Fragment-based de novo design of a cystathionine γ-lyase selective inhibitor blocking hydrogen sulfide production

Angela Corvino1,5, Beatrice Severino1, Ferdinando Fiorino1, Francesco Frecentese1, Elisa Magli2, Elisa Perissutti2, Vincenzo Santagada1, Mariarosaria Buccig, Giuseppe Cirino1, Geoff Kelly2, Luigi Servillo3, Grzegorz Popowicz4, Annalisa Pastore5 & Giuseppe Caliendo1

Hydrogen sulfide is an essential catabolite that intervenes in the pathophysiology of several diseases from hypertension to stroke, diabetes and pancreatitis. It is endogenously synthesized mainly by two pyridoxal-5'-phosphate-dependent enzymes involved in L-cysteine metabolism: cystathionine-β-synthase (CBS) and cystathionine-γ-lyase (CSE). Research in this field is currently impaired by the lack of pharmacological tools such as selective enzymatic inhibitors that could target specifically only one of these pathways. We used a novel approach based on a hybrid method that includes drug design, synthetic biology, metabolomics and pharmacological assays to rationally design a new inhibitor selective for the CSE enzyme. The identification of this compound opens new frontiers towards a better understanding of the role of CSE over CBS in the pathophysiology of diseases where a role for the H2S pathway has been proposed and the development of new lead compounds that could target the CSE enzyme.

Hydrogen sulfide (H2S), a colorless, flammable, water soluble gas with the characteristic smell of rotten eggs, has emerged as an important gaseous signaling molecule playing numerous roles in health and disease, along with CO and NO. Enzymatically generated H2S is mainly derived from two pyridoxal-5'-phosphate (PLP)-dependent enzymes responsible for the metabolism of L-cysteine (L-Cys): cystathionine-β-synthase (CBS) and cystathionine-γ-lyase (CSE). A third pathway that catalyses the production of H2S from L-Cys via the combined action of 3-mercaptopyruvate sulfurtransferase and cysteine aminotransferase has also been described. This pathway is less well characterized and its role in determining the H2S levels in tissues still poorly understood.

CBS and CSE are cytosolic enzymes which catalyse several H2S-generating reactions which convert L-Cys and/or homocysteine as substrates to L-cystathionine and pyruvate respectively. CBS was also originally considered to be the predominant enzyme for H2S production in the brain, indeed it is preferentially expressed in radial glia/astrocytes of adult and developing mouse brain, whereas H2S synthesis in the heart and vasculature was attributed to CSE. More recent studies with improved markers have provided a broader picture of enzyme distribution. Because of the co-presence of both enzymes in specific pathway it is important to find inhibitors that selectively target only one enzyme.

The most commonly used agents to inhibit H2S biosynthesis include propargylglycine (PAG), β-cyanoalanine (BCA) and aminooxyacetic acid (AOAA). PAG is described as CSE selective inhibitor in fact it will not inhibit recombinant CBS even if used at 10 mM; moreover the crystal structure of the covalent complex PAG-CSE is the only described so far. However all of these compounds have a relatively low potency and cell permeability and are used at high concentrations (mM). L-aminoethoxyvinylglycine (AVG) was also recently described as a potentially more potent and selective CSE inhibitor but its mechanism is still uncharacterised. The inherent

1Department of Pharmacy, University of Naples “Federico II” – Via D. Montesano, 49, 80131–Napoli, Italy. 2MRC Biomedical NMR Centre, NIMR, NW7 1AA, London. 3Department of Biochemistry, Biophysics and General Pathology, Second University of Naples, Italy. 4Technical University of Munich, Garching Campus, Munich, Germany. 5Department of Neuroscience, Wohl Institute, King’s College London, Denmark Hill Campus, London SE5, UK.

Correspondence and requests for materials should be addressed to A.P. (email: annalisa.pastore@crick.ac.uk) or G.C. (email: giuseppe.caliendo@unina.it)
flexibility of AVG makes it in fact difficult to rationalise the mode of binding of this compound until a crystal structure of its complex with CSE becomes available. Lack of biological data assessing the role of this compound in cell culture, tissue baths and in vivo further studies have also discouraged an interest on AVG, leaving PAG as the reference usually considered.

