

# Communication

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# Self-Immolative Thiocarbamates Provide Access to Triggered H<sub>2</sub>S Donors and Analyte Replacement Fluorescent Probes

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Supporting Information Placeholder

**ABSTRACT:** Hydrogen sulfide (H<sub>2</sub>S) is an important biological signaling molecule, and chemical tools for H<sub>2</sub>S delivery and detection have recently emerged as important investigative methods. Key challenges in these two related domains include developing H<sub>2</sub>S donors that are triggered to release H<sub>2</sub>S in response to stimuli, and developing H<sub>2</sub>S probes that do not irreversibly consume H<sub>2</sub>S. Here we report a new strategy for H<sub>2</sub>S donation based on the triggered self-immolation of benzyl thiocarbamates to release carbonyl sulfide (COS), which is rapidly converted to H<sub>2</sub>S by carbonic anhydrase (CA). We leverage the triggered selfimmolative motif to develop a new class of easily-modifiable donors that can be triggered to release H<sub>2</sub>S. We also demonstrate that this approach can be coupled with common H<sub>2</sub>S-sensing motifs to generate scaffolds which, upon reaction with H<sub>2</sub>S, generate a fluorescence response and also release caged H2S, thus addressing challenges of analyte homeostasis in reaction-based probes.

The advent of chemical tools to probe and manipulate biochemical processes has revolutionized how biological processes are investigated.<sup>1-3</sup> Spawning from initial investigations into fluorescent proteins,<sup>4-5</sup> small molecule fluorescent reporters now comprise a key pillar of investigative chemical biology with a remarkable diversity of fluorescent tagging and measurement technologies.<sup>6-7</sup> Recent years have witnessed a significant expansion of sensor development to include chemical tools for imaging transition metal, alkali, and alkali earth ions.<sup>8-10</sup> Many of these sensing platforms can provide real-time, quantitative measurements of ion fluxes due to the reversible interaction of the sensor with the analyte, thus providing methods for imaging the dynamic process of metal ion trafficking associated with important signaling events ranging from Ca<sup>2+</sup> sparks during muscle contraction<sup>11</sup> to Zn<sup>2+</sup> fluxes during mammalian egg fertilization.<sup>12</sup> Complementing such investigative tools are small molecule donors that release caged analytes with controllable rates.<sup>13-16</sup> Such platforms provide powerful methods for controlling levels of specific small molecules, which often include pro-drugs, metal ions, or small reactive sulfur, oxygen, and nitrogen species (RSONS), in different biological contexts.

In the last two decades, RSONS have emerged as important bioinorganic molecules involved in myriad biological processes, many of which have been elucidated by utilizing chemical tools for small molecule detection and delivery. RSONS are involved in the complex cellular redox landscape and are often involved in oxidative stress responses, immune responses and signaling pathways, as well as other emerging roles.<sup>17</sup> For example, NO, HNO,

and ONOO- play important roles ranging from smooth muscle relaxation to immune response<sup>18</sup> and are largely intertwined with reactive oxygen species, such as O<sub>2</sub><sup>-</sup> and H<sub>2</sub>O<sub>2</sub>, which are critical in oxidative stress responses and have been implicated in various aging mechanisms.<sup>19</sup> Similarly, reactive sulfur species, such as H<sub>2</sub>S, hydropolysulfides (HS<sub>n>1</sub><sup>-</sup>), and persulfides (RSSH) have recently garnered interest as important biological signaling molecules with roles in long term potentiation and cardiovascular health.<sup>20</sup> By contrast to their metal ion counterparts, RSONS are often fleeting in nature and often react irreversibly with cellular targets. This heightened reactivity has provided chemists with significant challenges in developing constructs that can release these reactive molecules under controlled conditions, but have also provided chemists with different strategies to devise chemical tools for their detection by engineering reactive groups onto sensing platforms that react selectively albeit irreversibly with the analyte of interest.21

Although small molecule donors and reaction-based probes have provided significant insights into the roles of RSONS in biology, key needs remain. For example, engineering donors with precise but modifiable triggers to enable analyte release in response to specific stimuli and developing reaction-based probes that do not irreversibly consume the analyte would enable new insights into RSONS biology. Motivated by these needs, as well as our interest H<sub>2</sub>S chemistry, we report here a new caged H<sub>2</sub>S releasing strategy and provide proof-of-concept applications in both small-molecule donor and reaction-based probe design. By leveraging triggerable self-immolative thiocarbamates, we demonstrate access to easily-modifiable H<sub>2</sub>S donors that can be triggered by external stimuli (Figure 1a), and address common issues of analyte consumption in reaction-based fluorescent probes (Figure 1b) by developing analyte-replacement reactionbased platforms (Figure 1c).





