Molecular bridge-mediated ultralow-power gas sensing

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Abstract
We report the electrical detection of captured gases through measurement of the quantum tunneling characteristics of gas-mediated molecular junctions formed across nanogaps. The gas-sensing nanogap device consists of a pair of vertically stacked gold electrodes separated by an insulating 6 nm spacer (~1.5 nm of sputtered α-Si and ~4.5 nm ALD SiO2), which is notched ~10 nm into the stack between the gold electrodes. The exposed gold surface is functionalized with a self-assembled monolayer (SAM) of conjugated thiol linker molecules. When the device is exposed to a target gas (1,5-diaminopentane), the SAM layer electrostatically captures the target gas molecules, forming a molecular bridge across the nanogap. The gas capture lowers the barrier potential for electron tunneling across the notched edge region, from ~5 eV to ~0.9 eV and establishes additional conducting paths for charge transport between the gold electrodes, leading to a substantial decrease in junction resistance. We demonstrated an output resistance change of >108 times upon exposure to 80 ppm diamine target gas as well as ultralow standby power consumption of <15 pW, confirming electron tunneling through molecular bridges for ultralow-power gas sensing.

Introduction
The development of low-power Internet-of-Things (IoT) sensor systems has been rigorously pursued by an increasing number of scientific communities to enable continuous access to various information around the globe. Detection of motion, humidity, and temperature was proposed for the purpose of creating a user-accessible database via the implementation of a low-power IoT wireless sensor network by Laubhan1. Improvements to gas detection techniques have been rigorously pursued to cover a wide range of commercial IoT applications, including indoor air quality monitoring, and industrial applications, such as the detection of hazardous or combustible gases. Additionally, similar gas sensors have been deployed for remote and continuous environmental monitoring to detect the presence of toxic gases such as volatile organic compounds (VOCs), CO, SO2, H2S, and O32.

Among such power-limited IoT applications, gas sensors remain one of the most challenging components due to the power-hungry nature of gas sensing, especially in the required continuous operation mode. Although gas sensors typically vary in their operating temperature requirements, most of them consume >10 μW of power due to the requirements of elevated temperature or heating3. Recently, some low-power (sub-10 μW) gas sensors have been demonstrated; however, they displayed only limited output signal changes of less than a few orders of magnitude, ultimately being limited in the minimum detection amount when utilizing ~μWatt level of power. These sensors achieved low-power operation by utilizing self-heating under a bias voltage or by minimizing electrical leakage paths within the device, such as self-heating carbon nanotubes4, Si-nanowire/TiO2 core–shell heterojunctions5, chemically gated Si nanowire/SnO2 thin film FETs6 and Pd-based Si nanomembrane sensors7. They were limited in the output signal
range mainly because the transducer output varied line-
arily with the analyte concentration in a physically con-
ected device structure. For example, a CNT NO₂ sensor
displayed only a < 10 times change in the output current
upon exposure to 0.9 ppm target gas while consuming
0.8 μW of DC power, and an FET gas sensor produced an
output of <10² times upon exposure to 104 ppm analyte
while consuming ~1 μW of DC power.

Such limited output signals can be further amplified
without additional power consumption by physically
forming an electrically conductive bridge across an other-
wise electrically nonconducting open space mediated by
the capture of target gas molecules. This capture would lead to
electron tunneling through the molecular bridge, resulting
in a current change of multiple orders of magnitude due to
the exponential dependence of the tunneling current on
parameters such as the tunneling distance and energy
barrier. An open-air gap could be realized by constructing a
parallel plate structure between metal electrodes and
matching the gap distance with the molecular size of the
target gas. The initial air gap provides an extremely high DC
off resistance between electrodes. This leads to a minimal
leakage current and hence a highly reduced standby power
consumption. A similar nanogap concept was previously
introduced for quantum tunneling-based gas sensing with preliminary results15–25.

This paper reports detailed proof-of-concept results of a
new family of ultralow-power nano-air-gap sensors based
on molecular bridge formation across nanogap electrodes.
These air-gap sensors evidently demonstrated very high
ON/OFF output signal ratios due to molecular bridge-
mediated electron quantum tunneling. Specifically, this
article reports the operation principle, device structure
and fabrication, and sensor test results.