The possibility to inhibit selectively only one of these enzymes has tremendous therapeutic potentialities. CBS inhibition has for instance been suggested as a potential therapeutic to the treatment of ischemic injury. Particularly interesting would be to be able to inhibit selectively the CSE pathway since this possibility would have important consequences in several pathologies, e.g. asthma and hemorrhagic shock. PAG, for instance, has been shown to inhibit H2S CSE-mediated production in animal models and to accelerate recovery of blood pressure after hemorrhagic shock14. PAG was also shown to prevent the increase in plasma levels of markers of liver and pancreas injury, reduce the tissue content of myeloperoxidase15 with consequent significant prolongation of animal survival16. An important challenge which holds the promise to allow a much better comprehension of the H2S metabolism and that could be used for therapeutic purposes is thus the search for selective inhibitors able to block only one of the two enzymes and the establishment of the mechanism of interaction. However, the goal of finding such inhibitor is not an easy task given the similarity between the substrates and the mechanism involved in the enzymatic activity of the two enzymes.

We set with the aim of producing new and more selective inhibitors of CSE. Using a semi-rational in silico drug screening, we went through the synthesis and structural characterization of a selective CSE inhibitor. We tested the new compounds in tissue using an ex vivo assay with intact rat aortic rings. The compound showing maximal inhibitory effects in this test is an oxothiazolidine derivative. The effects of this compound on the enzyme kinetics were further tested in vitro on the purified enzymes using a metabolomics approach based on nuclear magnetic resonance techniques. By setting up complementary and more comprehensive ex vivo and in vitro assays, we also established a more accurate way to study the H2S pathway and demonstrated the limitations of drawing conclusions only on the basis of the methylene blue assay, the most commonly used methodology for this purpose17,18. We could conclusively prove that the identified compound has properties of a new selective inhibitor of CSE. Our results bear crucial bio-technological and pathophysiological implications which will help further studies of the H2S pathway.

### Results and Discussion

**Rationale of the design of the new compounds.** We first carried out a preliminary in silico analysis of the structures of the two enzymes to understand the geometric properties of the active sites. Although both homo-tetrameric and PLP bound, their sequences and structures are very different. In CSE, essential for the reaction mechanism is a PLP covalently attached to Lys212 and stabilized by \( \pi \)-stacking interactions with the ring of Tyr114. In the complex with the inhibitor PAG, this is covalently bound to Tyr114 as a vinylether13. The amino and carboxyl groups of PAG form hydrogen bonds with Glu339 and with Arg119 and Arg62 from the adjacent monomer respectively. In CBS, PLP is bound to Lys11919,20 and is surrounded mostly by small amino acids, such as Ser349 and Asn149.

Informed by this structural analysis, we used a fragment-based approach in which we fused the structural features of the substrate cysteine with those of already known CSE inhibitors. We started by commercially available cysteine derivatives already used as substitutes of cysteine and selected L-thiazolidine-4-carboxylic acid and L-2-oxothiazolidine-4-carboxylic acid (indicated as compounds 1 and 2 in Table 1). We added to these scaffolds a propargyl group in \( \alpha \)-carboxyl position since this structural element is essential in the proposed PAG mechanism of inhibition13 (compounds 1a and 2a). Independently, we considered the commercially available L-2-amino-3- (thiophen-2-yl)propanoic acid and 5-amino-2-hydroxybenzoic acid which were interesting because the thiiazolidine moiety is substituted by aromatic groups (compounds 3 and 4). We reasoned that an important part of the

| Compd | Structure | Compd | Structure |
|-------|-----------|-------|-----------|
| 1     | ![Structure](image1) | 1a    | ![Structure](image2) |
| 2     | ![Structure](image3) | 2a    | ![Structure](image4) |
| 3     | ![Structure](image5) | 3a    | ![Structure](image6) |
| 4     | ![Structure](image7) | 4a    | ![Structure](image8) |

Table 1. Chemical formulas of the compounds commercially available (left two columns) and synthesized for the present work (right two columns).
CSE active site is constituted by Tyr114 and PLP which form π-stacking interaction. The presence of an aromatic group could thus facilitate interaction of the new compound21. We thus attached a propargyl moiety to these scaffolds obtaining compounds 3a and 4a. This allowed us to have novel molecular entities which fuse the structural features of both cysteine and the CSE inhibitor PAG.

Pharmacological assays suggest compound 2a as an inhibitor. The inhibitory properties of the commercially available compounds 1–4 and their newly synthesized derivatives were tested by an ex vivo assay on rat aortic rings by which the inhibitory activity on L-cysteine induced vasorelaxation was probed. All compounds reduced L-Cys induced vasorelaxation, with compounds 2, 3 and 4 showing a higher inhibitory activity (for the E_{max} values refer to Suppl. Information) (Fig. 1A). Incubation of aortic rings with compound 1a did not affect the L-cysteine induced vasorelaxation. In contrast, compound 2a completely abrogated L-cysteine-induced vasorelaxation at a concentration of 100 μM (Fig. 1B). An inhibitory effect was observed also with compound 3a. However, as compared to 2a, compound 3a was unable to completely abrogate relaxation even at the highest concentration tested (1 mM). Compound 4a, tested at a lower concentration (10 μM) because of its limited solubility, significantly inhibited L-cysteine induced vasorelaxation. However, its solubility limits the potential use of this compound as an inhibitor and makes it difficult to assay the compound in aqueous media in a vast range of concentrations.