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**Figure 1.** (a) Caged donors triggered by different stimuli. (b) Reaction-based probes typically consume the target analyte. (c) Analyte-replacement reaction-based probes enabled by incorporation of caged analytes into reaction-based motifs.

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59 60 Development of analyte-replacement sensing platforms requires two important components: a versatile H<sub>2</sub>S donation motif that releases H<sub>2</sub>S in response to a specific triggering event, and a method to couple this caged donor to a reaction-based sensing platform with an optical output. As a proof-of-concept design toward this objective, we chose to use H2S-mediated azide reduction for our sensing platform, which has emerged as the most common method for H<sub>2</sub>S detection and exhibits high selectivity for H<sub>2</sub>S over other RSONs (Figure 2a).<sup>22</sup> Although a number of H<sub>2</sub>S-donating motifs have been reported and have found utility as important research tools,<sup>23-25</sup> none of these fit the design requirement of our approach. To develop an H2S donating motif compatible with our design requirements, we reasoned that common strategies in drug and fluorophore release, namely the selfimmolative cascade decomposition of para-functionalized benzyl carbamates (Figure 2b),<sup>26, 27, 28</sup> could be modified to enable triggered H<sub>2</sub>S release. Because self-immolative carbamates release an amine-containing payload and extrude CO2 as a byproduct, we reasoned that replacing the carbonyl oxygen with a sulfur atom to generate a thiocarbamate would result in carbonyl sulfide (COS) rather than CO<sub>2</sub> release (Figure 2c). In a biological environment, COS is quickly hydrolyzed to H<sub>2</sub>S and CO<sub>2</sub> by carbonic anhydrase (CA), which is a ubiquitous enzyme present in plant and mammalian cells.<sup>29-30</sup> The second byproduct of the thiocarbamate self-immolation is a reactive quinone methide, which rapidly rearomatizes upon reaction with available nucleophiles, such as water or nucleophilic amino acids such as cysteine.<sup>31-32</sup> On the basis of the requirements outlined above, we expected that a quenched fluorophore could be functionalized with a pazidobenzylthiocarbamate to enable H2S-mediated azide reduction to form the transient aryl amine intermediate, which would subsequently undergo the self-immolative cascade reaction to extrude COS/H<sub>2</sub>S and liberate the fluorophore to access an analytereplacement sensing motif (Figure 2d). Previous work

(a)  $H_2S$ -mediated azide reduction FL  $N_3$   $H_2S$  FL  $NH_2$ (b) Self-immolative carbamates

This work (c) Thiocarbamate-based COS/H<sub>2</sub>S donor development



**Figure 2.** Established strategies for using (a)  $H_2S$ -mediated azide reduction in  $H_2S$  sensing, and (b) self-immolative carbamates to deliver an amine-bound payload, such as a fluorescent dye. Incorporation of self-immolative thiocarbamates enables access to (c) triggered  $H_2S$  donors, and (d) analyte replacement reaction-based probes.

To confirm that the COS could serve as a potential source of  $H_2S$  donation, we first established that independently prepared COS

could be efficiently hydrolyzed to H<sub>2</sub>S by CA. Upon addition of COS gas to deoxygenated aqueous buffer (PBS, 1 mM CTAB, pH 7.4) containing CA from bovine erythrocytes, we observed rapid H<sub>2</sub>S production using an H<sub>2</sub>S-responsive electrode. In the absence of CA, negligible current was observed from COS alone, which is consistent with slow and pH-dependent, nonenzymatic hydrolysis in water (Figure S4).<sup>33</sup> We also observed a dose-dependent reduction in H<sub>2</sub>S production upon addition of the CA inhibitor acetazo-lamide (AAA),<sup>34</sup> which confirmed the enzymatic hydrolysis of COS by CA (Figure 3).



**Figure 3.** Conversion of COS to H<sub>2</sub>S by carbonic anhydrase (CA) with varying concentrations of CA inhibitor acetazolamide (AAA) in PBS buffer, pH 7.4.

