sub>2</sub>S. Carbonyl sulfide (COS) is released through 1,6-elimination induced by H<sub>2</sub>S attack, which in turn releases H<sub>2</sub>S on catalysis by carbonic anhydrase (CA).<sup>27-30</sup> This H<sub>2</sub>S triggered and H<sub>2</sub>S releasing fluorescent probe has the potential to break through the current bottleneck and thus increase the accuracy in detecting ferroptosis. From our experimental results, the progression of erastin induced ferroptosis in cells with and without HL-H<sub>2</sub>S were not significantly different. In contrast, the progression of ferroptosis was significantly promoted when the H<sub>2</sub>S release from HL-H<sub>2</sub>S was inhibited using AZ. Our results indicate that HL-H<sub>2</sub>S with both H<sub>2</sub>S triggered and H<sub>2</sub>S releasing mechanisms does not induced ferroptosis. Therefore, our probe is capable of *in situ* high-fidelity ferroptosis analysis, making it a reliable tool for the comprehensive and accurate understanding of ferroptosis progression.

#### Results and discussion

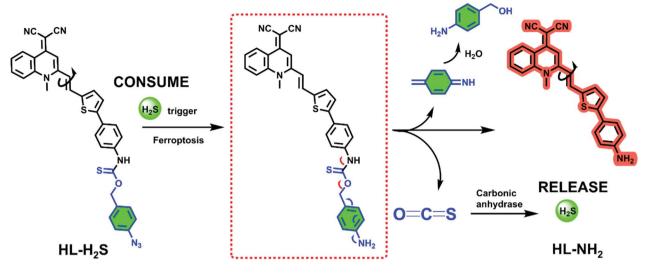
#### Design strategy and synthesis of fluorescent probe HL-H<sub>2</sub>S

The response mechanism of the H<sub>2</sub>S triggered and H<sub>2</sub>S releasing probe HL-H<sub>2</sub>S are shown in Scheme 1. With NIR emission, HL- $H_2S$  is endowed with good tissue penetration ability, which results in minor cell and tissue damage, and low interference from autofluorescence.31 An azido benzene was used as the H2S recognition site<sup>32,33</sup> which was linked to the fluorophore using a thiocarbamate (H<sub>2</sub>S precursor). When HL-H<sub>2</sub>S responds to H<sub>2</sub>S, COS is released through 1,6-elimination, which is then catalyzed by CA to release H<sub>2</sub>S. Thus, a novel H<sub>2</sub>S triggered and H<sub>2</sub>S releasing system results. Most notably, quinolinemalonitrile is linked to the thiophene in HL-NH<sub>2</sub> by  $\sigma$ -bonds, so molecular rotation of its structure will be regulated by viscosity. This enabled us to accurately detect the ferroptosis process, because cell viscosity increases during ferroptosis.6,8,34 HL-H2S could be used to detect the ferroptosis of cells of high viscosity, but there was no significant change in normal cells.<sup>6,35</sup> In a high-viscosity environment, the rotation of HL-NH2 is hindered, resulting in an increase of quantum yield from 0.01 to 0.67, as shown in Table S1.<sup>†</sup> As such the sensitivity of HL-NH<sub>2</sub>, is enhanced in a viscous environment facilitating accurate detection of ferroptosis. The detailed synthetic procedures and characterization data for probe HL-H<sub>2</sub>S are described in Scheme S1.<sup>†</sup> The response mechanism

of the probe was confirmed by monitoring the <sup>1</sup>H NMR and verifying the molecular weight before and after reaction using HR-MS (Fig. S1 and S2†). When **HL-H<sub>2</sub>S** reacts with NaHS (H<sub>2</sub>S donor), the signal at  $\delta$  11.4 (H<sub>a</sub>) disappears, and a molecular ion peak *m*/*z* 429.11169 using HR-MS confirms the formation of **HL-NH<sub>2</sub>** (*m*/*z* 429.11444, [M + Na]<sup>+</sup>).

