A highly selective and sensitive near-infrared fluorescent probe for imaging of hydrogen sulphide in living cells and mice

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Hydrogen sulphide (H₂S), the third endogenous gaseous signalling molecule, has attracted attention in biochemical research. The selective detection of H₂S in living systems is essential for studying its functions. Fluorescence detection methods have become useful tools to explore the physiological roles of H₂S because of their real-time and non-destructive characteristics. Herein we report a near-infrared fluorescent probe, NIR-HS, capable of tracking H₂S in living organisms. With high sensitivity, good selectivity and low cytotoxicity, NIR-HS was able to recognize both the exogenous and endogenous H₂S in living cells. More importantly, it realized the visualization of endogenous H₂S generated in cells overexpressing cystathionine β-synthase (CBS), one of the enzymes responsible for producing endogenous H₂S. The probe was also successfully applied to detect both the exogenous and endogenous H₂S in living mice. The superior sensing properties of the probe render it a valuable research tool in the H₂S-related medical research.

Hydrogen sulphide (H₂S) has emerged as the third endogenous gaseous signalling molecule in living organisms, along with nitric oxide (NO) and carbon monoxide (CO)¹⁻². It is mainly produced by enzymes such as cystathionine β-synthetase (CBS), cystathionine γ-lyase (CSE) and cysteine aminotransferase (CAT)/3-mercaptopyruvate sulphurtransferase (3-MST)³. Physiological levels of H₂S have diverse biological functions, including neurotransmission⁴, vasodilation⁵, apoptosis⁶, inflammation⁷, ischemia/reperfusion-induced injury⁸ and insulin secretion⁹. In addition, abnormal concentrations of H₂S appear to be involved in many diseases¹⁰,¹¹, such as Alzheimer’s disease and diabetes mellitus.

With increasing interest in understanding the chemical and biological properties of H₂S, sensitive and selective detection techniques for monitoring endogenous H₂S are urgently desirable, since the complex manifestations of H₂S in both physiological and pathological states, as well as its underlying molecular events are still not fully understood. The current approaches for H₂S detection, such as the methylene blue method, the monobromobimane (MBB) method, gas chromatography (GC), and the sulphide ion selective electrodes (ISE) method are not suitable for in situ analysis¹². Fluorescence-based assays, however, could offer convenience, high sensitivity, nondestructiveness, as well as real-time imaging¹³.

Recently, various sensing strategies have been focused on the design of H₂S-reactive probes, including nucleophilic addition¹⁴⁻¹⁹, copper sulphide precipitation²¹⁻²³, H₂S-mediated reduction²⁴⁻²⁹, and the thiolysis of dinitrophe nyl ether by H₂S³⁰,³¹, etc.³². However, most of these fluorescent probes are based on fluorophores with peaks in the ultraviolet-visible (UV/V is) region, which renders them difficult to be employed for imaging H₂S in living animals due to high absorption and autofluorescence of biomolecules.³³ By contrast, probes with absorption and emission in the near-infrared (NIR) region are more desirable for in vivo imaging because of minimal photo damage, deep
was lower than most of the reported NIR H₂S probes. Therefore, NIR-HS is highly sensitive to low-nanomolar
levels of sulphide, which facilitate the quantitative detection of endogenous/intracellular H₂S in complex biolog-
cal systems.

Results and Discussion

Synthesis and sensing mechanism of NIR-HS. In the design of a NIR probe for H₂S, the hemicya-
nine skeleton (a NIR dye) was selected as a fluorophore in the light of its NIR emission and high stability46. It is
known that the thiolysis of the dinitrophenyl ether reaction can be chemoselective for H₂S over biothiols38.
Thus, probe NIR-HS was constructed by connecting a dinitrophenyl group to hemicyanine (a NIR dye) via an
ether-linkage (Fig. 1). The fluorescence of NIR-HS was quenched due to alkylation on the hydroxyl group46. We
speculated that the reaction of sulphide with NIR-HS would cleave an ether group, and release the free fluoro-
phore, thereby achieving fluorescence detection of sulphide. On the basis of this design, the structure of NIR-HS
and the proposed sensing mechanism are illustrated in Fig. 1. The probe was readily synthesised in two steps.

