There is great interest in developing a low-power gas sensing technology that can sensitively and selectively quantify the chemical composition of a target atmosphere. Nanomaterials have emerged as extremely promising candidates for this technology due to their inherent low-dimensional nature and high surface-to-volume ratio. Among these, nanoscale silicon is of great interest because pristine silicon is largely inert on its own in the context of gas sensing, unless functionalized with an appropriate gas-sensitive material. We report a chemical-sensitive field-effect transistor (CS-FET) platform based on 3.5-nm-thin silicon channel transistors. Using industry-compatible processing techniques, the conventional electrically active gate stack is replaced by an ultrathin chemical-sensitive layer that is electrically nonconducting and coupled to the 3.5-nm-thin silicon channel. We demonstrate a low-power, sensitive, and selective multiplexed gas sensing technology using this platform by detecting H2S, H2, and NO2 at room temperature for environment, health, and safety in the oil and gas industry, offering significant advantages over existing technology. Moreover, the system described here can be readily integrated with mobile electronics for distributed sensor networks in environmental pollution mapping and personal air-quality monitors.

**INTRODUCTION**
To date, the gas sensor market has primarily been dominated by applications in oil and gas, automotive, and other process industries. This may no longer hold true, especially with the recent boom in low-power sensor technologies associated with the Internet of things. New applications in personal air-quality monitoring, pollution tracking, and breath analysis for preventive health care have created a growing demand for low-power chip-scale gas sensors. Although there are several categories of gas sensors, only three have indicated potential for chip-scale miniaturization and low-power operation, namely, resistive ceramic (1) and metallic (2) sensors, nanoelectromechanical systems (3), and field-effect transistors (FETs) (4). Of these, resistive ceramic gas sensors have been explored extensively in the past. These gas sensors are typically based on transition metal oxides that operate on the principle of resistance change due to gas-oxide redox reactions at high temperatures. Furthermore, resistance change is affected in the same material undergoing chemical change. Selective gas sensing using this technology is quite challenging because it is dependent on the temperature of gas-oxide interaction. By far, the FET technology has shown significant potential in overcoming limitations of size, power consumption, and sensitivity. The FET technology works on the principle of channel current modulation due to gas interaction in the linear amplifying region of a transistor. Strong and growing interest in this technology is very clear, especially with the advent of materials such as carbon nanotubes (5–9), semiconducting nanowires (4, 10–15), organic semiconductors (16–18), graphene (19), and transition metal dichalcogenides (20, 21). Silicon nanowires are of particular relevance to this work, where significant progress has been made in demonstrating them for gas sensing (4) and biosensing (11), indicating nanoscale silicon as a model material system for sensitive and selective sensors.

Here, we demonstrate chemical-sensitive FETs (CS-FETs) as a highly sensitive, selective, and low-power gas sensing platform, where a 3.5-nm-thin silicon channel on an insulator is the enabling technology (Fig. 1). Today, these ultrathin silicon channels (22) are used as fully depleted transistors to drive the current generation of low-power digital electronics (23). The CS-FET sensors operate at room temperature and are chemically gated with a chemical-sensitive layer, which upon exposure to a target gas, induces a channel threshold voltage shift. Room temperature gas sensors may typically suffer from effects such as poisoning and slow recovery to baseline after gas exposure. To address this, CS-FET gas sensors are integrated with local Au/Cr micro heaters on a silicon-on-insulator (SOI) platform. These microheaters are activated only after a gas detection event to assist in quick recovery to baseline and do not contribute in any way to the actual sensing mechanism of the CS-FET. Figure 1A depicts a perspective-view optical microscope image of a fabricated CS-FET sensor, whereas Fig. 1 (B and C) depicts the array concept of individually functionalized CS-FETs and a pictorial representation of a single device, respectively.