#### Photophysical properties of HL-NH<sub>2</sub>

With probe HL-H<sub>2</sub>S in hand, the photophysical properties of the fluorophore HL-NH<sub>2</sub> in different solvents were systematically investigated to optimize the detection system. Fig. S3a and b<sup>†</sup> show the absorption and fluorescence spectra of HL-NH<sub>2</sub>, respectively, and the photophysical parameters are given in Table S1.<sup>†</sup> In glycerol, the fluorescence intensity of HL-NH<sub>2</sub> was the highest at 646 nm (quantum yield = 0.67), the molar absorptivity reached  $0.64 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$  ( $\lambda_{abs} = 430 \text{ nm}$ ), and the Stokes shift was 216 nm. Therefore, light scattering due to excitation was avoided, and the resolution and accuracy of the probe improved. According to Fig. S3d,† the fluorescence intensity of HL-NH<sub>2</sub> was highest in glycerol, indicating that the fluorescence intensity of HL-NH<sub>2</sub> is affected by viscosity. The sensitivity of HL-NH2 to polarity was also analyzed. As shown in Fig. S3c,† the fluorescence intensity of HL-NH2 does not change as the dielectric constant of the solvent increases, indicating that fluorescence emission of HL-NH<sub>2</sub> is not affected by the polarity of the solvent. According to the literature,<sup>36,37</sup> quinolinemalonitrile derivates exhibit aggregation-induced emission (AIE). The AIE performance of HL-NH2 was investigated in a mixture of tetrahydrofuran (THF) and water, where THF was a good solvent for HL-NH<sub>2</sub> and water was a poor solvent. As shown in Fig. S4a,<sup>†</sup> the absorption intensity of HL-NH<sub>2</sub> remains low without significant changes as the water content  $(f_w)$ decreased from 100% to 70%. As  $f_w$  drops below 70%, the absorption intensity of HL-NH2 increases abruptly at 450 nm, which may be due to the scattering effect of the aggregates generated in situ.38 Unlike absorption intensity, the fluorescence intensity of HL-NH<sub>2</sub> was negligible at 0% and 100%  $f_w$ , and reaches a maximum at 40% (Fig. S4b-d†), which is different from conventional AIE luminescence.39 We believe that the reason for the difference observed for our system and traditional AIE fluorophores based on quinolinemalonitrile are due to the primary amino group of HL-NH<sub>2</sub> exhibiting enhanced solubility is PBS through hydrogen bonding with water, which results in reduced AIE.37 The response of HL-NH2 to viscosity was also evaluated. As shown in Fig. S5a-c,† the fluorescence intensity of HL-NH2 gradually increases as the viscosity of the solution is increased from 1.1 cP (0 vol% glycerol) to 1410 cP (99 vol% glycerol) by increasing the proportion of glycerol. According to Fig. S5b and c,† two different linear relationships emerge for volume fraction ranges from 0-40% and 50-99%. For each viscosity range the linear correlation coefficient between the fluorescence intensity of the probe and the viscosity are 0.996, indicating that the probe HL-H2S can quantify the viscosity of a target solution. Finally, the aqueous solubility of the probe HL-H2S was investigated. The probe was soluble in a water solution at a concentration of 20 µM and



Scheme 1 H<sub>2</sub>S triggered and H<sub>2</sub>S releasing probe for high-fidelity ferroptosis evaluation and mechanistic details for COS/H<sub>2</sub>S release by HL-H<sub>2</sub>S.

exhibited a good linear relationship between solubility and concentration ( $R^2 = 0.974$ ). Therefore, the solubility of **HL-H<sub>2</sub>S** is sufficient for *in vivo* and *in vitro* experiments (Fig. S6†).

#### Spectroscopic response of HL-H<sub>2</sub>S to H<sub>2</sub>S

The response of the probe  $HL-H_2S$  to  $H_2S$  (from NaHS) was investigated in phosphate buffer solution (PBS, pH = 7.4,

containing 80% glycerol and 2% DMSO). The addition of 100  $\mu$ M H<sub>2</sub>S resulted in a significant increase in the spectral intensity of both the UV and fluorescence (Fig. 1a and b), indicating excellent response of **HL-H<sub>2</sub>S** towards H<sub>2</sub>S. The fluorescence titration experiments (Fig. 1c) indicated that as the H<sub>2</sub>S concentration increased from 0 to 500  $\mu$ M, the fluorescence intensity of the solution ( $\lambda_{em} = 670$  nm) increased by around 25-fold before reaching a plateau at a H<sub>2</sub>S concentration of 100  $\mu$ M

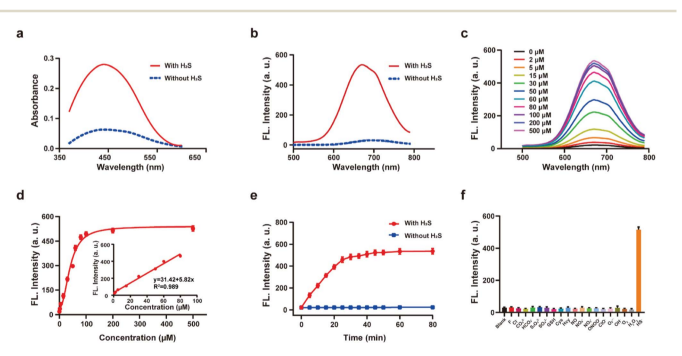


Fig. 1 Absorption spectra (a) and fluorescence spectra (b) of probe HL-H<sub>2</sub>S with H<sub>2</sub>S (from NaHS) concentration (100  $\mu$ M) with H<sub>2</sub>S (solid line) and without H<sub>2</sub>S (dotted line). (c) Fluorescence spectra of HL-H<sub>2</sub>S with the increasing H<sub>2</sub>S concentration (0–500  $\mu$ M). (d) Calibration curve for the determination of H<sub>2</sub>S (inset linear response at lower H<sub>2</sub>S concentrations). (e) Time dependent fluorescence intensity of HL-H<sub>2</sub>S in the absence and presence of H<sub>2</sub>S (100  $\mu$ M). (f) Response of HL-H<sub>2</sub>S to 200  $\mu$ M different interferents (HL-H<sub>2</sub>S alone (blank)); F<sup>-</sup>, Cl<sup>-</sup>, CO<sub>3</sub><sup>-2</sup>, HCO<sub>3</sub><sup>-2</sup>, SO<sub>4</sub><sup>2-</sup>, GSH, Cys, Hcy, NO, NO<sub>3</sub><sup>-</sup>, NO<sub>2</sub><sup>-</sup>, ONOO<sup>-</sup>, ClO<sup>-</sup>, O<sub>2</sub><sup>--</sup>, 'OH, <sup>1</sup>O<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>, HS<sup>-</sup>, respectively. For all experiments the concentration of HL-H<sub>2</sub>S was 10  $\mu$ M.  $\lambda_{ex}$  = 450 nm, in PBS (pH = 7.4, containing 2% DMSO and 80% glycerol). In (d–f), data represent the mean of three replicates and the error bars indicate the SD.