Materials and methods

Device design and fabrication

Details of the device fabrication method have been
extensively discussed in our previous article26. Figure a
shows the simplified fabrication process flow. The nano-
gap dimension is defined by the thickness of the deposited
spacer layer and is suitably engineered to match the
length of the linker molecule and the target molecule for
successful and specific capture of the target analyte. Fig-
ure 1b shows SEM images of the fabricated nanogap
device and overlap area and a high-resolution SEM image
of the nanometer-scale air gap.

Structure of target and linker molecule

To demonstrate the validity of the proposed sensing
mechanism, we aimed to detect a vapor of 1,5-
pentanediamine, commonly known as cadaverine, as the
target gas molecule. Cadaverine was purchased from
Millipore Sigma in liquid form (97% purity). Figure S1 in
Supplementary section describes the synthesis of the lin-
kern molecules. Figure 2 shows the chemical structures of
the target gas, cadaverine, and the synthesized linker
molecule. The thiol linker molecule is comprised of three
components: the sulfur head group, the aromatic and
alkyl spacer chain, and the benzoic acid tail group. The
head group, consisting of thiol molecules (-SH), is capable
of covalently bonding to the gold surface. The spacer
chain defines the length of the linker molecule and, due to
its conjugated nature, allows augmented electron flow
throughout the molecule. Capture of our targets is
achieved by the tail segment of the linker, which in this
case is composed of benzoic acid. The terminal carboxylic
acid from the benzoic acid and the matching length of the
spacer chain with the nanogap distance together deter-
mine the specificity of the target gas molecules to be
captured. The IUPAC name of the synthesized linker
molecule is (4-((4-((4-mercaptophenyl)ethynyl)phenyl)
ethynyl)benzoic acid). Additional details of the SAM lin-
kern molecule structure and synthesis are provided in the
supplementary section.

SAM coating procedure, coating characterization, and
target capture specifics, and linker conductivity
measurements

Thiol SAM chemisorption on gold surfaces, first reported
by Bell Laboratories in 198327, was performed by dissolving
30 mg of the linker chemical in 5 mL of dimethyl sulfoxide
(99.9% purity as purchased from Millipore Sigma) and
~15 mL of ethanol and immersing the gold-coated device
for a specified amount of time. It should be noted that
ethanol does not chemically influence the coating proce-
dure, and the required volume is simply to ensure complete
immersion of the device within the solution. Figure 3a–c
shows images taken under a fluorescence microscope of a
sample exposed to similar concentrations of the fluorescent
cadaverine target (Alexa Flou 488, Sigma Aldrich) for
2 min after being immersed in the linker chemical solution
for different periods of time (12, 24 and 48 h). Figure 3c
shows the highest density of captured fluorescent cadaver-
ine molecules and, by extension, the formation of an SAM
upon immersing the sample for 48 h. Leaving the sample
submerged within the solution for more than 48 h resulted
in the gold layer peeling off and was therefore avoided.
Hence, a 48-h coating procedure was chosen to be part of
the protocol. Conductivity measurements of the linker
solution performed using Peak force tunneling atomic force
microscopy (PF-TUNA) revealed higher conductivity than
commercially available alkane-thiol linker molecules, as
shown in Fig. 3d, e.
Estimation of the target gas concentration

The concentration of the cadaverine target gas was estimated by first establishing a calibration curve between known cadaverine liquid volumes and corresponding signal intensities (areas) and then comparing the obtained calibration curve to the signal intensities obtained by injecting an unknown concentration of the gas sample. Here, a gas chromatography-flame ionization detector (GC-FID) system was utilized to inject different masses of liquid cadaverine and monitor the chromatogram signal intensities at each mass. The utilized masses of cadaverine were 6.2, 12.4, 18.6, 24.8, and 31 ng, with corresponding diluted cadaverine samples of 0.1, 0.2, 0.3, 0.4, and 0.5 μL.

Note that the cadaverine sample was diluted to 1% with a mixture of water to reach a much lower level (nanogram) of liquid injection and, more importantly, to maintain the signal intensity in a similar range as later gas injection. Finally, the injected mass \((m_{\text{inj}})\) vs. signal intensity \((A_{\text{FID}})\) data were fitted with a linear equation: \(A_{\text{FID}} = 31.18 \times m_{\text{inj}} - 0.03794\).