Treatment of IR-780 with resorcin in the presence of K₂CO₃ afforded compound 1, which was then condensed
and the proposed sensing mechanism are illustrated in Fig. 1. The probe was readily synthesised in two steps.

Fluorescent properties of NIR-HS. The fluorescent properties of NIR-HS (10 μM) in the absence and
presence of Na₂S were determined. The free probe was almost nonfluorescent (Fig. 2A). However, treatment of
Na₂S (100 μM) led to a large fluorescence enhancement at 723 nm (50 fold, Φ = 0.13). Figure 2B depicted elevated
fluorescence intensities with increasing amounts of Na₂S (0–300 μM) until a plateau reached at 100 μM Na₂S. An
excellent linear correlation between the observed fluorescence intensities and various concentrations of Na₂S
(0–100 μM) was observed in PBS buffer (Fig. 2B inset). The in vitro detection limit for sulphide was 38 nM, which
was lower than most of the reported NIR H₂S probes. Therefore, NIR-HS is highly sensitive to low-nanomolar
levels of sulphide, which facilitate the quantitative detection of endogenous/intracellular H₂S in complex biological
systems.

The fluorescence intensity in reaction of NIR-HS with Na₂S reached the maximum value within approximately
20 min (Supplementary Fig. S2). The effects of pH on the detection of sulphide were then evaluated (Supplementary
Fig. S3). In the pH range from 5.8 to 6.0, the emission intensities were quite low and did not change significantly.
From pH 6.2 to 6.8, the fluorescence intensities were gradually increased, and the maximal fluorescence intensities
were observed from pH 7.0 to 9.0. The emission profile of fluorophore (compound 1) (Supplementary Fig. S4) are
consistent with the results of treating the probe with Na₂S in different pH PBS buffer, indicating that the observed
pH profile is due to the fluorophore itself. Taken together, NIR-HS is suitable for the detection of sulphide between
pH 7.0 and 9.0.

Selectivity to sulphide of NIR-HS. To investigate the selectivity of NIR-HS towards sulphide, NIR-HS was
treated with various species. NIR-HS displayed high selectivity for H₂S over physiological concentrations of

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Figure 1. Design and synthesis of NIR fluorescent turn-on probe NIR-HS.
biological thiols (Fig. 3A and Supplementary Fig. S5), including glutathione (10 mM GSH), cysteine (1 mM Cys) and homocysteine (1 mM Hcy). The good selectivity of NIR-HS attribute to the stronger nucleophilic properties of HS\(^{-}\), smaller size and a lower pKa at neutral pH compared to other thiols. The remaining non-thiol amino acids (Ala, Glu, Trp, Met, Tyr, Leu, Val, Ser, Pro, Arg, Gly, Phe, His, Gin, Asn, Ile and Thr), reactive oxygen species (H\(_2\)O\(_2\), ·OCl\(^{-}\), O\(_2\)\(^{-}\), ·OH and tBuOOH), reactive nitrogen species (NO\(_2\)\(^{-}\) and NO), sulphur-containing inorganic ions (S\(_2\)O\(_3\)\(^{2-}\), S\(_2\)O\(_5\)\(^{2-}\), SO\(_4\)\(^{2-}\), S\(_2\)O\(_4\)\(^{2-}\), SO\(_3\)\(^{2-}\) and SCN\(^{-}\)), S-nitroso glutathione (SNG), reducing agents (NADH and glucose) and inorganic salts (KCl, CaCl\(_2\), NaCl, MgCl\(_2\), FeCl\(_3\), ZnSO\(_4\) and NaH\(_2\)PO\(_4\)) induced negligible responses (Fig. 3B and Supplementary Fig. S6). Additionally, competitive experiments also revealed hardly any interference to sulphide detection in the coexistence of Na\(_2\)S and various species (Supplementary Fig. S6 and Fig. S7). The fluorescence intensity decreased in the presence of H\(_2\)O\(_2\) and ZnSO\(_4\) due to the oxidation of H\(_2\)S by H\(_2\)O\(_2\) and sulphide precipitation of H\(_2\)S by ZnSO\(_4\). Thus, NIR-HS can be used for the selective detection of sulphide with minimum interference from other biological species.