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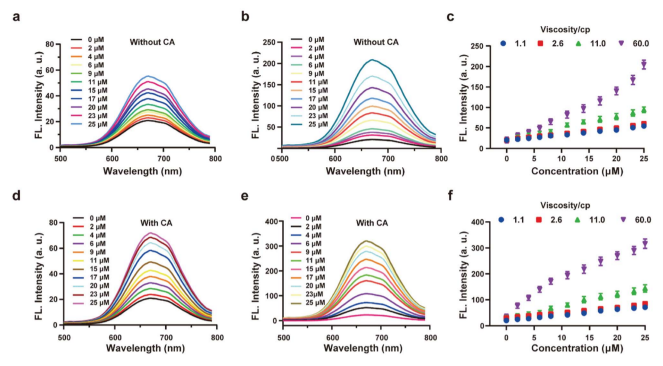


Fig. 2 Probe HL-H<sub>2</sub>S (10  $\mu$ M) with increasing H<sub>2</sub>S concentrations (0–25  $\mu$ M) in PBS (pH = 7.4, containing 2% DMSO). Fluorescence spectra of HL-H<sub>2</sub>S (10  $\mu$ M) in the absence of carbonic anhydrases (CA) with the increasing H<sub>2</sub>S concentration (0–25  $\mu$ M) in the absence (a) and presence (b) of glycerol (80% in volume, viscosity: 60.0 cP). (c) Dose-dependent titrations between HL-H<sub>2</sub>S and H<sub>2</sub>S concentrations (0–25  $\mu$ M) with different glycerol level (1.1, 2.6, 11.0 and 66.0 cP). Fluorescence spectra of HL-H<sub>2</sub>S (10  $\mu$ M) in the presence of carbonic anhydrases (CA) with the increasing H<sub>2</sub>S concentration (0–25  $\mu$ M) with different glycerol level (1.1, 2.6, 11.0 and 66.0 cP). Fluorescence spectra of HL-H<sub>2</sub>S (10  $\mu$ M) in the presence of carbonic anhydrases (CA) with the increasing H<sub>2</sub>S concentration (0–25  $\mu$ M) in the absence (d) and presence (e) of glycerol (80% in volume, viscosity: 60.0 cP). (f) Dose-dependent titrations between HL-H<sub>2</sub>S and H<sub>2</sub>S concentration (0, 2, 4, 6, 9, 11, 15, 17, 20, 23 and 25  $\mu$ M);  $\lambda_{ex} = 450$  nm. In c and f, data represent the mean of three replicates and the error bars indicate the SD.

(Fig. 1d). While the fluorescence intensity exhibited a good linear relationship with concentrations of H<sub>2</sub>S over a range from 1-80 µM (Fig. 1d inset). The detection limit was calculated to be 1.3 nM. The above experiments indicated that HL-H<sub>2</sub>S exhibits a dual response capability for viscosity and H<sub>2</sub>S. The response of HL-H<sub>2</sub>S was then analyzed in terms of kinetics, pH stability, and selectivity to further investigate the performance towards H<sub>2</sub>S. Kinetic analysis indicated that the reaction between HL-H2S and H2S (100 µM) was completed within 40 min, indicating that HL-H<sub>2</sub>S is capable of quickly identifying H<sub>2</sub>S (Fig. 1e). The effect of different pH environments on HL-H<sub>2</sub>S and its response to H<sub>2</sub>S were then evaluated. As shown in Fig. S7,<sup>†</sup> over a pH range from 5.0 to 8.5, the fluorescence intensity of HL-H<sub>2</sub>S does not change significantly regardless of the presence of H<sub>2</sub>S, indicating that the response of HL-H<sub>2</sub>S to H<sub>2</sub>S is not affected by changes in environmental pH. Finally, the influence of reactive oxygen (ClO<sup>-</sup>, O<sub>2</sub><sup>•-</sup>, <sup>•</sup>OH, <sup>1</sup>O<sub>2</sub>, and H<sub>2</sub>O<sub>2</sub>), reactive nitrogen (NO<sub>2</sub><sup>-</sup>, NO<sub>3</sub><sup>-</sup>, NO, and ONOO<sup>-</sup>), reactive sulfur (GSH, Cys, Hcy, S<sub>2</sub>O<sub>3</sub><sup>2-</sup>, SO<sub>4</sub><sup>2-</sup>and HS<sup>-</sup>) and other anions (F<sup>-</sup>, Cl<sup>-</sup>, CO<sub>3</sub><sup>2-</sup>, HCO<sub>3</sub><sup>-</sup>) on the response of HL-H<sub>2</sub>S were evaluated. As shown in Fig. 1f, these interfering species hardly affect the fluorescence intensity of the system, while the introduction of  $H_2S$  significantly enhanced the fluorescence intensity of the system, indicating that HL-H<sub>2</sub>S exhibits a high selectivity for H<sub>2</sub>S.