These results showed that only conversion of compound 2 into the propargyl derivative 2a results in a compound with good solubility which leads to an improvement of the inhibitory activity.

The inhibitory effect of compound 2a is direct and higher than that of known inhibitors. Compound 2a, which shows appreciable inhibition, was tested by the well-known methylene blue assay which measures H_{2}S production (Fig. 2). Under basal conditions, tissue homogenates produced 1.4 ± 0.1% nmol of H_{2}S per mg of protein. H_{2}S production was stimulated by adding exogenous L-Cys into the vehicle, as routinely performed in this kind of experiment22,23. The modification doubled H_{2}S production. Incubation with compound 2a significantly inhibited the increased H_{2}S production stimulated by L-Cys showing an effect comparable to that obtained with PAG but at a lower dose (100 μM for compound 2a and 10 mM for PAG). We also compared the inhibitory activity of compound 2a with that of PAG (Fig. 3A,B) by using isolated aortic rings. Incubation with compound 2a completely abrogated L-cysteine-induced vasorelaxation, while PAG reduced but not completely abrogated L-cysteine-induced vasorelaxation, despite the compound was tested at a higher concentration (10 mM and 100 μM for PAG and 2a respectively).

To compare the efficacy of the two inhibitors, IC_{50} values were calculated determining the effect on the aorta rings relaxation in semilogarithmic scale at three different concentrations of the tested compounds. The obtained results, expressed as percent of relaxation (Fig. 3) allowed us to get the IC_{50} values. We obtained 3.3 ± 0.3 mM for PAG and 31.6 ± 0.6 μM for the compound 2a indicating a two order magnitude improvement of the latter as compared to PAG.

These results propose compound 2a as a more potent inhibitor than previously known ones.

To estimate the plasma membrane crossing properties of the new inhibitor we calculated the partition n-octanol/aqueous phase coefficients (LogP_{w} values) of the neutral form of 2a (c log P = 0.27) as compared to PAG (c log P = −3.02) where logP is a theoretically calculated lipophilicity parameter that refers to the partition coefficient (P) in n-octanol/water of the neutral species. This tells us that the compound 2a is more hydrophobic than PAG and this property justifies the higher concentration (10 mM) of PAG required to cross the membranes and to exert, consequently, a maximal inhibitory effect.
We also tested the importance of the propargyl group by substituting this with a propyl group which bears approximately the same volume, although having a different geometry (compound 2b). When we tested this compound on rat aortic rings (Figure S3 of Suppl. Information), we observed no inhibitory effects indicating that the propargyl group is essential.

Inhibition is caused by the direct and selective interaction with the CSE enzyme. The effects of the selected compound were then tested on the purified recombinant CSE and CBS enzymes as this is the only way to prove that the effects observed in ex vivo assays are solely the consequence of a direct and specific interaction between the compound and the enzymes. The CBS and CSE enzymes were first expressed in Escherichia coli, purified and characterized by SDS-PAGE, absorbance and circular dichroism techniques which confirmed that the enzymes are correctly folded and bind the respective prosthetic groups (Figure S4 of Suppl. Information).

We used metabolomics to follow directly the conversion of L-Cys into the final products, which are respectively cystathionine and pyruvate for CBS and CSE. This strategy, which relies on high accuracy techniques such as
as nuclear magnetic resonance (NMR) and/or mass spectrometry (MS), has by now been vastly validated for the
detection of metabolites, enzymatic products and metabolic pathways. Because of its ability to identify all
products rather than only specific ones, as in the methylene blue assay, metabolomics can powerfully complement
these assays and provide new information. NMR spectra were recorded to detect the progressive formation of the
products starting from 15N,13C-labelled substrate. For the reaction catalyzed by CBS, unlabeled L-homocysteine
was also added as co-substrate. In the interest of following the kinetics at close intervals, we recorded only the first
increment of 13C HSQC spectra at fixed time intervals (Figure S5 of Suppl. Information). Complete 2D 13C HSQC
spectra were collected before and after each experiment to confirm completeness of the reactions. The results were
best visualized as a pseudo 3D experiment in which each spectrum is plotted along the x and y axes and the third
axis reflects time (Fig. 4).