We next prepared model thiocarbamates to confirm that the proposed decomposition cascade to release COS occurs efficiently and to demonstrate the biological compatibility of this new H2S donor motif. We incorporated an azide in the para position of the benzylthiocarbamate to function as the H2S-responsive trigger for self-immolation and COS release. To facilitate NMR identification of the products, we first prepared azidobenzylthiocarbamate 1 with a *p*-fluoroaniline payload, and the corresponding carbamate 2 as a control compound (Figure 4a-c). Although 2 should undergo the same self-immolative decomposition upon azide reduction, it releases CO2 rather than COS, and thus should not result in H2S donation upon reaction with CA. To monitor the reactivity of the model compounds under controlled reaction conditions, we used tris(2-carboxyethyl)phosphine (TCEP), an azide-reducing agent, to trigger self-immolation, due to its near-instantaneous reduction of azides. In each case, <sup>1</sup>H, <sup>13</sup>C{<sup>1</sup>H}, and <sup>19</sup>F NMR spectroscopy was used to monitor the reaction after reduction of the model complexes by TCEP. Consistent with our design hypothesis, we observed the disappearance of the benzylic peak, loss of the thiocarbonyl carbon peak, and formation of new resonances upon self-immolation by NMR spectroscopy (Figure S1-S3). All such changes were observed within 5 minutes of TCEP addition, confirming the rapid self-immolation of the scaffold upon reduction, and were consistent with COS release from the thiocarbamate scaffold upon azide reduction.

Having confirmed that CA rapidly catalyzes the hydrolysis of COS, we next investigated the H<sub>2</sub>S-donating ability of model compounds 1 and 2 under identical conditions. Monitoring thiocarbamate 1 in buffer containing CA did not result in H2S formation, confirming that the thiocarbamates do not react directly with CA and that aryl azides are stable in the presence of CA (Figure S5). Upon injection of TCEP, however, rapid release of H<sub>2</sub>S was observed, indicating that reduction of the azide to an amine is essential to trigger self-immolation and COS release. Additionally, repetition of the experiment with added AAA significantly reduced the rate of H2S production, confirming that uninhibited CA is required for significant H<sub>2</sub>S production from the triggered thiocarbamate scaffold (Figure 4d). Finally, the analogous carbamate (2) was investigated under identical conditions, and as expected no H<sub>2</sub>S was produced upon addition of TCEP, confirming that the sulfur-containing thiocarbamate is required for H<sub>2</sub>S formation. In total, these experiments demonstrate the validity of using the thiocarbamate group as a triggerable source of H<sub>2</sub>S release in aqueous solution, which we expect will prove fruitful

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Figure 4. (a,b) Synthesis of model thiocarbamates and carbamates. (c) Model compounds. (d) H<sub>2</sub>S release from 1 after reduction by TCEP in the presence of CA, under identical conditions with the addition of AAA (2.5  $\mu$ M,), and from carbamate 2. (e) Quantification of total sulfide in whole mouse blood after treatment 25  $\mu$ M 3 and 4 after 30 min of incubation time in the presence of excess TCEP.

Expanding on our cuvette-based studies, we also investigated H<sub>2</sub>S release from model thiocarbamates in whole mouse blood to expand on the efficacy of H<sub>2</sub>S release from thiocarbamates in biologically-relevant contexts. Although murine systems provide a convenient model, mice have among the lowest CA levels in mammals, with murine blood only containing about 15% of the CA present in human blood,<sup>35</sup> and thus represent a challenging target for sulfide release mediated by CA. To quantify total sulfide levels, we used the monobromobimane (mBB) method which allows for the analytical measurement of different sulfide pools and is compatible with many types of biological samples.<sup>36</sup> Measurement of the total sulfide, which includes free sulfide as well as bound sulfane-sulfur, revealed background levels of 8 µM, which are higher than total sulfide levels commonly observed and reported in plasma, but are consistent with the high sulfane-sulfur content in red blood cells.<sup>37, 36</sup> We prepared thiocarbamate 3, which lacks the azide trigger, to confirm that the thiocarbamate group was stable in whole blood and did not release COS without activation of the trigger group, and compared results obtained with this model compound with azide-functionalized 4. Total sulfide levels were measured for each compound, as well as the control, after 30 minutes of incubation with excess TCEP (Figure 4e). Consistent with our expected results, only samples containing donor 4 with the azide trigger increased total sulfide levels in blood ( $p \leq 0.0001$ ). These results establish the stability of the thiocarbamate in biological milieu and confirm that endogenous CA in murine blood, even though significantly lower than in most other biological environments,<sup>35</sup> is sufficient to hydrolyze the COS released from thiocarbamates after the self-immolation cascade is triggered, highlighting the efficacy of this H<sub>2</sub>S-releasing strategy in biological environments.

