Generation of controllable gaseous H$_2$S concentrations using microfluidics

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Abstract

Hydrogen sulfide (H$_2$S) plays an important role as an intercellular and intracellular signaling molecule, yet its targets are not well understood. As a molecule it easily evaporates and it is hard to acquire stable concentration for in vitro studies, constituting a major problem for the field to identify its downstream targets and function. Here we develop a microfluidic system that can provide consistent and controllable H$_2$S levels in contrast to the current method of delivering large bolus doses to cells. The system relies on the permeability of H$_2$S gas through a polydimethylsiloxane thin membrane. A hydrogen sulfide donor, sodium hydrosulfide, is perfused in the microchannels below the gas permeable membrane and gaseous H$_2$S diffuses across the membrane, providing a stable concentration for up to 5 hours. Using electrochemical sensors within 3 ppm range, we found that H$_2$S concentration was dependent on two parameters, the concentration of H$_2$S donor, sodium hydrosulfide and the flow rate of the solution in the microchannels. Additionally, different H$_2$S concentration profiles can be obtained by alternating the flow rate, providing an easy means to control the H$_2$S concentration. Our approach constitutes a unique method for H$_2$S delivery for in vitro and ex vivo studies and is ideally suited to identify novel biological processes and cellular mechanisms regulated by H$_2$S.

Introduction

Hydrogen sulfide (H$_2$S) is a colorless gas with a pungent smell of ‘rotten eggs’. In humans, hydrogen sulfide (H$_2$S) is known to be a toxic gas that is lethal in high concentrations over 500 ppm. However, emerging research demonstrates that H$_2$S is generated endogenously in low concentrations and serves as a signalling molecule which regulates cellular homeostasis. H$_2$S, thus, belongs to the family of ‘gasotransmitters’ along with nitric oxide (NO) and carbon monoxide (CO). Additionally, its biological role is evident in plants and bacteria, either as a bi-product or a power source.
H₂S is endogenously produced in mammals mainly by three enzymes, cystathionine – β – synthase (CBS), cystathionine-γ-lyase (CSE or CGL) and 3-mercaptopyruvate sulfur transferase (3-MST) which are located in different organs.³ H₂S is additionally produced non-enzymatically, but is considered negligible in comparison to the enzymatic production.⁴ In 1996 H₂S was firstly introduced as a signalling molecule and since then, H₂S has attracted increased attention for its potential therapeutic effects in mammals. For example, it has been found that H₂S is cytoprotective with antioxidant properties,¹¹ vasodilatory,¹²,¹³ angiogenic,¹⁴,¹⁵ and neuromodulatory,¹⁶ and its repertoire of biological functions is steadily expanding.¹,²,¹⁷–¹⁹

Hydrogen sulfide gas can be diluted in liquid at a ratio 1mg/L=717μM [STP]. In an aqueous solution hydrogen sulfide follows the equilibrium:

$$H_2S \rightleftharpoons H^+ + HS^- \rightleftharpoons H^+ + S^{2-}$$

With pKa varying depending on the temperature. At room temperature, the pKa for the left equilibrium is almost 7 while the pKa for the second equilibrium varies in different studies which is over 12. Hydrogen sulfide in its protonated form evaporates from the solution in a gaseous form, a phenomenon that causes the pH to increase. Therefore, the equation is constantly driven to the left, diminishing the concentration of H₂S in the solution.²⁰,²¹

Using different measurement techniques, a discrepancy in in vivo H₂S concentrations has been reported which is mostly attributed to the techniques themselves¹,²² with older studies reporting elevated H₂S endogenous concentrations.²³,²⁴ Additionally, for in vivo measurements it is assumed that H₂S is rapidly metabolized within a tissue, thus the concentrations vary depending on the measuring location.²⁰,²⁵ Thus the reported physiological concentrations vary from the submicromolar range to 200 μM.²⁶

A key challenge for H₂S research is typical H₂S donors provide H₂S molecules in an unstable and uncontrolled manner. In most of the studies, hydrogen sulfide is generated by inorganic salts, either sodium hydrosulfide (NaSH) or sodium sulfide (Na₂S), or by purging a solution with gaseous H₂S. When these solutions are added in vivo or in cell cultures, a certain concentration of H₂S is achieved, which sometimes can reach cytotoxic levels.²⁷,²⁸ This concentration of H₂S is achieved for seconds as it degrades with time mostly by volatilization, and by oxidation in a pH manner, where oxidation increases for physiological pH.²,²⁵,²⁹ DeLeon et al. showed the significance of this fact for three different cases, a cell culture plate, a myograph bath, and a Langendorff isolated heart apparatus. It is shown that the concentration of H₂S in a Na₂S solution declines rapidly with a half-life in the order of minutes. This loss of H₂S is attributed mostly to volatilization and minimally to oxidation.²⁵