#### In vitro H<sub>2</sub>S release of HL-H<sub>2</sub>S

The above experiments confirmed that the probe  $HL-H_2S$  is capable of H<sub>2</sub>S-specific recognition in a high-viscosity environment. Next, ferroptosis was simulated in vitro and the detection and release capacity of the probe for H<sub>2</sub>S was evaluated to facilitate the accurate detection of ferroptosis using the probe. CA has been reported to catalyze the conversion of COS, to produce H<sub>2</sub>S.<sup>27,28</sup> As shown by Fig. 2a, in the absence of CA and glycerol, a ca. ~3-fold enhancement of the fluorescence with increasing H<sub>2</sub>S concentration was achieved. However, the fluorescence increases 10-fold as the glycerol content increases to 80% (viscosity: 60 cP), and the effect of environmental viscosity on the response of HL-H<sub>2</sub>S to H<sub>2</sub>S is significant (Fig. 2b and c). Compared to the absence of CA, the sensitivity of HL-H<sub>2</sub>S to H<sub>2</sub>S is slightly enhanced in the presence of CA (Fig. 2d). Importantly, Fig. 2e and f show that the fluorescence intensity increases 15fold with addition of H<sub>2</sub>S, which was attributed to the release of H<sub>2</sub>S from HL-H<sub>2</sub>S following reaction with H<sub>2</sub>S. To verify this conjecture, acetazolamide (AZ), an inhibitor of CA,40 was used to regulate the release of  $H_2S$ . Fig. S8 a and b<sup>+</sup> shows that the fluorescence intensity of the solution with AZ and CA is almost identical to that of the solution without AZ or CA, and both are lower than that of the solution with CA, indicating that under these conditions HL-H<sub>2</sub>S can be catalyzed by CA to produce H<sub>2</sub>S. To verify the conversion of decomposed HL-H<sub>2</sub>S to H<sub>2</sub>S in the

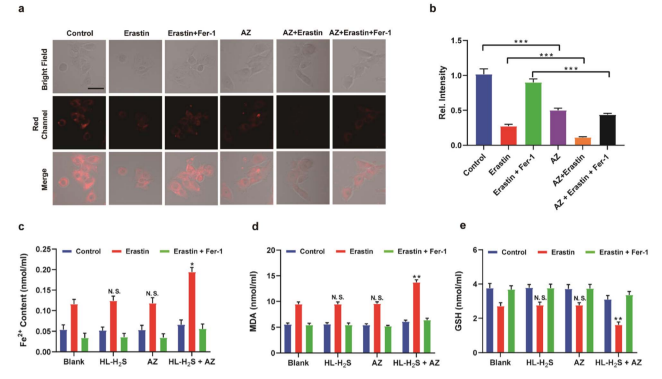


Fig. 3 Confocal images of  $H_2S$  in ferroptosis. (a) PC12 cells under different conditions: control group (only  $HL-H_2S$ , 10  $\mu$ M);  $HL-H_2S$  (10  $\mu$ M) + erastin (10  $\mu$ M);  $HL-H_2S$  (10  $\mu$ M) + Fer-1 (5  $\mu$ M). (b) Histograms of average fluorescent intensities in (a) Difference was analyzed by one-way ANOVA. \*\*\*p < 0.001.  $\lambda_{ex} = 450$  nm,  $\lambda_{em} = 630-710$  nm. Scale bars: 40  $\mu$ m. (c) PC12 cells under different conditions: control group (HL-H\_2S (10  $\mu$ M) + acetazolamide (AZ, 50  $\mu$ M)); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M); HL-H\_2S (10  $\mu$ M) + AZ (50  $\mu$ M) + erastin (10  $\mu$ M) + Fer-1 (5  $\mu$ M). (d) Histograms of average fluorescent intensities in (c); (e) the content of Fe<sup>2+</sup>. (f) The content of MDA. (g) The content of GSH. For the fluorescence change: data are presented as the mean  $\pm$  SD (control: n = 44 cells from three cultures; HL-H\_2S + erastin: n = 43 cells from three cultures; HL-H\_2S + erastin + Fer-1: n = 33 cells from three cultures; HL-H\_2S + AZ + erastin: n = 35 cells from three cultures; HL-H\_2S + AZ + erastin + Fer-1: n = 33 cells from three cultures). \*p < 0.05, \*\*P < 0.01 vs. blank. In (b) and (d-g), the error bars indicate the SD.

presence of CA, the well-known methylene blue (MB) method<sup>29</sup> was used to measure the generation of  $H_2S$  under the above conditions. The results indicated that the changes of  $H_2S$  monitored using MB were consistent with the fluorescence results obtained for the probe, indicating that under these conditions **HL-H\_2S** can be catalyzed by CA to produce  $H_2S$  (Fig. S8c†), which confirms that **HL-H\_2S** can release  $H_2S$  in the presence of CA. Fig. S8d† is the calibration curve between concentrations of  $H_2S$  (0–20  $\mu$ M) and absorption intensity at 670 nm.