Detection of H\(_2\)S in living cells. We thereafter assessed the potential utility of NIR-HS to monitor H\(_2\)S in living MCF-7 cells. Prior to cell imaging, MTT assays were performed to evaluate the cytotoxicity of the probe. NIR-HS and compound 1 exhibited IC\(_{50}\) of 96.9 ± 3.2 μM and 99.4 ± 1.7 μM, respectively (Supplementary Figs S8 and S9). These results indicated that cells were variable after incubation with NIR-HS (5 μM) for 24 h.
MCF-7 cells incubated with the free probe (5 μM) showed relatively weak fluorescence emission (Fig. 4, panel 1A). In contrast, upon addition of Na2S (50 μM) to the above cells, a strong red fluorescence was observed (Fig. 4, panel 1B). The sulphide-induced increase of the fluorescence intensity in cells of panel 1B was finished after approximately 20 min (Fig. 4, panel 1C). Subsequently, cells were pretreated with ZnCl2 (an efficient eliminator of H2S). With the addition of Na2S (50 μM) to the ZnCl2-pretreated cells, no fluorescence intensity increases were observed (Fig. 4, panel 2A). The results indicated that the fluorescence change of NIR-HS in the cells arises from H2S. Moreover, the treatment of probe-loaded cells with Na2S (25 μM) yielded lower fluorescence emission (Fig. 4, panel 2B) compared to the fluorescence of the cells in panel 1C, implying that NIR-HS is capable of imaging different sulphide concentrations in living cells (Fig. 4, panel 2C).

Next, we tested the abilities of NIR-HS to visualize the endogenous H2S. MCF-7 cells express H2S-producing enzyme such as CSE47. NO could upregulate the CSE expression and stimulate the CSE activity, resulting in increased endogenous H2S level48. Therefore, SNP (Sodium Nitroprusside, a NO donor) was employed to induce the production of endogenous H2S in MCF-7 cells. The probe-loaded cells exhibited faint fluorescence emission without the addition of SNP (Fig. 5, panel 1A). After incubation of probe-treated cells with SNP (Fig. 5, panel 1B) for another 20 min, the fluorescence signal increased significantly, indicating the generation of endogenous H2S within the cells. Whereas the cells preincubated with DL-propargylglycine (PPG, an inhibitor for CSE48) provided almost no fluorescence enhancement (Fig. 5, panel 1C), demonstrating that the fluorescence change is triggered by endogenously generated H2S.

CSE and CBS are major enzymes for H2S production, and the overexpression of CBS or CSE could result in the elevation of endogenous H2S level1,2. We thus constructed the cells with CBS overexpression (Fig. 5, panel 2A). Cells transfected with empty vector (pCMV6) were set as control group (Fig. 5, panel 2B). As shown in Fig. 5, MCF-7 cells that were overexpressing CBS showed much stronger fluorescence (Fig. 5, panel 2A) than that from cells of the control group (Fig. 5, panel 2B), suggesting the increased endogenous level of H2S in CBS overexpressed cells. The western blot assay proved the overexpression of CBS in cells of panel 2A (Supplementary Fig. S12). We also quantified the fluorescence intensities of these cells, and found that the CBS overexpressed cells showed 2-fold enhanced fluorescence intensity compared to the control cells (Fig. 5, panel 2C). These results revealed the capability of NIR-HS to recognize endogenous H2S in living cells.

Detection of H2S in living mice. The prominent NIR features of NIR-HS render the probe highly favorable for fluorescence imaging of H2S in living animals. Inspired by these data, we further examined the suitability of the sensor to visualize exogenous and endogenous H2S in living mice. Kunming mice were divided into several groups. The mice were given i.p. injection of DMSO as the negative control group (Supplementary Fig. S13, panel