In the control kinetics of CBS catalyzed production of L-cystathionine (Fig. 4A), the 15N,13C-labelled L-Cys
resonances gradually decrease up to complete disappearance, while new resonances increase in intensity. The
expected cystathionine resonances were not all detected, because of the unlabeled homocysteine contribution. In
the control kinetics of the CSE-catalyzed reaction, the 15N,13C, L-Cys peaks decrease up to complete disappear-
ance, while the pyruvate product increases (Fig. 4B). The presence of compound 2a solutions pre-incubated with
CBS did not alter the kinetics and the reaction proceeded with the practically complete conversion of L-Cys to
L-cystathionine (Fig. 4C), whereas inhibited CSE catalyzed pyruvate production (Fig. 4D). These results conclu-
sively prove that compound 2a is a powerful inhibitor selective for CSE with no effects on the kinetics of CBS.
Metabolomics reveals the limitations of the methylene blue assay. During our NMR based metabolomics studies we observed something peculiar: when PAG was used in the NMR assay, no appreciable formation of pyruvate was observed, as expected, but the resonances of L-Cys surprisingly decreased and new product resonances, different from pyruvate, appeared and progressively increased (Fig. 5A). This behaviour was also confirmed in a $^{13}$C 2D HSQC spectrum acquired at the end of the kinetics: when PAG was present, the cysteine resonances at 3.90 ppm ($C_\alpha$ and 55 ppm in the carbon spectrum not shown) and at 3.05 ppm ($C_\beta$H$^2$ and 25 ppm) were absent at the end of the reaction. This means that, although PAG inhibits $H_2S$ formation, it is not able to block cysteine consumption. This is per se an important result which warns us against the inefficacy of PAG. New peaks at 3.92 ppm, 3.40 ppm and 3.25 ppm appeared instead (and at 55 ppm, 32 ppm and 32 ppm in the carbon spectrum respectively). These chemical shifts were tentatively assigned to cystine, lanthionine or a...
similar compound. To confirm which of these compounds is the additional product we used the high accuracy LC-MS technique\(^2\). We found that, during the reaction, a predominant quantity of cysteine is converted into cystine with a minor quantity of lanthionine. Identification of this reaction product has so far been overlooked likely because all previous studies solely relied on the methylene blue assay which allows only detection of H\(_2\)S production. Since the presence of an excess of freshly prepared DTT makes it unlikely chemical formation of cystine by oxidation, a possible explanation is that cystine is formed enzymatically.

**In vitro assays suggest that the new inhibitor is not suicidal and competitive.** We characterized the complex of CSE with compound \(2a\) by mass spectrometry to test whether also this inhibitor, like PAG, is suicidal. We found no increase of the mass of CSE suggesting no covalent attachment of compound \(2a\) (Figure S6 of Suppl. Information). This result agrees with the suggested mechanism of inhibition which postulates the need of a NH\(_2\) group for the reaction to occur\(^1\). To quantify the inhibitory potency towards CSE of compound \(2a\) with respect to PAG, we used an adapted version of a previously described spectrophotometric assay\(^1\). The IC\(_{50}\) measured on the purified protein is 6.3 \(\pm\) 1.0 \(\mu\)M for \(2a\) versus 42.2 \(\pm\) 1.1 \(\mu\)M for PAG (Figure S7 of Suppl. Information). This value is in excellent agreement with the data from the ex vivo assay.

With the same spectrophotometer assay described above, we also determined the Michaelis-Menten constants \(V_{\text{max}}\) and \(K_m\), the maximum reaction rate at maximum saturation of the substrate concentration, and \(K_m\). For the data at 0 \(\mu\)M, 5 \(\mu\)M, 10 \(\mu\)M and 20 \(\mu\)M of inhibitor, respectively. (B) Slopes of the double reciprocal plot vs. the concentrations of inhibitor (0, 5, 10 and 20 \(\mu\)M) (Fig. 6B). The intercept gives the value of \(K_m/V_{\text{max}}\) and the slope is \(K_m/(V_{\text{max}} K_i)\). \(K_i\) is obtained as the intercept/slope ratio.