# Imaging of $\mathrm{H_2S}$ and cytoplasmic viscosity in living cells with HL-H\_2S

After confirming the  $H_2S$  response and  $H_2S$  release capacity of  $HL-H_2S$  for *in vitro* experiments, we anticipated that  $HL-H_2S$  can keep the level of cell ferroptosis form being aggravated during ferroptosis monitoring, which excludes the possibility of  $HL-H_2S$ -induced ferroptosis, thus enabling high-fidelity imaging and analysis of ferroptosis. Before performing cellular confocal imaging, the biocompatibility of  $HL-H_2S$  was evaluated. As shown in Fig. S9,† the survival rate of PC12 cells is over 90% after 24 h coincubation with  $HL-H_2S$  at different concentrations (0–30  $\mu$ M) according to MTT assay, indicating that  $HL-H_2S$  has

good biocompatibility and is suitable for in situ imaging analysis. The dual response of HL-H<sub>2</sub>S to H<sub>2</sub>S and viscosity was then evaluated at the cellular level. Intracellular H<sub>2</sub>S content and viscosity levels were regulated using NaHS (10 µM) to release  $H_2S$ , viscosity inducer monensin<sup>41</sup> (10  $\mu$ M), and different cell incubation temperatures (25 °C and 4 °C) since lowering the temperature increases the intracellular viscosity.42,43 As shown in Fig. S10a and b,<sup>†</sup> the red channel fluorescence intensity of the groups with NaHS or monensin were slightly higher than those of the control group. However, the red channel fluorescence intensity of the group with NaHS and monensin was significantly higher than that of the first three groups. In addition, as the incubation temperature decreased, the red channel fluorescence intensity in the group with NaHS increased significantly, while that of the group without NaHS remained unchanged (Fig. S10c and d†). The above experiments indicate that HL-H<sub>2</sub>S has dual response to viscosity and H<sub>2</sub>S at the cellular level. The ability of HL-H2S to detect exogenous and endogenous H<sub>2</sub>S was then assessed. As shown in Fig. S11,<sup>†</sup> PC12 cells were incubated with HL-H<sub>2</sub>S (10  $\mu$ M) for 10 min to ensure complete entry into the cells. Then the cells were incubated for 40 min with monensin (10 µM), sulfhydryl scavenger N-ethylmaleimide44 (NEM, 0.5 mM), and different concentrations of

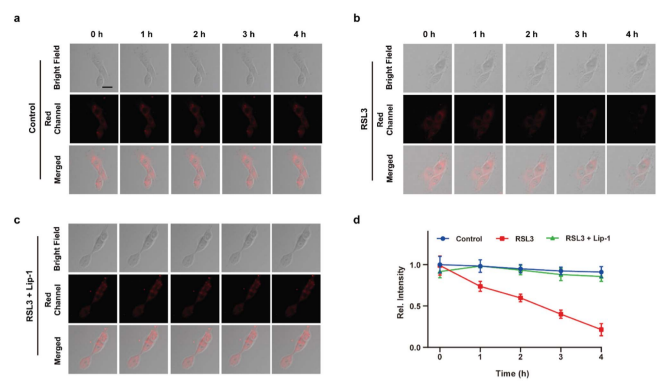


Fig. 4 Confocal images of PC12 cells for: (a) control (only HL-H<sub>2</sub>S), (b) RSL3 (1.0  $\mu$ M, ferroptosis inducer), and (c) RSL3 + Lip-1 (0.5  $\mu$ M, ferroptosis inhibitor), respectively. Cells were stained with HL-H<sub>2</sub>S (10.0  $\mu$ M) for 30 min before imaging. (d) Histograms of average fluorescence intensity of HL-H<sub>2</sub>S in control (blue line), RSL3 (red line) and RSL3 + Lip-1 (green line) at different times (0, 1, 2, 3, 4 h). For the fluorescence change: control group: n = 43 cells from three cultures; RSL3 group: n = 41 cells from three cultures; RSL3 + Lip-1 group: n = 51 cells from three cultures.  $\lambda_{ex} = 450$  nm,  $\lambda_{em} = 630-710$  nm. Scale bars: 20  $\mu$ m. In (d), data represent the mean of three replicates and the error bars indicate the SD.