Recently, several compounds that slowly release hydrogen sulfide have been reported. The compound most commonly used is GYY 4137 which is commercially available. GYY 4137 releases hydrogen sulfide slowly for several minutes, but the production rate cannot be controlled and the release increases with time, therefore, high concentrations of H₂S can be achieved after a few days.³⁰ The only study that partial control of the release of hydrogen
sulfide is shown in a pH-controlled manner using the slow releasing chemical JK-1 – JK-5.\textsuperscript{31} Using this method, the concentration of H$_2$S can be controlled by changing the pH, unfortunately this is not applicable for biological applications. Additionally, it is reported that other impurities of the donor or the chemical properties of the donor itself might implicate the already complicated phenomenon.\textsuperscript{27,32} For example, impure NaSH can be a source of solid sulfur in astrocyte studies with a developing Ca$^{2+}$ influx due to sulfur.\textsuperscript{33} The combination of these factors constitute a major problem for the field and thus the exact mechanism of H$_2$S activity is still unclear.\textsuperscript{29,32}

Recently, Faccenda et al. showed that H$_2$S penetrates through a thin polydimethylsiloxane (PDMS) membrane allowing a continuous measurement in which the solutions are not influenced by impurities of the donor as they cannot penetrate the PDMS membrane. In this study the donor, either Na$_2$S or CSE, is inserted in a well and the H$_2$S molecules that have diffused through the 100 μm membrane can be measured with different methods.\textsuperscript{34}

Microfluidic devices offer new approaches to stably generate environment gas landscapes such as defined stable oxygen concentration environments for both cell cultures and brain tissue ex-vivo studies. These devices rely on a microfluidic network that is bonded to a thin PDMS membrane (figure 1). Previously our lab has published many studies using similar devices to deliver controlled oxygen landscapes to cells and tissues\textsuperscript{35–41} Similarly, we found that these devices can be used to establish environments of different concentrations of hydrogen sulfide. As the solution of NaSH is introduced in the microfluidic network, the gaseous H$_2$S diffuses across the PDMS membrane and a steady state concentration is achieved. This method delivers H$_2$S in the reservoir for a long period of time (hours), in comparison to a conventional method, where a solution of NaSH is directly injected in the reservoir and diminishes rapidly (minutes) (figure 2). The established concentration of a specific pH NaSH solution flowing in the microfluidic network provides hydrogen sulfide depending on two main factors: the concentration of the NaSH solution and the flow rate of the liquid. In order to quantify the method, we measured the mean value of H$_2$S in the reservoir for 5 hours and we found a linear relationship with the flow rate (figure 3). Additionally, the concentration of hydrogen sulfide can be modulated by alternating the flow rate (figure 4). Finally, different microfluidic networks of different areas were compared in order to further validate the phenomenon (figure 5). Using these microfluidic devices different concentrations of H$_2$S are achieved where only pure gaseous H$_2$S molecules are provided in the system and can be controlled.

**Methods & Materials**

**Fabrication of microfluidic device**

The microfluidic device consists of three different layers, the microfluidic channels superimposed by a 100 μm thick PDMS membrane and a reservoir, where liquid/media/solution can be introduced for different studies. The microchannels are fabricated using typical soft lithography techniques. Specifically, photosensitive polymer SU-8 resist was spun on a silicon wafer and exposed to UV light under a photomask (Fineline Imaging, Colorado Springs, CO). The design was then developed and polydimethylsiloxane (PDMS) (Sylgard 184, Dow Corning, Midland, MI) was poured on top followed by curing at 85°C.
for two hours. The 100 μm PDMS membrane was fabricated by spinning uncured PDMS at 800 rpm and then curing at 85° C for 45 minutes. A PDMS slab of 6.5 mm thickness was punched with 22.6 mm diameter holes that serve as potential liquid reservoirs, and 1 mm holes that served as inlet and outlet holes for the microchannels. The microfluidic channels were bonded on the PDMS membrane and the PDMS piece was bonded on top of the membrane using Corona plasma treatment (Electro Technic Products, Chicago, IL).