**Figure 6.** Enzymatic characterization of the effects of compound \(2a\) on purified CSE. (A) Lineweaver-Burk plot of CSE catalyzed H\(_2\)S production in the absence (control) and presence of various concentrations of compound \(2a\) (5\(\mu\)M, 10\(\mu\)M and 20\(\mu\)M). The data suggest that \(2a\) is a competitive inhibitor, since the plots are linear and intersect at the y-axis (\(K_m\) remains unaffected). For reference, non-competitive inhibition would produce plots with the same x-intercept as uninhibited enzyme (\(K_m\) is unaffected) but different slopes and y-intercepts. Uncompetitive inhibition would cause different intercepts on both the y- and x-axes. The estimated \(V_{\text{max}}\) and \(K_m\) values are 0.50 \(\pm\) 0.02 U/mg and 1.1 \(\pm\) 0.1 mM. The \(K_{m,\text{apparent}}\) values are 1.5 \(\pm\) 0.1 mM, 1.9 \(\pm\) 0.3 mM and 2.8 \(\pm\) 0.3 mM for the data at 0 \(\mu\)M, 5 \(\mu\)M, 10 \(\mu\)M and 20 \(\mu\)M of inhibitor, respectively. (B) Slopes of the double reciprocal plot vs. the concentrations of inhibitor (0, 5, 10 and 20 \(\mu\)M) (Fig. 6B). The intercept gives the value of \(K_m/V_{\text{max}}\) and the slope is \(K_m/(V_{\text{max}} K_i)\). \(K_i\) is obtained as the intercept/slope ratio.

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Rationalizing the results. We rationalized our findings analysing the structure of CSE. Starting from the coordinates of CSE (PDB code 3cog) which contain the covalently bound PAG, we built in manually the coordinates of compound 2a using the software MAESTRO (Schroedinger, LLC, New York, NY). The cavity which contains PAG in the active site, which is also where the substrate cysteine should bind, is big enough to accommodate also compound 2a (Fig. 7).

This can easily form multiple interactions with the surrounding groups and hydrogen bond with the side chains of Ser63, Tyr114, Gly116, Arg119, Asn241, and Ser358. Amongst these interactions, particularly important would be those formed by the carbonyl group of the L-2-oxothiazolidine moiety which could form a bifurcated hydrogen bond with Gly116 and Ser358. The triple bond would occupy a position similar (<2Å difference) to that of PAG without being covalently bound and could not easily be replaced because the pocket between the two tyrosines is very deep and narrow. A triple bond forms a long thin “rod” of very small van der Waals radius which could thus fit in, while double or single bonds could not be allocated between the tyrosines for lack of space. Pi-stacking of the triple bond with the two tyrosine rings could additionally contribute to the stability of the interaction.

This configuration would also explain why compounds 1 and 1a do not fit: they have no carboxy group which makes the ring to be bent while it is quite flat in 2a and this prevents the possibility of hydrogen bonding to the carbonyl oxygen. The bent ring will most likely not fit in the pocket. Compounds 3 and 3a have a CH2 linker which positions the ring too far from the pocket to form interactions and changes its angle relative to the triple bond. Compounds 4 and 4a are simply too big for the pocket.

We can thus conclude that we have identified a new compound which has all features to optimally fit in CSE and inhibit selectively its function.

Conclusion

Investigation of the pathophysiological and pharmacological roles of hydrogen sulfide has rapidly become an important research field still widely unexplored which is attracting increasing attention. H2S is now accepted as a novel gasotransmitter of potential importance in several areas from asthma to cancer, impotence and stroke. A major obstacle in the rapid development of this field has however been the lack of pharmacological tools such as selective enzymatic inhibitors which would allow the study of only one among the several possible pathways. The already established inhibitors exhibit low potency and selectivity. We have thus aimed at developing more specific and potent inhibitors of CBS and CSE towards H2S production. We used a semi-rational strategy which combines two highly successful routes, the structural based and the fragment-based approaches and went all way from drug design to ex vivo and in vitro studies. We identified an oxothiazolidine derivative which showed maximal inhibitory effects in tissue and proved to be more efficient than PAG and completely selective for CSE by metabolomics assays on the purified enzymes. As compared to PAG, the new compound has a significant difference: both compounds inhibit CSE from producing pyruvate, ammonia and hydrogen sulfide, but surprisingly, in the presence of PAG, CSE remains able to consume the substrate L-Cys and produces cystine and lanthionine. This is an important result previously under looked. In contrast, compound 2a inhibits the enzyme completely, keeping the concentration of substrate unaffected. This finding suggests the possibility of using lower doses of compound 2a compared to PAG, consequently also reducing all possible adverse effects. PAG is also a suicidal inhibitor while 2a does not bind irreversibly offering a further advantage.