NaHS (H<sub>2</sub>S donor) (0, 5, 10, or 20  $\mu$ M). The experimental results indicated that the red channel fluorescence intensity gradually increased with an increase of H<sub>2</sub>S concentration (Fig. S11a and b<sup> $\dagger$ </sup>), indicating that HL-H<sub>2</sub>S could detect exogenous H<sub>2</sub>S. It should be noted that the fluorescence intensity of the groups with NEM dropped significantly compared to the control group, which was due to the removal of endogenous H<sub>2</sub>S by NEM. Therefore, HL-H<sub>2</sub>S could detect endogenous H<sub>2</sub>S. CA activity was modulated using AZ to illustrate the H<sub>2</sub>S releasing capacity of the probe. According to the results shown in Fig. S12,† the red channel fluorescence intensity of the group with AZ decreases significantly compared to the groups without AZ. Meanwhile, since exogenous H<sub>2</sub>S is present in cells, regardless of whether the cells were co-incubated with AZ, the result is consistent with Fig. S8.† However, the fluorescence intensity difference is negligible between the groups with and without NEM. The above experiments show that HL-H<sub>2</sub>S is capable of H<sub>2</sub>S response and release at the cellular level.

#### Imaging of ferroptosis during OGD/R

Inspired by the above experiments, the high-fidelity response of the H<sub>2</sub>S triggered and H<sub>2</sub>S releasing probe **HL-H<sub>2</sub>S** to ferroptosis was evaluated. As shown in Fig. 3a and b, a ferroptosis model was built by incubating PC12 cells with the ferroptosis inducer erastin,<sup>45</sup> in which the red channel fluorescence intensity decreases significant compared to the control group, indicating that endogenous H<sub>2</sub>S is consumed in the ferroptosis process. Meanwhile, the addition of ferroptosis inhibitor ferrostatin-1 (Fer-1)45 resulted in a significant increase of red channel fluorescence intensity, indicating that HL-H<sub>2</sub>S is capable of the in situ imaging of ferroptosis. Given that Fe<sup>2+</sup>, MDA, and GSH levels are common indicators of ferroptosis progression, the simultaneous increase in Fe<sup>2+</sup> and MDA and decrease in GSH indicate that a ferroptosis model was successfully constructed<sup>46,47</sup> (Fig. 3c-e, blank group). Importantly, the Fe<sup>2+</sup>, MDA, and GSH levels were not significantly different between cells with and without HL-H2S (As shown in Fig. 3c-e, blank group vs. HL-H<sub>2</sub>S group). In addition, regardless of whether the cells undergo ferroptosis, the red channel fluorescence intensity was significantly decreased in cells where H<sub>2</sub>S release was inhibited from HL-H<sub>2</sub>S by AZ (Fig. 3a and b). While Fe<sup>2+</sup> and MDA levels increased and the GSH level decreased significantly compared to the groups without HL-H<sub>2</sub>S (as shown in Fig. 3c-e, blank group vs.  $HL-H_2S + AZ$  group). Moreover, there is no change of Fe, MDA, or GSH content for the AZ only control experiment compared to the control or probe alone, further indicating that AZ has no effect on the degree of cell ferroptosis, and only inhibits the ability of the probe to release H<sub>2</sub>S by inhibiting the activity of CA (as shown in Fig. 3c-e, blank group vs. AZ group). These results indicated that the H<sub>2</sub>S triggered and H<sub>2</sub>S releasing fluorescent probe HL-H<sub>2</sub>S maintained the progression of ferroptosis, thus enabling high-fidelity in situ

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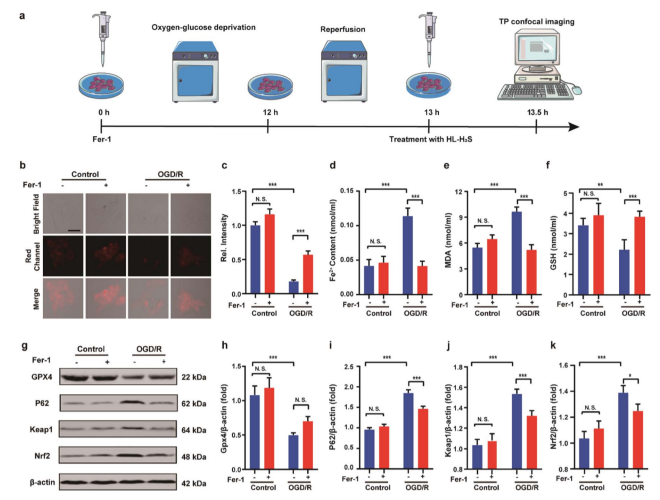
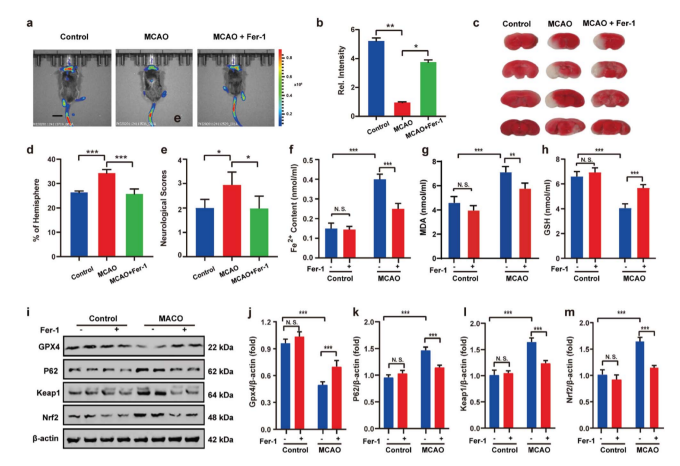