Solution preparation

A solution of distilled water was purged with nitrogen for at least 1 hour in an aspirator bottle with a bottom hose outlet of 1 L (Cole Parmer, Vernon Hills, IL). As soon as the sodium hydrosulfide (NaSH) (Sigma Aldrich) was added to the solution, the bottle was corked with a two-hole rubber stopper, and ¼” barb male luer lock fittings attached to a large bore stopcock were placed in the holes of the stopper. The solution was extracted from the bottle by opening a stopcock attached on a tube connected to the bottom hose outlet. This minimizes air introduced into the bottle, minimizing both volatilization and oxidation of H₂S gas. The solution was then titrated into 220 mL of a weak sulfuric acid solution (2 μL in 220 cc of DI water) keeping the pH level below 3.7, within a 250 cc glass bottle sealed similarly with a rubber stopper. Each specific concentration of NaSH solution was loaded to the syringes via the luer lock stopcock. For each experiment a fresh solution was prepared as the concentration of H₂S in this solution diminishes over time.

Experimental setup

NaSH solution was loaded in plastic 60 mL syringes (60 cc luer lock BD, Fischer Scientific, Hanover Park, IL). A syringe pump (PHD 2000, Harvard Apparatus, Holliston, MA) was used to drive the fluid through a needle connected with a thin tygon tubing (tygon microbore tubing, 0.020” × 0.060” OD, Cole Parmer, Vernon Hills, IL). The outer diameter of the tubing is bigger than the inlet and outlet holes, securing a leak free flow. All the experiments were carried out in ambient air under a fume hood.

Validation

The detection of gaseous H₂S concentration on top of the membrane was achieved using electrochemical sensors (Alphasense, Essex, UK). The electrical signal is acquired at a sampling rate of 1 Hz with a data acquisition system (FS-1608, Measurement Computing, Norton, MA) using DAQami software. The sensors were calibrated using premixed gases of H₂S in nitrogen (Cal Gas Direct Incorporated, Huntington Beach, CA) (figure S1).

Results and discussion

In order to characterize our method, the electrochemical sensors were placed on top of the membrane of our microfluidic devices and H₂S gas concentration was measured. The microfluidic network consists of a serpentine of rectangular cross section of 800 μm width and 100 μm height, while the microfluidic channels were 800 μm apart for most of the studies aside from those investigating other microchannel geometries. It is shown that the concentration of H₂S increases when NaSH solution is introduced in the microfluidic network. The flow rate is the main factor that influences the rate of increase of the
concentration to reach a steady equilibrium. The levels reach equilibrium within the first hour of the initial flow and remains constant with time (figure 2). In comparison, in figure 2, \( \text{H}_2\text{S} \) concentration of dispensing a 20 \( \mu \text{M} \) NaSH PBS solution in an open well is depicted. This measurement was taken by placing the electrochemical sensor over the well supported and sealed by a PDMS piece of the same diameter as the sensors to maintain consistent position and to contain diffused gas in the reservoir. With the conventional method, \( \text{H}_2\text{S} \) concentration is diminished within 30 minutes, while with our method the concentration of \( \text{H}_2\text{S} \) increases and remains constant. Additionally, using our method three different molarities, 20 \( \mu \text{M} \) (figure 3a), 50 \( \mu \text{M} \) (figure 3b), and 100 \( \mu \text{M} \) (figure 3c) were tested for different flow rates for at least 6 hours. These equilibrium concentrations for each molarity solutions and different flow rates were averaged for 5 hours in the steady \( \text{H}_2\text{S} \) concentration (from the first to the sixth hour), shown in figure 3d. It is shown that the concentration has a linear relationship \( (R^2=0.99) \) with the flow rate of NaHS solution, while the slope of the line is different for higher concentration solutions of NaHS. In specific the slope is 0.017 ppm/\( \mu \text{L}/\text{min} \) for 20 \( \mu \text{M} \), 0.041 ppm/\( \mu \text{L}/\text{min} \) for 50 \( \mu \text{M} \), and 0.069 ppm/\( \mu \text{L}/\text{min} \) for 100 \( \mu \text{M} \). Error bars show the standard deviation of three different devices. The deviation is likely caused by subtle shifts of the sensor as the reservoirs' diameter was slightly bigger than the sensors' diameter. The concentration of \( \text{H}_2\text{S} \) has a linear relationship \( (R^2=0.99) \) with the flow rate of NaSH solution, while the slope of the line correlates with the concentration of NaSH.