In conclusion, it is clear that before using the new compound for human treatment, it will be essential to carry out a careful pharmacological study to assess whether the new compound is toxic or is associated with any unwanted side effects. For the time being, we are nevertheless confident that identification of this highly selective CSE inhibitor will help to better define the role of CSE vs CBS in the pathophysiology of the diseases where a role
for the H$_{2}$S pathway has been proposed. Furthermore, the development of such agents, particularly water-soluble, will allow us to evaluate the cross-talk of H$_{2}$S pathways with other relevant pathways (e.g. NO, COX).

**Experimental section**

**Compound synthesis and their identification.** The synthesis strategy is detailed below and summarized in Figures S1 and S2 of Suppl. Information. The identity of the resulting products was proven using mass spectrometry, $^{1}$H and $^{13}$C NMR. Melting points, determined using a Buchi Melting Point B-540 instrument, are uncorrected and represent values obtained on recrystallized or chromatographically purified material. $^{1}$H-NMR spectra for compound characterization were recorded on a Varian Mercury Plus 400 MHz instrument and in DMSO solutions. Chemical shifts are reported in ppm using Me$_{4}$Si as internal standard. The following abbreviations are used to describe peak patterns when appropriate: s (singlet), d (doublet), t (triplet), q (quartet), qt (quintet), dd (double doublet), m (multiplet). Mass spectra of the final products were performed on API 2000 Applied Biosystem mass spectrometer. Elemental analyses were carried out on a Carlo Erba model 1106. Analyses indicated by the symbols of the elements were within ±0.4% of the theoretical values. All reactions were followed by TLC, carried out on Merck silica gel 60 F254 plates with fluorescent indicator and the plates were visualized with UV light (254 nm). Preparative chromatographic purifications were performed using silica gel column (Kieselgel 60). Samples were dried over Na$_{2}$SO$_{4}$ and concentrated with a Buchi R-114 rotary evaporator at low pressure.

**Isolated mouse aortic ring assay.** The animals were sacrificed according to the regulations of University of Naples. Thoracic aorta from CD-1 mice was rapidly dissected and cleaned from fat and connective tissue. Animal care was in accordance with Italian and European regulation on the protection of animals used for experimental and other scientific purposes. For endothelium-denuded rings, the endothelial layer was removed by gently rubbing the internal surface of the vascular lumen. Rings of 1.5–2 mm length were cut and mounted in isolated organ baths filled with gassed Krebs solution containing (in mol/L) NaCl 0.118, KCl 0.0047, MgSO$_{4}$ 0.0023, KH$_{2}$PO$_{4}$ 0.0012, CaCl$_{2}$ 0.0025, NaHCO$_{3}$ 0.025 and Glucose 0.010 (95% O$_{2}$ + 5% CO$_{2}$) at 37 °C linked to isometric force transducers (UgoBasile, Comerio, Varese, Italy). Rings were initially stretched until a resting tension of 1.5 g was reached and equilibrated for at least 45 min during which tension was adjusted to 1.5 g. The bathing solution was periodically changed. In each experiment, rings were contracted with L-phenylephrine (PE) at 3μmol/L and then at 1 μmol/L until response was reproducible. To verify the integrity of the endothelium, aortic rings were contracted with PE (1 μM), once plateaus was reached and a cumulative concentration-response curve to commercially available precursors or the newly synthesized compounds (10 nM – 300μM) was performed. The rings were contracted with PE (1μM) and a cumulative concentration-response curve to L-cysteine (1μM - 3 mM) was performed. All data are expressed as means ± SEM. Statistical analysis was performed by two-way ANOVA followed by Bonferroni multiple comparison test. Differences were considered statistically significant when P-value was <0.05. GraphPad Prism software (GraphPad Software, Inc., La Jolla, CA, USA) was used for all the statistical analysis. IC$_{50}$ values were determined ex vivo, using at least three different concentrations of the tested compounds. All the experiments were performed in triplicate. The obtained data, expressed in semilogarithmic scale, were used by the ED$_{50}$ Plus v1.0 online software to get the IC$_{50}$ values. All experimental protocols were approved by the licencing committee of University of Naples.