Fig. 5 Confocal images of H<sub>2</sub>S in OGD/R. (a) Schematic illustration of oxygen glucose deprivation/reoxygenation (OGD/R) modeling method and probe treatment. (b) PC12 cells using OGD/R models with untreated cells, Fer-1 treated cells, OGD/R treated cells and Fer-1 treated cells during OGD/R. (c) Histograms of average fluorescent intensities in (b); for the fluorescence change: for untreated cells: n = 48 cells from three cultures; for Fer-1 treated cells: n = 44 cells from three cultures; for Fer-1 treated cells: n = 51 cells from three cultures. (d) The content of Fe<sup>2+</sup>; (e) the content of MDA; (f) the content of GSH. (g-k) Western blotting analysis of GPX4, p62, Keap1, and Nrf2 in OGD/R model pre-treated with untreated cells, Fer-1 treated cells, OGD/R treated cells and Fer-1 treated cells and Fer-1 treated cells and Fer-1 treated cells during OGD/R. Difference was analyzed by one-way ANOVA. \*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001.  $\lambda_{ex} = 450$  nm,  $\lambda_{em} = 630-710$  nm. Scale bars: 40 µm. In (c)–(f) and (h)–(k), the error bars indicate the SD.

imaging of ferroptosis. To further evaluate the capacity of HL-H<sub>2</sub>S for real time monitoring of ferroptosis, confocal imaging of PC12 cells stained with HL-H<sub>2</sub>S were recorded at different times (0, 1, 2, 3, 4 h) under different cell culture conditions including control, RSL3, and RSL3 + liproxstatin-1 (Lip-1). As shown in Fig. 4, the fluorescence of cells with RSL3 (ferroptosis inducer) decreased gradually with time. However, the fluorescence for control cells (only HL-H<sub>2</sub>S) and those pre-treated with Lip-1 (ferroptosis inhibitor<sup>48</sup>) remained constant. Thus, fluorescence intensity change of HL-H<sub>2</sub>S could be used to monitor RSL3induced ferroptosis in live cells.

Finally, PC12 cells were used to construct an oxygen glucose deprivation/re-oxygenation (OGD/R) model and simulate cells undergoing ischemia-reperfusion,<sup>49</sup> with which the relationship between cellular OGD/R and ferroptosis were explored (Fig. 5a). According to Fig. 5b and c, the red channel fluorescence intensity in the OGD/R group decreases significantly compared

to the control group (p < 0.001). However, the fluorescence intensity of the group with Fer-1 is again significantly higher compared to the group without Fer-1 (p < 0.001). Therefore, the cellular ischemia-reperfusion process was accompanied by ferroptosis. It should be noted that the red channel fluorescence intensity of the control group with the addition of Fer-1 is slightly increased compared to the group without Fer-1, indicating the presence of ferroptosis among normal cells. The trends of Fe<sup>2+</sup>, MDA, and GSH levels are comparable with those observed for the fluorescence intensity (Fig. 5d-f), indicating the capacity of the probe to monitor ferroptosis progression induced by OGD/R. The p62-Keap1-Nrf2 signaling pathway was examined to further explore the pathological mechanisms by which the cellular ischemia-reperfusion process induces ferroptosis.47,50 Western blotting assay shown in Fig. 5g-k indicated that the p62-Keap1-Nrf2 signaling pathway was activated in the OGD/R cells, while GPX4 expression was significantly



**Fig. 6** Visual imaging MCAO in living mice model. (a) Mice using MCAO models with different treatments: control group (mice not undergoing MCAO); MCAO group (mice undergoing MCAO); Fer-1 group (injection of Fer-1 to mice tail veins). Scale bar = 2 cm. (b) Histograms of average fluorescent intensities in (a); (c) infarct size in the ipsilateral hemisphere was measured 3 days after stroke onset using TTC staining, then normalized to the contralateral hemisphere and expressed as a percentage. (d) Bar graphs represent the statistic results. (e) Neurological score test; the content of (f) Fe<sup>2+</sup>, (g) MDA and (h) GSH in living mice MCAO model under different conditions in (a). (i–m) Western blotting analysis of GPX4, p62, Keap1, and Nrf2 in living mice MCAO model under different conditions: control group (mice not undergoing MCAO); MCAO group (mice undergoing MCAO); Fer-1 group (injection of Fer-1 to mice tail veins). In (b), (d)–(h) and (j)–(m), the error bars indicate the SD. Difference was analyzed by one-way ANOVA. \*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001.  $\lambda_{ex} = 450$  nm,  $\lambda_{em} = 630-710$  nm.

downregulated. After incubation with Fer-1, the p62–Keap1– Nrf2 signaling pathway was suppressed and GPX4 was upregulated. The p62–Keap1–Nrf2 signaling pathway and the GPX4 expression in the control group indicated no significant changes with or without the addition of Fer-1. Therefore, the ischemia-reperfusion process inhibited GPX4 and in turn affected the p62–Keap1–Nrf2 signaling pathway to induce ferroptosis.