**RESULTS**
Although the CS-FET is a generic gas sensing technology that can be configured for different applications by selection of proper sensing films, the targeted application for this work is in the detection of H2S, H2, and NO2 for personnel and environment safety in the oil and gas industry. H2S is a colorless, flammable, and corrosive gas that poses significant safety concerns in this industry. It has a distinct rotten egg smell at very low concentrations [approximately 0.01 to 1.5 parts per million (ppm)] and becomes odorless and toxic beyond this concentration range with instant death at 800 ppm. In the United States, the National Institute for Occupational Safety and Health (NIOSH) has
characterized a 60-min exposure to 100 ppm of H₂S as an immedi-
ately dangerous to life or health (IDLH) condition (24). Because of
this, the H₂S sensing range of interest in the oil and gas industry is 0
to 100 ppm. The state of the art uses a combination of resistive ce-
ramic and electrochemical H₂S sensors, with response times (t₉₀) of
around 15 to 120 s and detection limits of 0.4 to 20 ppm, depending
on the manufacturer. Similar to H₂S, H₂ is also a major concern in
these industries, where it is used in petroleum refining that involves
desulfurization. NIOSH characterizes H₂ as a flammable gas with a
lower explosion limit of 4% by volume in air (24). Catalytic-type
as well as optical infrared sensors are currently used for detecting
H₂ leaks with response times of around 30 s or less and detection
ranges of 0 to 100%. In addition to H₂S and H₂, NO₂ emissions from
such industries are becoming increasingly important to monitor
because they are common indicators of air quality. NO₂, which is
a respiratory irritant, is a reddish brown gas with a pungent odor
that is released during the burning of complex hydrocarbons. The
IDLH limit for NO₂, as specified by NIOSH, is 20 ppm for a 30-min
exposure (24). The sensing range of interest is also 0 to 100 ppm, and
the state of the art uses both resistive ceramic and electrochemical gas
sensor technology. Depending on the specific manufacturer, response
times are around a minute or less with detection ranges of 0.2 to 20 ppm.

The technologies described above are typically very bulky in size
and weight as well as power-hungry, rendering them mostly as fixed
point detectors. It is also quite common for these sensors to detect false
positives due to cross-sensitivity from interfering gases. The effective
sensing of H₂S is of importance to personnel health and safety in the
oil and gas industry, where the current standard practice is for person-
nel to carry bulky and power-hungry monitors to detect the presence
of the mentioned gases. In the following sections, we demonstrate that
the CS-FET gas sensor technology can offer significant advantages in
terms of sensitivity, selectivity, size, and power consumption that can
augment or potentially replace existing technology.

CS-FETs are gated chemically with gas-specific ultrathin sensing
layers (sub–5 nm) that are electrically nonconductive. Interactions
with a target gas induce changes in the sensing layer work function,
which in turn causes large modulations in the current (I_D) flowing
through the ultrathin silicon channel, leading to highly sensitive detection
events. Unlike other highly sensitive nanomaterials, the CS-FET silicon channel on its own is comparatively inert and unresponsive un-
less functionalized with appropriate sensing layers, enabling selective
detection (see fig. S1). Depending on the nature of the target gas,
sensing layers can be composed of metal or metal alloys, transition
metal oxides, or even polymers. Here, mono- and bimetallic compo-
sites are used as chemical-sensitive layers. The underlying transistors
in the CS-FET platform are fabricated using a complete top-down ap-
proach, which is fully compatible with silicon complementary metal-
oxide semiconductor (CMOS) processing (23). This enables scalable
batch fabrication of large ultrathin channel transistor arrays, where
each transistor can be functionalized with different gas-specific sensing
layers (Fig. 1B), paving the way to single-chip multiplexing for selective
gas sensing. Selectivity is a major challenge in any gas sensing tech-
nology. It is the property by which a gas sensor can accurately detect
the presence of a specific gas in the presence of other interferents.
Although absolute selectivity may be difficult to achieve, relative se-
lectivity is highly possible if selectivity is defined contextually, that is,
previous knowledge of known interferents in a targeted application or
environment (25).
Using multiplexed CS-FETs, we show that it is possible to actively track and detect the gases of the target application described earlier, sensitively and selectively at low power. By iterative elimination, three sensing layers were identified for the abovementioned three gases of interest and deposited on individual CS-FETs: Pd$_{0.3}$nmAu$_{1}$nm for H$_2$S, Ni$_{0.3}$nmPd$_{1}$nm for H$_2$, and Ni$_{1}$nm for NO$_2$. As described in the previous section, all three CS-FET gas sensors in this work operate at room temperature. Figure 1D shows a cross-sectional transmission electron microscopy (TEM) image and an associated energy-dispersive x-ray spectroscopy (EDS) profile of a completed device with an ultrathin composite sensing layer of Pd-Au capacitively coupled to a 3.5-nm-thin silicon channel (see also fig. S2). This CS-FET (discussed below) is sensitive to the detection of H$_2$S.