**Estimate of the Lipophilic parameters.** To estimate the plasma membrane crossing properties of the new inhibitor, the log $P^D$ values, i.e. partition coefficients n-octanol/aqueous phase of the neutral form of selected compounds, were calculated (c log$P$ and in vitro) modifying previous protocols$^{17}$. For the ex vivo assay, thoracic aorta was dissected, placed in sterile phosphate buffer solution and cleaned of fat and connective tissue. The aortic rings were homogenized in a lysis buffer (potassium phosphate buffer 100 mM pH 7.4, sodium orthovanadate 10 mM and protease inhibitor) and treated with the different concentrations of compounds for 30 minutes and added in a reaction mixture (total volume 500μL) containing peridoxal-5′-phosphate (2 mM, 20μL), L-cysteine (10 mM, 20μL) and saline (30μL). The reaction was initiated by transferring tubes from ice to a water bath at 37 °C. ZnAc$_{1}$ (1%, 250μL) was added to trap evolved H$_{2}$S followed by TCA (10%, 250μL) and N,N-dimethyl-p-phenylelenediamine-sulfate (20μM, 133μL) in 7.2 M HCl and FeCl$_{3}$ (30μM, 133μL) in 1.2 M HCl were added. After 20 minutes absorbance values were measured at a wavelength range of 670 nm. All samples were assayed in duplicate and L,S concentration was calculated against a calibration curve of NaHS (3.12–250μM). Basal release of H$_{2}$S was measured in absence of L-Cys. Vehicle indicates the used buffer, containing exogenous L-Cys.

For the in vitro assay, each reaction mixture (250μL) consisted of 5μg of human CSE, 0.01 mM PLP, 1 mM L-cysteine and 50 mM sodium phosphate pH 8.2 buffer. The inhibitors, at different concentrations, were added to each reaction 15 min before L-cys was added to the solution. Reaction was initiated by transferring the Eppendorf tubes, tightly paraffilmed, from ice to a 37 °C water bath. After 30 min of incubation, the evolved H$_{2}$S was trapped via addition of ZnAc (1% w/v, 100μL) and proteins were precipitated via addition of TCA (10% w/v, 100μL). Subsequently, N,N-dimethyl-p-phenylelenediamine-sulfate in 7.2 M HCl was immediately followed by addition of FeCl$_{3}$ in 1.2 M HCl, for development of methylene blue. The amount of H$_{2}$S produced was determined via absorbance measurements of the centrifugal supernatants. The absorbance of the resulting solution was measured at 670 nm and the amount of H$_{2}$S was calculated against a calibration curve of standard H$_{2}$S solutions.

The determination of IC$_{50}$ on purified CSE was done by a concentration-response curve obtained by plotting the percentage of inhibition versus the logarithm of inhibitor concentrations. The IC$_{50}$ values were estimated from the resulting graph using GraphPad Prism v6.07 (GraphPad Software, Inc. La Jolla, CA).
Protein expression and purification. E. coli BL21 (DE3) Codon Plus was used as the host strain to express recombinant human CSE or CBS. CSE cDNA was cloned into pGEX-4T3 and CBS into pGEX-Kg to create N-terminal GSH-S-transferase (GST) fusion proteins. The expression vectors were transformed and plated on LB-agar plates, supplemented with ampicillin (100 µg/mL). The expression and purification of CBS and CSE was adapted from previously described protocols. Colonies of pGEX-Kg/GST-CBS or pGEX-4T3/GST-CSE transformed BL21 cells were separately inoculated into 100 mL of steam-autoclaved Luria-Bertani (LB) broth supplemented with 100 µg/mL of ampicillin (LB-Amp100), and incubated at 37 °C overnight shaking. After induction and harvesting, the lysates were centrifuged at 16000 rpm for 30 min at 4 °C using a Beckman JA20 rotor. The soluble fractions were loaded onto Glutathione sepharose 4B columns equilibrated with binding buffer PBS. The flow-throughs were collected, and the beads washed. Proteins attached to the columns were eluted with five column volumes of elution buffer (50 mM Tris–HCl, 10 mM reduced GSH, 5mM DTT, pH 8.0), dialyzed and concentrated in 10 mM sodium phosphate buffer pH 8.2 and DTT 1 mM. The eluates were passed through a Superdex-200 column connected to the AKTA purifier FPLC equilibrated in the same buffer at a flow rate of 1 mL/min. Fractions containing the enzymes were identified by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE), pooled together and concentrated using an ultra-centrifugal filter (Millipore Amicon Ultra-4 10000 MWCO) at 4 °C.

Structural characterization of recombinant enzymes. Far-UV CD measurements were performed on a J-715 spectropolarimeter (JASCO, Oklahoma City, OK) equipped with a PTC-348 Peltier temperature control system using an enzyme concentration of 4 µM in 10 mM sodium phosphate pH 8.2 buffer. The spectra were recorded in quartz cuvettes (Hellma, Plainview, NY) with path-lengths of 1 mm and averaging five separate wavelength scans on independent protein preparations to ensure reproducibility. Baseline correction was performed by subtraction of the appropriate buffer spectrum. Adsorbance spectroscopy was carried out on a Varian Cary 50 Bio UV-Visible Spectrophotometer. Measurements were performed in a 1 cm cuvette cell and monitored spectroscopically from 650nm to 250 nm.