To determine an unknown concentration of cadaverine, first, a gas sample was prepared inside a closed chamber, and then, by injecting the gas sample into the same GC-FID system, the mass of the gas sample was calculated according to the above equation. The calculated mass was then converted to ppm considering complete evaporation of cadaverine at STP. For example, if the estimated mass was 1.3 ng, this corresponded to 12.31 picomoles of liquid mass or 301 pL of gas at STP. For this example, the gas injection volume was 100 μL, and therefore, the ppm was calculated as 3.01 ppm.

Sensor measurements in the test chamber

The test setup utilized a stainless steel cylindrical structure (height: 15 cm and diameter: 5.5 cm) with an injection hole sealed with a rubber septum seal and two
shows a schematic of the test setup. The electrical feedthroughs connected to the fabricated sensor located inside the chamber were also connected to an impedance analyzer, a Keithley 4200A-SCS, outside the chamber. The Keithley impedance analyzer had a resolution of 100 fA. The chip was supplied with a DC bias of 0.7 V across the nanogap-separated electrodes. The ambient test conditions were continuously monitored using a BME280 chip, which is a commercially available humidity, temperature, and pressure sensor. The chamber was not subject to any vacuum modifications, and it was kept under typical atmospheric conditions at atmospheric pressure and maintained at a room temperature of ~23 °C during all tests. A μ-syringe was used to introduce cadaverine gas into the test chamber through the septum seal. The concentration of cadaverine within the chamber was calculated from the volume of cadaverine injected as described in Section 2.4, and the sensor response was measured using the Keithley impedance analyzer. Figure 4 shows a schematic of the test setup.

**Tunneling operation based on molecular bridging**

Figure 5a illustrates the sensor operating principle based on molecular bridging. The two electrodes are originally disconnected, resulting in a high tunneling resistance on the order of 10⁹ Ω as the 'OFF' state. These electrodes are coated with a self-assembled monolayer (SAM) of conjugated thiols that capture the target gas molecules. After molecular capture, additional current conduction paths are created between the electrodes via the formation of capture-mediated molecular bridges. This leads to an enhanced quantum tunneling current and a significant reduction in the junction resistance as the 'ON' state.

The electrical I–V characteristics and device resistance are determined by the species trapped in the nanogap, which can be two organic SAM layers separated by an air gap (OFF state) or an SAM-captured gas molecule-SAM chain (ON state). The electrical conductivity of these is examined below.

**Thiol SAM capture layer**

Generally, the electrical properties of organic molecules can be determined by using AFM/STM, or (mechanically controllable breaking) MCB techniques or can be simulated using computational chemistry methods. Electrical transport through conventional alkane(di)thiol SAMs has been described as pure quantum tunneling across a thin dielectric film that has a rectangular potential barrier with image charge effects included, as described by J. Simmons in 1963. The dielectric constant of a typical alkane-thiol SAM layer was determined by impedance measurements to be 2.1 according to Akkerman. Akkerman also verified that the barrier height of alkane-thiol SAMs sandwiched between a pair of Au electrodes and protected by a layer of PEDOT:PSS (within the spacer stack) was in the range of 4–5 eV. Although alkane thiols are poor conductors of electricity, the conductivity of thiol molecules can also be engineered by the addition of alternative alkyne and aromatic rings within the molecule itself. This modification leads to a conjugated molecular arrangement that results in amplified charge transport along the entire molecule because of delocalized n-electron orbitals, which can freely move along the whole molecule. Bower also verified that conjugation in SAMs containing oligophenyl groups led to a reduction in the attenuation constant, which essentially meant an augmented conductance for molecules having a higher degree of conjugation. This exponential dependence of the resistance of SAMs on the SAM length (RSAM = k · eγ length) has been extensively studied and verified. Using a combination of the non-equilibrium Green’s function, Density functional theory (DFT), and confirmed I–V characteristics, Ratner concluded that the degree of conjugation and overlap of the orbital densities of the molecular end group and the contact electrodes together determine the heightened conductivity of these molecular chains because of constant delocalized charge transport. Alkane diamine molecules have also been studied using common computational chemistry algorithms, where the length dependence of the electrical conductance has been observed to be exponential, similar to thiol linker molecules. Generally, most organic molecules demonstrate an exponential dependence of the electrical properties on the molecule length. However, it must be noted that there is a transition molecular length beyond which the resistance dependence is almost ohmic in nature. Although this critical length depends on the specific organic molecule forming the molecular chain, generally, for conjugated molecules, it is ~4.0 nm. This exponential dependence of the electrical resistance on the length of the molecular chain is a lower limit of the device resistance; hence, for higher resistance changes, one should engineer