In terms of device operation, the CS-FET works as a junctionless n-channel transistor, with the channel degenerately n-doped to similar levels as the source and the drain (26). Most charge carriers in the 3.5-nm-thin silicon channel can be fully depleted upon gas exposure, leading to large modulations in the output current. Sensing principle in the CS-FET is based on the channel threshold voltage shift ($V_t$ shift), depicted in Fig. 2 (A and B) by a simplified band alignment representation across the sensing layer and silicon channel before and after exposure to a target gas (in this case, H$_2$S). Gas interaction with the Pd-Au sensing layer can lead to a change in its work function ($\Phi_m$) as well as charge density. This net change increases the channel potential, leading to a decrease in device threshold voltage (more negative $V_t$) and an increase in $I_D$. Sensing is highly dependent on the nature of the interacting gas and the sensing layer, decreased $V_t$ for reducing gases (for example, H$_2$S) and increased $V_t$ for oxidizing gases (for example, NO$_2$). Diagnostic back gate measurements ($I_D$ versus $V_{BG}$ and $I_D$ versus $V_{DS}$) in Fig. 2 (C and D) experimentally validate this $V_t$ shift sensing principle upon targeted gas exposure.

Figure 3A shows time-dependent measurements of the Pd-Au-based H$_2$S sensor operated at a supply voltage of 100 mV. With a sampling time of 1 s, the baseline current is ~30 nA, leading to a low standby power consumption of ~3 nW. This particular sensor has a very low detection limit of 1 part per billion (ppb) with a saturation concentration of above 10 ppm. The response versus concentration curve (Fig. 3B) shows a linear region from 1 to 10 ppm with a sensitivity of 280%/ppm [($\Delta G/G_o$) per ppm]. This high sensitivity is attributed to the thin sensing layer as well as the thin silicon channel (as illustrated in fig. S3, thick silicon channel junctionless transistors results in a much lower sensitivity). Response times ($t_{90}$; time taken by the sensor current to reach 90% of its maximum value upon exposure to gas) are longer for smaller gas concentrations, with ~144 s for 1 ppb of H$_2$S and ~112 s for 100 ppm of H$_2$S. Microheater-assisted recovery times ($t_{50}$; time taken by the sensor current to drop to within 50% of its...
original baseline value after removal of gas) of H2S sensors increases with concentration, with ~20 s for 1 ppb of H2S and ~800 s for 100 ppm of H2S. A possible hypothesis for this is that H2S tends to strongly bond with the Pd-Au sensing layer, making it harder to desorb excess H2S molecules at larger concentrations (see fig. S4).

Similar to the H2S CS-FET sensor, the Ni-Pd-based H2 sensor also exhibits linear and saturation characteristics in its time-dependent response curve (Fig. 3C). A supply voltage of 2.5 mV was used to drive the sensor at a baseline current of ~135 nA, consuming just ~0.34 nW of power. The lowest limit of detection is approximately 0.3% H2 (~3000 ppm), with a saturation occurring above 0.5% H2 (~5000 ppm) and a maximum sensitivity of 0.24%/ppm observed within this linear region. The interaction mechanism between H2 and Ni-Pd is based on the dissociation and diffusion of the atomic hydrogen in palladium (27). Ni content in the sensing layer blocks H2S interaction with palladium (28), as will be demonstrated later. Compared to the H2S sensor, response times are much shorter with a minimum t90 of ~17 s for 2% H2 (Fig. 3D). Microheater-assisted recovery time is also much shorter and is quite similar throughout the entire H2 sensing range with a t90 of ~30 s.