Enzymologic analysis. Kinetics studies were performed using an adapted version of a spectrophotometric assay described by Stipanuk and Beck. Assay mixtures (250 µL) contained 5 µg of purified CSE, 2 mM PLP and L-cysteine (1–4 mM) previously incubated at 37 °C. For each L-cysteine substrate concentration, the amount of H₂S was measured every 3 min over 30 min. Zinc acetate (1% w/v) was added to trap the produced H₂S gas. The enzymatic reaction was stopped by the addition of 10% (w/v) trichloroacetic acid. N,N-dimethyl-p-phenylendiamine dihydrochloride dye (20 mM) and 30 mM FeCl₃ were added and the absorbance at 670 nm was measured using a microplate reader (Thermo Scientific). All determinations were carried out in triplicates. The amount of H₂S produced was calculated against a calibration curve of sodium hydrosulfide (0–250 µM). The initial reaction velocity, V₀ (units/mg of human CSE, where 1 unit is the amount of enzyme that catalyzes the reaction of 1 µmol of H₂S per minute) was plotted against the cysteine substrate concentration [S]. Rates were fit with the Michaelis-Menten equation using GraphPad Prism software version 7 for Windows (San Diego, USA).

NMR spectroscopy. Samples were composed of 15N,13C-L-cysteine (125 µL, 1 mM), unlabeled L-homocysteine (125 µL, 1 mM) (added only for the CBS time course as co-substrate), DTT (0.5 µL, 1M), D₂O (50 µL), and enzyme (100 µL, 110 µM) in 10 mM sodium phosphate pH 8.2 and filled to a final volume of 500 µL. NMR experiments were performed at 300 K on a Bruker AVANCE spectrometer operating at 800 MHz with TCI Cryoprobes. Controls were run excluding either the enzyme or the tested compounds pre-incubated with each enzyme. For each sample, pseudo-2D experiments were collected in which the first increment of a 1H HSQC spectrum was acquired with 5 minutes repetition time for a total of 60 spectra, using 15N,13C, L-cysteine and unlabeled L-homocysteine as CBS substrate and 15N,13C, L-cysteine as a unique substrate for CSE enzyme. 2D 1H,13C HSQC spectra were acquired before and after the enzyme reaction was completed.

HPLC-ESI-MS/MS Analyses. Chromatographic analyses were performed using a HPLC Agilent 1100 series equipped with an auto-sampler coupled online with an Agilent LC-MSD SL quadrupole ion trap. The liquid chromatography separation was carried out on a Supelco Discovery-C8 column (250 × 3.0 mm, particle size 5 µm) eluted isocratically with 0.1% formic acid in water, at a constant flow rate of 0.1 mL/min. The injection volumes were 10–20 µL for each sample. Electrospray ionization (ESI) was in positive ion mode, with nitrogen as the nebulizing and drying gas using a pressure of 30 psi, a drying temperature of 350 °C and drying gas at 7 L/min. The ion charge control (ICC) was applied with the target set at 30000 and maximum accumulation time at 20 ms. Optimization of instrumental parameters was performed by continuous infusion of 5 µM standard solution in 0.1% formic acid. Mass cutoff and amplitude were optimized to obtain the most efficient MS/MS transitions from the positively charged precursor ion [M + H]⁺ to the fragment ions. Multiple reaction monitoring (MRM) was used with the MS/MS transitions utilized for the quantitation of analytes of 209.1→120 for lanthionine, 241.0→152 for cystine, 122.0→105 for cysteine. Quantitation was achieved by comparison with calibration curves obtained from standard solutions. Measurements were performed from the peak area of the extracted ion chromatogram (EIC). Retention times and peak areas of the monitored fragment ions were determined by the Agilent software Chemstation version 4.2.

Structural analysis. Structural analysis and manual docking was possible with the support of the software PYMOL and MAESTRO (Schroedinger, LLC, New York, NY). The position of PAG in the X-ray structure 3cog was taken as pose of reference.
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Author Contributions
A.C., B.S. and F. Frencentesi carried out the synthesis. A.C. also performed the metabolomics experiments by NMR and produced the recombinant enzymes. G.K. provided technical expertise on the NMR data acquisition and analysis. E.M., E.P. and F. Fiorino characterized the compounds. M.B. and G.Cirino performed the experiments ex vivo. A.P., V.S. and G. Caliendo directed the structural and synthetic research respectively. A.C. and A.P. wrote the manuscript. L.S. performed the Mass spectrometry data. G.P. performed the in silico analysis.

Additional Information
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