#### In vivo imaging of cerebral apoplexy in mice

After uncovering the relationship between ischemiareperfusion and ferroptosis, the cerebral apoplexy in mice was simulated using a middle cerebral artery occlusion (MCAO) model. Prior to imaging, the biocompatibility of **HL-** $H_2S$  in each tissue was assessed. After intravenous injection of **HL-H**<sub>2</sub>S, the different organs, including brain, heart, liver, spleen, lung, and kidney were stained by hematoxylin and eosin (H&E). As shown in Fig. S13,† **HL-H**<sub>2</sub>S exerted no obvious organ or tissue damage, indicating its good tissue

biocompatibility for in vivo imaging analysis in mice. The findings acquired from the mouse MCAO model were like those obtained from the cellular model. The fluorescence intensity in the brains of mice in the MCAO model group decreased significantly compared to that of normal mice (p <0.01). However, adding ferroptosis inhibitor Fer-1 caused a significant rebound of fluorescence intensity in the brains of mice (Fig. 6a and b, p < 0.05), indicating that MCAO in mice could lead to ferroptosis in their brains. 2,3,5-Triphenyltetrazolium chloride (TTC, measures tissue viability used to evaluate infarct size) staining and behavioral experiment results are in high agreement with in vivo imaging, indicating the viability of the mouse MCAO model (Fig. 6c-e). The changes in Fe<sup>2+</sup>, MDA, and GSH levels again demonstrate that ferroptosis occurred in the brain of mice during stroke (Fig. 6f-h). Similarly, the relationship between ferroptosis and GPX4 activity and the p62–Keap1–Nrf2 signaling pathway was explored in mice, and the results were consistent with those at the cellular level. Therefore, the MCAO process inhibits GPX4

activity in mice, which in turn affects the p62-Keap1-Nrf2 signaling pathway and ultimately leads to ferroptosis (Fig. 6i-m).

### Conclusions

In summary, a H<sub>2</sub>S triggered and H<sub>2</sub>S releasing NIR fluorescent probe HL-H<sub>2</sub>S was developed, which exhibits dual response to viscosity and H<sub>2</sub>S. After reacting with H<sub>2</sub>S, the probe releases COS through 1,6-elimination, which in turn releases H<sub>2</sub>S catalyzed by CA. In vitro experiments confirm a low detection limit of 1.3 nM, significant fluorescence enhancement and good selectivity amongst various ROS/RNS species. In addition, the probe exhibited excellent characteristics in terms of large Stokes shift (216 nm), favourable water solubility, excellent pH stability, and low cytotoxicity. While the cell experiments indicated that the progression of erastin induced ferroptosis in cells with and without HL-H<sub>2</sub>S were not significantly different. In contrast, the progression of ferroptosis was significantly promoted when the H<sub>2</sub>S release from HL-H<sub>2</sub>S was inhibited using AZ. Significantly, by using H<sub>2</sub>S triggered and H<sub>2</sub>S releasing mechanisms during the imaging process, the probe could avoid the induction of ferroptosis. Therefore, the probe is capable of in situ high-fidelity ferroptosis analysis, making it a reliable tool for the comprehensive and accurate understanding of ferroptosis progression. A cellular OGD/R model was constructed, and a mouse MCAO model was built to simulate MCAO in mice, and ferroptosis inducer erastin and ferroptosis inhibitor Fer-1 were used to regulate the progression of ferroptosis during MCAO. Confocal imaging revealed that the MCAO process could induce ferroptosis.

### Data availability

All data supporting this study are provided as ESI<sup>†</sup> accompanying this paper.

### Author contributions

Tianyu Liang: methodology, investigation, data curation, validation, formal analysis, writing – original draft. Taotao Qiang: project administration, writing – review & editing. Longfang Ren: data curation, investigation. Fei Cheng: validation. Baoshuai Wang: data curation, formal analysis. Mingli Li: software, data curation. Wei Hu: investigation, data curation, formal analysis, writing – original draft. Tony D. James: conceptualization, writing – review & editing, supervision.

### Conflicts of interest

TDJ acts as an academic consultant for TQ as part of a guest professorship at SUST.

### Acknowledgements

All animal procedures were performed in accordance with the Guidelines for Care and Use of Laboratory Animals of South-Central University of Nationalities and experiments were approved by the Animal Ethics Committee of College of Biology (South-Central University of Nationalities). This work was funded by the Open Project of Shaanxi Collaborative Innovation Center of Industrial Auxiliary Chemistry & Technology (No. XTKF-2020-05); the Key Scientific Research Group of Shaanxi Province (2020TD-009), Key Scientific Research Program of Shaanxi Provincial Education Department (Collaborative Innovation Center project) (20JY003), Science and Technology Plan Project of Xi'an Weiyang District (No. 201907) and the Youth Innovation Team of Shaanxi Universities. TDJ wishes to thank the Royal Society for a Wolfson Research Merit Award and the Open Research Fund of the School of Chemistry and Chemical Engineering, Henan Normal University for support (2020ZD01).

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