Figure 4. Confocal fluorescence imaging of exogenous sulphide in living MCF-7 cells using NIR-HS. Cells were incubated with NIR-HS (5 μM) alone for 10 min (1A). Cells in panel 1A were thereafter treated with Na2S (50 μM) for 10 min (1B) and 20 min (1C). Cells were pretreated with 1 mM ZnCl2, then incubated with NIR-HS (5 μM) for 10 min and Na2S (50 μM) for 20 min (2A). Cells were incubated with NIR-HS (5 μM) for 10 min and further incubated with Na2S (25 μM) for additional 20 min (2B). Scale bars = 10 μm. The average fluorescence intensity of the above images (2C). Data are presented as the mean ± SD (n = 3). *p < 0.001 vs. (1A) column, **p < 0.001 vs. (1C) column.
A), and the mice were given i.p. injection of free probe as the probe-loaded group (Fig. 6, panels A). One group were pretreated with ZnCl2, and then injected with free probe (Supplementary Fig. S13, panel B). The other three groups were injected with different amounts of Na2S (1, 5 and 10 equiv.) after i.p. injection of probe (Fig. 6, panels B, C and D). The last group were given i.p. injection of SNP and followed by i.p. injection with the probe. The mice were imaged using a Night OWL II LB 983 small animal \textit{in vivo} imaging system. The fluorescent images showed almost no background fluorescence in the negative control group (Supplementary Fig. S13, panel C, \( R = 0.12 \) in column A), and weak fluorescence in the probe-loaded group (Fig. 6, panel A; panel F, \( R = 1.0 \) in column A), which suggests weak fluorescence signals in probe-loaded mice may be caused by endogenous H2S. To confirm this assumption, we pretreated (i.p. injection) another group of mice with ZnCl2 (an efficient eliminator of H2S). After 10 min of ZnCl2-treatment, the mice were given i.p. injection of free probe. Compared with the free probe-loaded mice, the fluorescence of ZnCl2-treated group is remarkably weakened (Supplementary Fig. S13, panel B; panel C, \( R = 0.18 \) in column B), indicating that the weak fluorescence in probe-loaded mice is triggered by physiological concentration of endogenous H2S. The mice treated with both Na2S (1, 5 and 10 equiv.) and the probe displayed much higher fluorescence (Fig. 6, panels B, C and D) than the mice treated with only the probe, which demonstrate that NIR-HS could respond to exogenous sulphide in mice. Moreover, the mice injected with SNP (Sodium Nitroprusside, a NO donor, could induce the production of endogenous H2S) and probe showed a maked elevation in the fluorescence intensities from the abdominal area of the mice (Fig. 6, panel E, \( R = 3.4 \) in column E), indicating that NIR-HS was sensitive enough to detect endogenous H2S in living mice. Importantly, the fluorescence intensities from the abdominal area of the mice were quantified, and the data showed that the fluorescence intensities triggered by Na2S were concentration-dependent (\( R = 1.0 \) in column A, \( R = 1.6 \) in column B, \( R = 3.5 \) in column C, \( R = 4.8 \) in column D) (Fig. 6, panel F). Figure 7 demonstrated that the fluorescence intensities became strong gradually within 20 min, consistent with the results of titrating the probe with Na2S at different time in PBS buffer (Supplementary Fig. S2). These experiments suggested that NIR-HS is suitable for monitoring exogenous and endogenous H2S in living mice.

Recently, the development of fluorescent probes for H2S \textit{in vivo} is of high interest. A few fluorescent probes have been successfully discovered for imaging of H2S in living animals, such as mice\(^{49-51}\), zebrafish\(^{52-55}\) and Caenorhabditis elegans\(^{56,57}\), et al. In addition to fluorescent probes, luminescent probe and chemiluminescent probe have been applied to determining H2S in living mice\(^{58,59}\). Despite these progresses, the NIR fluorescence imaging of endogenous H2S \textit{in vivo} is still highly desirable. Wallace et al. utilized fluorescent probe SF5 to investigate the regulation of leukocyte H2S synthesis \textit{in vivo}\(^{44}\). However, probe SF5 emitted around 520 nm, the visible-light range limited its application for \textit{in vivo} imaging due to the interference of background autofluorescence. Lu et al. prepared a novel bioluminescence probe for detection of endogenous H2S in nude mice\(^{45}\). Nevertheless, for this
bioluminescence probe, Cys at 15 μM triggered weak bioluminescence. It is well known that the concentrations of Cys in cells/tissue are much higher compared to the concentrations of endogenous H2S, and small response induced by Cys may interference the detection of H2S. Moreover, the H2S reaction site of this bioluminescence probe was azide group. The azide-containing H2S probes could undergo photoactivation under continuous excitation, rendering them unsuitable for in vivo imaging. Thus, NIR probes with high sensitivity, good selectivity and favourable properties to monitor endogenous H2S in vivo are highly needed. Our probe NIR-HS is more suitable for biological imaging endogenous H2S in living mice.