By simply changing the flow rate of the syringe pump, our method allows for the concentration of \( \text{H}_2\text{S} \) to be precisely alternated during the same experiment. As a demonstration of hypothetical \( \text{H}_2\text{S} \) exposure experiments, two cycles of different flow rates of a 50 \( \mu \text{M} \) NaSH solution are alternated every 45 minutes (figure 4). On the first, the flow rate is alternated from 20 to 30 to 40 to 50 to 60 to 70 \( \mu \text{L}/\text{min} \) (figure 4a) and on the second, the flow rate is changed from 60 to 40 to 20 \( \mu \text{L}/\text{min} \) (figure 4b). At the end of each cycle the flow rate returns to the initial value according to the program that was defined with the syringe pump. For drastic changes in flow rate, the 45 minute interval is not enough for the concentration to reach equilibrium concentration. This occurs in the first cycle when the flow rate returns from 70 \( \mu \text{L}/\text{min} \) to 20 \( \mu \text{L}/\text{min} \) and the concentration of \( \text{H}_2\text{S} \) remains above the concentration that corresponds to 20 \( \mu \text{L}/\text{min} \). Similarly, on the second cycle the concentration does not reach equilibrium for the initial flow rate of 60 \( \mu \text{L}/\text{min} \). Nevertheless, this demonstrates the ability to control \( \text{H}_2\text{S} \) levels by varying flow rates through the microchannels.

The dimensions of the microchannel and how they related to \( \text{H}_2\text{S} \) values in the reservoir were next investigated by changing the geometry of the serpentine microfluidic network. We compared the concentration profiles for microchannels with width ratio \( W_c/W_s=800/800 \), with those of \( W_c/W_s=1200/200 \) for microchannel height of 100 \( \mu \text{m} \), where \( W_c \) is the width of the microchannel and \( W_s \) is the width of the spacing between the microchannels. Surprisingly, the results are not significantly different for all concentrations of NaSH (Figure 5a). In addition, the serpentine with \( W_c/W_s=800/800 \) was compared to a 400 \( \mu \text{m} \) height and the results were almost identical (Figure 5a). Thus, we conclude that the only factors that significantly influence \( \text{H}_2\text{S} \) concentration are NaSH concentration and the flow rate,
indicating that microchannels of different geometries can be used to generate similar concentrations within the given range of dimensions tested.

**Conclusion**

To date, studying the impact of hydrogen sulfide in biological systems has been stymied by poor control over its concentration in solution. Using our microfluidic approach we show that physiological H$_2$S concentrations of up to 3 ppm can be achieved over many hours. In addition, compared to existing technology (or procedures) which deliver deprotonated HS$^-$, our method uses an acidic solution, which provides a more consistent H$_2$S concentration with time as the solution is separated from the reservoir by a PDMS membrane. By simply changing the flow rate, our microfluidic approach permits control of the H$_2$S concentration within the same experiment. To our knowledge this is the first microfluidic device that allows for the precise and stable control of gasotransmitter levels. This method can be adapted for in vitro and ex vivo studies, and could also enable the use of other gasotransmitters NO and CO, which could be delivered by flowing the gas in the microchannels, either individually or by combining different signalling gases. The use of microfluidic devices to study H$_2$S signalling has the potential to significantly advance the field by providing precise control of its concentration and identify novel H$_2$S-dependent processes in biological systems where current methods of H$_2$S control are lacking.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

This work was supported by National Science Foundation 1253060, DTE and NIH P01-HL60678.

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Fig. 1.
The microfluidic device assembly. The microchannel layer is bonded on a thin H$_2$S permeable membrane. The adjacent side of the membrane is bonded to the reservoir layer that the inlet and outlet holes of the microchannels are punched. On the right, the microchannels in a serpentine shape have a width $W_c$ which are separated by a spacing with width $W_s$. The reservoir is outlined with the black circle while the grey circle in the middle denotes the measuring region of each sensor.
Fig. 2.

a. The concentration using the microfluidic method is increasing with time until it reaches a stable concentration. The NaSH solution introduced in the syringe pump is 20 μM and different concentrations can be achieved by flowing the NaSH solution at different rates. Conventionally, dispensing a 20 μM NaSH PBS solution in a well the concentration spikes up in the beginning and it is totally lost within approximately 70 minutes. b. Graphical representation of the conventional introduction of H2S in a cell culture well plate. The H2S molecules are c. Graphical representation of the microfluidic method of H2S. The H2S molecules are introduced in the well as the solution is flown within the microchannels.
Fig. 3.
H₂S concentration for different flow rates for a. 20 μM b. 50 μM and c. 100 μM NaSH (error bars shown every hour). The average between 1-6 hours for different flow rates is depicted in d.
Fig. 4.
Alternating H$_2$S concentrations can be obtained by alternating the flow rate of a 50 μM NaSH solution in the microchannels. Two cycles of flow rates with an interval of 45 minutes are shown.
Fig. 5.
Mean H$_2$S concentration in the steady concentration for different flow rates for a serpentine with (a) wide microchannels (width = 1200 μm, spacing = 200 μm and height = 100 μm) (b) high height microchannels (width = 1200 μm, spacing = 800 μm and height = 400 μm) in comparison to the model serpentine microchannels (width = 800 μm, spacing = 800 μm and height = 100 μm).