Having confirmed the viability of triggered  $H_2S$  release with the model compounds, we next applied this design to incorporate a fluorophore to access an  $H_2S$ -responsive fluorescent probe that releases  $H_2S$  upon  $H_2S$  detection. Our primary goal was to demonstrate that the thiocarbamate group could be appended to common fluorophore motifs and efficiently quench the fluorescence. We chose to use methylrhodol (MeRho)<sup>38</sup> as the fluorophore due to its high quantum yield and single fluorogenic amine, which could be readily converted into the desired thiocarbamate. Since the azide-functionalized scaffold would be triggered by H<sub>2</sub>S to release both MeRho and COS, this would function as a fluorescent H<sub>2</sub>S probe that would replenish sulfide through the release of COS. To access the desired scaffold, we treated MeRho with thiocarbonyldiimidazole (TCDI) and NEt3 in DMF to afford methylrhodol isothiocyanate (MeRho-NCS) in 60% yield. Subsequent treatment with 4-azidobenzyl alcohol and NaH afforded the methylrhodol thiocarbamate azide (MeRho-TCA) in 35% yield (Figure 5a). We note that one benefit of this simple synthetic route is that almost any fluorophore containing a fluorogenic nitrogen can readily be functionalized with the benzylazide thiocarbamate group, thus providing access to a diverse library of fluorophores.

With a sulfide-replenishing H<sub>2</sub>S probe in hand, we investigated the fluorescence response upon addition of sulfide. Treatment of MeRho-TCA with 50 equiv. of NaSH in aqueous buffer (PBS, 1 mM CTAB, pH 7.4) resulted in a 65-fold fluorescence turn-on over 90 minutes (Figure 5b). Additionally, we confirmed that the MeRho-TCA scaffold was selective for HS<sup>-</sup> over other RSONs, by measuring the fluorescence response to Cys, GSH, Hcy, S<sub>2</sub>O<sub>3</sub><sup>2-</sup> , SO32-, SO42-, H2O2, and NO (Figure 5c). As expected, the MeRho-TCA scaffold exhibited excellent selectivity for sulfide over other RSONs, demonstrating that the thiocarbamate linker group did not erode the selectivity of the azide trigger, and also establishing that the MeRho-TCA scaffold can function as a viable H<sub>2</sub>S reporter. Because MeRho-TCA releases H2S upon reaction with H<sub>2</sub>S, we note that one consequence of this analyte replacement approach is that the resultant fluorescence response is not directly proportional to the initial H<sub>2</sub>S concentration. Additionally, in isolated systems, two equiv. of HS- are required for complete azide reduction, suggesting that the first-generation analytereplacement scaffolds only replace one half of the consumed sulfide.<sup>39</sup> It is also possible, however, that in biological media one equiv. of a thiol may play a role in H<sub>2</sub>S-mediated azide reduction, which remains a question for future investigations. In the present system, preliminary mechanistic investigations indicate that H<sub>2</sub>Smediated azide reduction is the rate-limiting step of the selfimmolative process, and that the subsequent release of COS and hydrolysis by CA to form H<sub>2</sub>S is rapid. Taken together, these data highlight the potential of this strategy to access analytereplacement, reaction-based fluorescent scaffolds.

In summary, we have outlined and demonstrated a new strategy for triggered H<sub>2</sub>S release based on self-immolative thiocarbamates. Importantly, this strategy provides solutions to key challenges associated with both H<sub>2</sub>S delivery and detection. Thiocarbamate-based H<sub>2</sub>S donors provide a new, versatile, and readily modifiable platform for developing new H<sub>2</sub>S donor motifs that can be triggered by endogenous or biorthogonal triggers. Similarly, this same H<sub>2</sub>S donation strategy can be coupled to fluorescent probe development to access reaction-based fluorescence reporters that replace the analyte that has been consumed by the detection event. In a broader context, we expect that the selfimmolative thiocarbamate donors will find significant utility as a potential platform for academic and potentially therapeutic H<sub>2</sub>S donors. Moreover, we anticipate that similar strategies can be applied to other RSONS reaction-based sensing methods to provide access to analyte-replacement sensing strategies for reactionbased sensing motifs.

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**Figure 5.** (a) Synthesis of **MeRho-TCA**. (b) Fluorescence response of **MeRho-TCA** to H<sub>2</sub>S. Inset shows integrated fluorescence over time by comparison to **MeRho-TCA** in the absence of NaSH. (c) Selectivity of **MeRho-TCA** for H<sub>2</sub>S over other RSONs. Conditions: 5  $\mu$ M probe, 250  $\mu$ M RSONs unless noted otherwise, in PBS buffer, 1 mM CTAB, pH 7.4, 37 °C.  $\lambda_{ex} = 476$  nm,  $\lambda_{em} = 480-650$  nm.

# ASSOCIATED CONTENT

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#### Supporting Information

Experimental details, H<sub>2</sub>S release profiles, spectra. This material is available free of charge via the Internet at http://pubs.acs.org. **ACKNOWLEDGMENT** 

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#### Conflict of Interest Disclosure

The authors have filed a provisional patent on thiocarbamatebased and other COS-related donation strategies, including those outlined within this manuscript.

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