In the case of the Ni sensing layer–based NO2 sensor (Fig. 3E), baseline power consumption is ~2 nW based on a sensor current of ~820 nA at a supply voltage of 2.5 mV. Here, the CS-FET output current swings in the opposite direction compared to the H2 and H2S sensor. As described previously, this indicates that the transistor Vt has increased (becomes more positive). The understanding here is that electron donation from NO2 to the Ni sensing layer occurs because of its oxidizing nature. This makes the Ni sensing layer electronegative, leading to a decrease in its effective work function (~Δϕ) and hence decreasing the potential of the capacitively coupled channel. Consequently, the transistor Vt becomes more positive and leads to a decrease in the CS-FET current (I3). As can be observed, the NO2 CS-FET has a detection limit of 1 ppm and a saturation concentration of 50 ppm. The sensitivity of NO2 is approximately 1%/ppm (Fig. 3F), and the minimum t90 is ~83 s for 100 ppm of NO2. Sensor recovery is near instantaneous with t90 dominated by the microheater on time (20 s).

As was previously discussed, the CS-FETs are fabricated using a top-down CMOS-compatible fabrication process that can facilitate the integration of large arrays of 3.5-nm-thin silicon channel transistors. Through multiple processing steps, different types of chemical-sensitive layers can be deposited on each transistor in an array, enabling a powerful multiplexed gas sensing platform. We demonstrate this using multiplexed H2S (Pd0.3nmAu1nm), H2 (Ni0.3nmPd1nm), and NO2...
(Ni1nm) CS-FETs in Fig. 4, which is used to detect a sequence of eight gas pulses. Note that heat from the local microheaters manifests as current spikes in the CS-FET in response to temperature change during the recovery process, which is similar to that in Fig. 3. In this experiment, three successive 10-ppm H2S pulses were injected for 5 min, which were sensitively detected by the Pd-Au-based H2S CS-FET. The H2 and NO2 sensors had minimal effects due to the Ni content in sensing layers, which blocks any H2S interaction. Next, three consecutive injections of 0.5% H2 pulses caused the Ni-Pd-based CS-FET to respond markedly. The NO2 and H2S sensors showed only slight response to the 0.5% H2 pulses, confirming their sensing layers as being relatively selective over hydrogen as well. Although the Pd-Au alloy-based sensing layer for the H2S CS-FET may respond to much larger concentrations of H2, theoretical and experimental works have hypothesized that Pd-Au clusters tend to interact weakly with H2 compared to pure Pd (29, 30). To further check the selectivity between the three CS-FETs, we sequentially carried out 10-min mixed-gas test of H2S and H2. For the first 5 min, we injected 10 ppm of H2S from the H2S delivery line, and only dry air from the hydrogen delivery line was flowing. A 0.5% H2 pulse was then injected for the next 5 min in the H2S ambient. As observed, the H2S and H2 CS-FET sensors were able to respond to their respective target gases. During this mixed-gas test, the NO2 CS-FET sensor responded minimally. Toward the end of the multiplexed measurements, a 100-ppm NO2 pulse was injected for 5 min, which was quickly picked up by the Ni-based NO2 CS-FET. The H2 and H2S sensors responded minimally to NO2, with a decrease in sensor currents. After each detection event, the local microheaters of all three CS-FETs were turned on simultaneously to recover and reset to sensor baseline.

On the basis of the results in Figs. 3 and 4, the silicon CS-FET framework provides a unique platform that can be used to integrate multiple gas sensors onto a single chip, beyond the three gases of interest in this work. Furthermore, the low power consumption of the underlying 3.5-nm-thin silicon transistor makes it possible for integration with mobile wireless electronics. To demonstrate this, we present a proof-of-concept application of our CS-FET gas sensor technology by integrating a H2-sensitive CS-FET sensor onto a wireless drone (Fig. 5). There is significant interest in unmanned aerial vehicle-based aerial mapping of harmful chemicals in the environment for threat detection/neuralization and ozone mapping, with a few early attempts using metal-oxide semiconductor gas sensors (31). Here, we used an off-the-shelf microdrone equipped with an ARM Cortex microprocessor, an inertial measurement unit, a barometric pressure sensor, and a radio frequency (RF) communication link (Crazyflie 2.0) (Fig. 5A). The Ni-Pd H2-sensitive CS-FET was integrated onto this drone using a simple voltage divider circuit. In a contained environment, a controlled leak of pure H2 was carried out (estimated resulting concentration in air, 1%). The test environment is shown in Fig. 5B, where the drone was tethered to high tension strings on either of its sides, such that its flight path was restricted in the vertical direction (z axis) only. A pre-programmed flight path was defined from a laptop that is enabled with an RF transceiver, where the drone would take off and climb at a steady...
rate to the source of the hydrogen leak, hover for a set period, and then descend, all while transmitting back the raw sensor data, as indicated in Fig. 5C. Figure 5D shows the real-time H$_2$ sensor measurement result and altimetry data corresponding to the experiment in Fig. 5C. Note that, because this was a proof-of-concept demonstration, no attempt was made to calibrate the raw sensor data. Furthermore, microheaters were not used for sensor recovery, indicating the slow H$_2$ sensor recovery to baseline at the end of the experiment.