![Chemical structures and molecule lengths of the target molecule (on the left) and linker molecule (on the right). The IUPAC name of the linker molecule is (4-(4-((4-mercaptophenyl)ethynyl)phenyl)ethynyl)benzoic acid)](image-url)
the nanogap and SAM dimensions such that the SAM is as short as possible and the air gap is the same length as the captured gas molecule. This combination maximizes the resistance ratio \( R_{OFF}/R_{ON} \) and the dynamic range of the device. The electrical equivalent model for the device is discussed below.

**Electrical equivalent model**

The mechanism for electrical current conduction across the junction of our device is considered to be electron tunneling. In this device, as shown in the schematic of Fig. 5b, there are two possible electron conduction paths represented by current sources \( I_S \) and \( I_{E(C g)} \). Current \( I_S \) represents electrical conduction through the dielectric spacer stack under the overlap region of the two electrodes. Current \( I_{E(C g)} \) represents electrical conduction through the multiple molecular bridges formed along the edges of the top electrode. This current is a function of the gas concentration \( C_g \). In the absence of an analyte target gas (\( C_g = 0 \)), current mainly flows through the spacer stack in the overlap region, with an overlap region of \( \sim 9 \) \( \mu \)m \(^2\); thus, \( I_T \approx I_S \). On the other hand, when the target gas is captured between the nanogaps along the edges, current \( I_{E(C g > 0)} \) will additionally flow through the newly formed molecular junction in the edge undercut area of \( \sim 0.06 \) \( \mu \)m \(^2\). The device gap and linkers should be designed such that \( I_{E(C g > 0)} \gg I_S \) and \( I_T \approx I_{E(C g)} \) after the target gas is captured.

The total current flowing across the junction can be written as

\[
I_T = I_S(V) + I_E(V, C_g)
\]

(1)

where \( I_T \) is the total current flowing across the nanogap sensor and \( V \) is the bias voltage across the device. The substrate and edge current components have different functional forms. The substrate component, which corresponds to conduction through thin dielectric stacks, is readily modeled using the generalized Simmons’
formula for tunneling current. The current \(I_0\) across the overlap support region can be expressed as

\[
I_0(V) = G_{SO} \left[ \frac{\Phi_d}{q} \cdot e^{-\frac{2\pi}{\sqrt{qV}}} + \left( \frac{\Phi_d}{q} + V \right) \cdot e^{-\frac{2\pi}{\sqrt{q(V+\Phi)}}} \right]
\]

where \(\Phi_d\) is the barrier energy of the spacer dielectric, \(d_s\) is the spacer layer thickness, \(m\) is the effective mass of tunneling electrons, \(\hbar\) is the reduced Planck’s constant, and \(q\) is the charge of an electron. The parameter \(G_{SO} = A_g G_{SO}\) is a conductance fitting parameter proportional to the area, \(A_g = W^2\), of the spacer dielectric, and \(g_{SO}\) is a junction conductance factor.

The derivation of a mathematical expression for \(I_E\) is considerably more difficult, and in general, it involves the calculation of nonequilibrium Green’s functions that specify the ballistic electron transport across the various molecular levels of the trapped gas molecule. Such calculations can lead to tunneling resonances and negative resistance regimes. The model that we have adopted below is based on our experimental observations, which did not display any resonant tunneling characteristics.