Taken together, we have prepared a novel fluorescent probe NIR-HS for H2S detection in living cells and mice. Advantages of this H2S-specific probe include emission in the NIR region, a low detection limit, high sensitivity, good selectivity and low cytotoxicity. This probe not only enables fluorescence imaging of endogenous H2S induced by SNP in living cells, but also detects endogenous H2S generated in cells overexpressing cystathionine β-synthase (CBS). The probe was also successfully applied to visualizing both the exogenous and endogenous H2S in living mice. Probe NIR-HS shows the potential to be used as a valuable research tool in studying biological roles of H2S. We are currently pursuing other strategies to develop more sensitive and specific fluorescent sensors for monitoring H2S in living animals, as well as the H2S-related medical studies.

Methods

Fluorometric analysis. All fluorescence measurements were conducted at room temperature on a Hitachi F4600 Fluorescence Spectrophotometer. The probe solution (CH3CN) was added to a quartz cuvette. With the probe diluted to 10 μM with 20 mM PBS buffer, Na2S was added (Na2S·9H2O serving as the H2S source in all experiments). The resulting solution was then incubated for 20 min. The samples were excited at 670 nm with the excitation and emission slit widths set at 5 nm and 10 nm, respectively. The emission spectrum was scanned from 690 nm to 850 nm at a velocity of 1200 nm/min. The photomultiplier voltage was set at 1000 V. Data are presented as the mean ± SD (n = 3).

Cell culture and confocal fluorescence imaging. The MCF-7 cells were grown up in DMEM media supplemented with 10% (v/v) FBS (foetal bovine serum) and penicillin/streptomycin (100 μg/mL) at 37 °C in
Cells were permitted to grow to 80% confluence before harvesting and transferring to a coverglass (Lab-Tek® II Chambered Coverglass, NaleNunc, Naperville, USA). A final concentration of 5 μM NIR-HS (1.0 mM stock solution in CH3CN) was added to the cell media and incubated at the previous conditions for 10 min. For exogenous sulphide imaging, the cells were thrice rinsed with PBS solution (pH = 7.4) to remove excess NIR-HS, which was followed by the addition of Na2S (20 mM stock solution in DI H2O, final concentration 25 μM or 50 μM) for incubation 10 min or 20 min at 37 °C. For endogenous sulphide imaging, cells were pretreated with NIR-HS (5 μM) for 10 min and then stimulated with SNP (sodium nitroprusside, 20 mM stock solution in DI H2O, final concentration 50 μM) for 20 min. In addition, cells with CBS overexpression were incubated with NIR-HS (5 μM) for 10 min. All the cells were thrice rinsed with PBS buffer prior to imaging. Confocal fluorescence imaging was performed on an Olympus FV1000 confocal laser scanning microscope with ×60 oil objectives. The excitation wavelength was 635 nm. The fluorescence images (660 nm-760 nm) were obtained at 1024 × 1024 pixels, and were analysed with Olympus software (FV10-ASW). All data are expressed as the mean ± SD (n = 3).

Animals and administration. Adult male Kunming mice weighing 20–25 g were provided by the Experimental Animal Centre of Xuzhou Medical College. All of the experiments were performed in compliance with the Chinese legislation on the use and care of laboratory animals and were approved by the Institutional Animal Care and Use of Xuzhou Medical College. The animals were housed in a room with regulated temperature (22 ± 2 °C) and humidity (50 ± 10%) on a 12 h light/dark cycle; the animals had ad libitum access to standard commercial animal feed and pure water. Mice were acclimatised for 1 week prior to the experiment.