DISCUSSION

We have shown the CS-FET technology as a sensitive, selective, and low-power gas sensing platform based on ultrathin silicon channel transistors using conventional CMOS processing. Because of its relatively inert nature, pristine silicon can be engineered to be selective to a wide variety of gases. Using this, we have shown the possibility of contextual selectivity and multiplexed gas sensing by using CS-FETs functionalized with sub-5-nm metallic sensing layers, Pd-Au, Ni-Pd, and Ni, that are selective to three target gases, H$_2$S, H$_2$, and NO$_2$, respectively, over each other. Finally, as a proof of concept for integration with mobile electronics, we demonstrated a H$_2$-sensitive CS-FET integrated with a wireless microdrone for aerial detection of hydrogen.

From a practical large-scale fabrication standpoint, thickness uniformity control of the SOI wafers is an important challenge to the ultrathin silicon CS-FET technology presented here. However, the silicon integrated circuit (IC) industry is already addressing this, where these ultrathin and fully depleted SOI wafers are being explored to develop next-generation low-power digital electronics. From an economic perspective, the cost of these devices will be minimal compared to the silicon IC industry because advanced nanofabrication processing techniques are not required for the CS-FET gas sensor technology. The final cost will be determined by chip size, number of chips per wafer, and packaging. In summary, our silicon CS-FET gas sensor platform can provide a robust platform for the research and development of new sensing layers for a wide variety of target chemicals, which will enable multigas detection on a single chip and at low power, paving the way to applications in environmental pollution mapping and personal air-quality monitors.

MATERIALS AND METHODS

Sensor fabrication process flow

The starting substrates, <100> SOI wafers 200 mm in diameter, were used for CS-FET fabrication, which were procured from Soitec, with 70-nm-thick top silicon, 145-nm-thick buried oxide, and 725-μm-thick silicon handle wafer. The active top silicon layer has a thickness variation of ±7.5 nm. Both the top silicon and handle wafer silicon were lightly boron (p~)-doped, with a starting resistivity range of 8.5 to 11.5 ohm·cm. All experiments, except fig. S5, were carried out with devices made from these Soitec wafers. New and different devices were used each time for all experiments. All fabrication steps of the CS-FET sensor devices were carried out in an academic laboratory, because of which there are variations in channel thickness and doping concentration, resulting in sensor response and sensitivity variation. The CS-FET sensors for the experiment in fig. S5 were made using 300-mm <100> SOI wafers procured from Shin-Etsu with a 12-nm-thick top silicon layer (resistivity,
9 to 18 ohm-cm; boron), a 145-nm-thick buried oxide, and a 775-mm-thick silicon handle wafer (resistivity, 9 to 18 ohm-cm; boron).

The fabrication process is depicted in fig. S6. To achieve ultrathin silicon on insulator transistors, the initial 70-nm top silicon of the SOI wafer was trimmed by oxidation (Tystar furnaces) at 1000°C in atmosphere using a three-step dry (5 min)–wet (13:45 min)–dry (5 min) process. This ensured rapid oxidation with a resultant low-stress SiO2 and a clean SiO2/Si interface. After this, a dilute (1:10) hydrofluoric acid (HF) solution was used to strip the grown oxide, leaving a thin top silicon layer. Six-point, fixed-angle (70°) ellipsometry indicated that the top silicon average thickness was approximately 5 to 6 nm, which was later confirmed to be approximately 3.5 nm under TEM. After this, 250-μm-long and 5-μm-wide mesas were patterned using i-line photolithography with a clear field quartz mask and image reversal photoresist (MicroChemicals GmbH AZ nLOF 2020). Next, using a TCP Poly-Si etcher (Lam Research Corp.), silicon etching in Cl2 and HBr completed thin-body silicon mesa etching. The etching process resulted in a slight over-etch in the buried oxide (<10 nm). Postetching, the photoresist atop the silicon mesa patterns, was removed in acetone. An ensuing wet cleanse was performed in a standard peroxide (Piranha) mixture at 120°C followed by a 10-s dilute HF (1:10) dip. Source/drain (S/D) contacts were then defined with physical gate lengths (S/D separation) of 2 and 5 μm. Ni (50 nm) was deposited using thermal evaporation, before which the wafers were stripped of native SiO2 in a 1:50 dilute HF solution for 20 s. Ni lift-off was carried using thermal evaporation, before which the wafers were stripped of native SiO2 in a 1:50 dilute HF solution for 20 s. Ni lift-off was carried out in acetone after which the wafers were annealed in forming gas at 350°C for 1 min to form a nickel silicide (NiSi). After this, microheater patterns were defined adjacent to the CS-FETs, after which Au (400 nm)/Cr (20 nm) was deposited by electron beam evaporation followed up lift-off in acetone. Functionalization of individual CS-FETs was carried out by a sequential blanket evaporation of different materials using a combination of thermal and electron beam, at a very slow rate (~0.1 to 0.2 Å/s) at room temperature. No annealing or chemical treatment was carried out on the sensing layers and was used as deposited.

Sensor measurement apparatus and setup
To ensure health and safety, all gas sensing measurements were carried out in a walk-in fume hood. The CS-FET chip was wire-bonded to a 64-pin chip carrier. Different power supply sources were used for the CS-FET sensor devices and the individual microheaters, with all CS-FETs tied to a common ground and the individual microheaters tied to a different common ground. The chip carrier containing the CS-FET sample was housed in a three-dimensional printed minichamber (47 mm × 47 mm × 23 mm) made of chemically resistant polylactic acid (PLA) material. The minichamber was later confirmed to be approximately 3.5 mm under TEM. After this, 250-μm-long and 5-μm-wide mesas were patterned using i-line photolithography with a clear field quartz mask and image reversal photoresist (MicroChemicals GmbH AZ nLOF 2020). Next, using a TCP Poly-Si etcher (Lam Research Corp.), silicon etching in Cl2 and HBr completed thin-body silicon mesa etching. The etching process resulted in a slight over-etch in the buried oxide (<10 nm). Postetching, the photoresist atop the silicon mesa patterns, was removed in acetone. An ensuing wet cleanse was performed in a standard peroxide (Piranha) mixture at 120°C followed by a 10-s dilute HF (1:10) dip. Source/drain (S/D) contacts were then defined with physical gate lengths (S/D separation) of 2 and 5 μm. Ni (50 nm) was deposited using thermal evaporation, before which the wafers were stripped of native SiO2 in a 1:50 dilute HF solution for 20 s. Ni lift-off was carried out in acetone after which the wafers were annealed in forming gas at 350°C for 1 min to form a nickel silicide (NiSi). After this, microheater patterns were defined adjacent to the CS-FETs, after which Au (400 nm)/Cr (20 nm) was deposited by electron beam evaporation followed up lift-off in acetone. Functionalization of individual CS-FETs was carried out by a sequential blanket evaporation of different materials using a combination of thermal and electron beam, at a very slow rate (~0.1 to 0.2 Å/s) at room temperature. No annealing or chemical treatment was carried out on the sensing layers and was used as deposited.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/3/3/e1602557/DC1
51. Sensor characteristics of an unfunctionalized CS-FET sensor
52. Nanoparticle sensing layer sparsity analysis
53. Effect of silicon channel thickness on CS-FET sensitivity
54. CS-FET sensor response to temperature, relative humidity, and room temperature recovery characteristics
55. Individual sensor cycling from low and high gas concentrations
56. Fabrication process of chemical-sensitive 3.5-nm-thin silicon transistors
57. Microheater characterization
58. Transfer characteristics of H2S, H2, and NO2 CS-FETs after each gas exposure for a 20-s period with a draw current of 100 mA from each individual heater.

REFERENCES AND NOTES


Room temperature multiplexed gas sensing using chemical-sensitive 3.5-nm-thin silicon transistors

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