Fluorescent imaging in living mice. The mice were anesthetized by i.p. injection of 10% chloral hydrate (0.04 mL/10g), and their abdominal fur was removed. The mice were random selected and divided into several groups. Subsequently, one group were given i.p. injection of DMSO (50 μL) as the negative control group, and the mice were given i.p. injection of free probe (50 μM, in 50 μL DMSO) as the probe-loaded group. One group were pretreated with ZnCl2 (10 mM, in 100 μL saline), and then injected with free probe. The other three groups were i.p. injected with the probe NIR-HS (50 μM, in 50 μL DMSO), and followed by i.p. injection with different amount of Na2S (50 μM, 250 μM and 500 μM in saline). The last group were i.p. injected with SNP (200 μM, in 100 μL saline), followed by i.p. injection of NIR-HS (50 μM, in 50 μL DMSO). After 20 min, the mice were then imaged by using a Night OWL ILIB 983 small animal in vivo imaging system, with an excitation filter of 670 nm and an emission filter of 690 nm–740 nm. For the time-dependent experiment, the mice were given i.p. injection

Figure 7. Representative fluorescence images of visualizing H2S levels at different times in living mice using NIR-HS. The mice were i.p. injected with the probe NIR-HS (50 μM, in 50 μL DMSO), followed by i.p. injection of 10 equiv. Na2S (500 μM, in 100 μL saline). Images were taken after incubation of Na2S at: 10 min (A); 20 min (B); 30 min (C). Quantification of the fluorescence emission intensities from the abdominal area of the mice of the above groups (D). Data are presented as the mean ± SD (n = 3).
of 10 equiv. Na$_2$S (500 μM in 100 μL saline) after the same disposal of the control mice. Images were then taken at different times (10, 20, and 30 min).

Western blot. For western blot analysis, the cells were washed with cold PBS and lysed with RIPAa buffer (1% Triton X-100, 1% deoxycholate, 0.1% SDS) containing protease inhibitors (1 mM PMSF, 20 mM NaF, 1 mM NaVO$_3$). Protein concentrations were determined using BCA protein assay kits. The cells samples were separated by 10% SDS-polyacrylamide gel electrophoresis (SDS-PAGE) and then transferred onto nitrocellulose membranes (Amersham Pharmacia Biotech). The membranes were blocked with 5% skim milk powder in a washing buffer (Tris-buffered saline containing 0.05% (v/v) Tween 20) for 2 h at 25°C and subsequently incubated overnight with the primary antibodies specific for CBS (1:1000) and β-actin (1:1000). Each membrane was thrice rinsed for 15 min and incubated with either alkaline phosphatase-conjugated secondary antibodies (1:1000, Goat anti-Rabbit IgG antibody) or alkaline phosphatase-conjugated secondary antibodies (1:1000, Horse anti-Mouse IgG antibody), which was followed by visualization by BCIP/NBT alkaline phosphatase colour development kits. Protein bands were scanned and quantified by densitometric analysis using ImageJ version 1.34a software.

Construction of CBS overexpressing cells. The cDNA clones for human CBS were purchased from Origene (lot no.: RC207555). MCF-7 cells were grown to 90% confluency before being transiently transfected with pCMV6-control and pCMV6-CBS expression plasmids using Lipofectamine 2000 (Invitrogen, Shanghai, China) according to the manufacturer’s instructions. Six hours after transfection, the medium containing transfection reagents was removed and incubated in fresh medium. The CBS overexpressed cells were harvested for subsequent experiments.

Statistical analyses. All statistical analyses were performed using SPSS software, version 16.0 (SPSS Inc., Chicago, IL, USA). Values are expressed as the mean ± SD (standard deviation of the mean). The data were analysed with one-way analysis of variance (ANOVA). Statistical significance was set at $p < 0.05$.

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J.Z., L.Z. and Y.L. conceived the idea and directed the work. L.Z. designed the experiments. L.Z., Y.S. and W.M. contributed to the development of experimental schemes. L.Z. performed the experiments. L.Z. and Y.S. contributed to the data analysis. L.Z. and Y.L. contributed to the writing of the paper. Y.S., W.M. and Y.L. critically read the paper.

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Author Contributions

J.Z., L.Z. and Y.L. conceived the idea and directed the work. L.Z. designed the experiments. L.Z., Y.S. and W.M. performed the synthesis. L.Z., F.Z. and Z.X. performed the cell-based imaging. L.Z. and E.L. performed the data analysis. Y.S., W.M. and Y.L. critically read the paper.
in vitro fluorescence tests. L.Z. and X.Z. performed the in vivo imaging. All authors contributed to the data analysis and to writing the manuscript.

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