In the absence of a target gas, the \(I-V\) characteristics resemble those provided by a Simmons-like model. However, when investigating current conduction across molecular chains, Ghosh found that Simmons’ expression for the tunneling current was unable to describe the \(I-V\) characteristics accurately. The reason was that certain assumptions made during the derivation of Simmons’ expression did not hold for a molecular chain. N. Zimovskaya found that the conduction through molecular junctions can be modeled by combining a Wentzel–Kramers–Brillouin (WKB) approximation of the transmission coefficient and the Landauer formula, which leads to a mathematical expression of the tunneling current similar to Simmons’ formula. The tunneling current depends exponentially on a function of the average energy barrier, as seen by the tunneling electrons. When the analyte gas is captured by our sensor, the dominant current is the gas-dependent edge current through the captured molecules, and using the Zimovskaya formula, it is given by

\[
I_E(V, C_g) = \left[ \frac{\beta \cdot C_g}{1 + \beta \cdot C_g} \right] \cdot \Gamma_{IE} \cdot \left[ \sqrt{\left( \frac{\Phi_d}{q} - \frac{V}{2} \right) - e^{-2d_s \sqrt{\frac{q}{2}} (\Phi_d + \frac{V}{2})}} \right]
\]

where \(\beta\) is a fitting parameter, \(C_g\) is the concentration of the target analyte gas in the test chamber, \(\Phi_E\) is the average barrier potential of the hybrid molecular chain, and the distance \(d_s = 2 \times (\text{the length of the linker molecule}) + \text{the length of the analyte molecule}\). Two parameters are expected to change in this equation when a gas molecule is captured. The first and most important gas concentration-dependent parameter is the average energy barrier \(\Phi_E\), as shown in the zero-bias energy diagram of Fig. 5c. In Fig. 5c, \(\Psi_{SAM}\) and \(\Psi_{alu}\) are the work functions of the SAM layer and the gold electrode, respectively. \(x_{\text{target}}\) is the electron affinity of the target gas.

Note that the average energy barrier depends on what is inside the edge gap. The energy barrier of the...
molecular bridge is obtained by fitting and comparing curves at different gas concentrations with the Zimbovskaya formula to obtain an estimate of the average energy barrier that electrons encounter. If no molecule is captured, then the average barrier is at the maximum given by the work function of the linker. If a molecule is captured, then the average barrier established on that junction depends on the location of the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) energy levels of the captured molecule. Curve fitting and parameter extraction reveal that in general, capture results in a lower average energy barrier $\Phi_E$ and hence a large enhancement of the device current.

The second concentration-dependent factor is the leading term

$$[\beta * C_G] \Gamma_{EO} \frac{1}{1 + \beta * C_G}$$

which is representative of the number of molecular junctions formed and is indicative of the saturation-type Langmuir adsorption characteristic. The parameter $\beta$ is a fitting parameter representing the adsorption characteristic, has a unit of 1/gas concentration and determines the low-concentration limit behavior of the edge current. This equation implies that the higher the surface concentration ($C_G$ of the target gas), the larger the number of molecular junctions formed ($\beta * C_G$) $/(1 + \beta * C_G)$, ultimately leading to a higher edge current $I_E(V,C_G)$. The edge current saturates when all possible absorption sites are full. The parameter $\Gamma_{ED} = p g_{EO}$ is a fitting parameter proportional to the edge perimeter $p = 4(W + 2A)$. This parameter specifies the magnitude of the nonlinear edge conductance in units of $AV^{-1/2}$.

Note that Eq. 3 also displays an exponential dependence of the molecular junction resistance on the total molecular length, in agreement with observations made by others.

**Results and discussion**

**Experimental $I$-$V$ characteristics before and after gas exposure**

The fabricated sensor, without being exposed to a target gas, showed a minimal leakage current across the tunneling junction (on the order of $\sim$50 pA with a
molecules is facilitated by the electrostatic attraction between the end benzoic acid group of the linker molecule and the highly polar NH₂ end group of the cadaverine molecule. Additionally, the nano-air-gap dimension is designed to be ~6 nm. The linker molecule is ~2.5 nm in length. Therefore, after proper functionalization, the nano-air-gap dimension reduces to ~1.0 nm, making it ideal for capture of cadaverine since its length is ~1.0 nm. We believe that the electrostatic attraction combined with careful fabrication of the nanogap is the reason cadaverine is captured and thereby detected strongly. Successful capture is accompanied by the formation of hydrogen bonds between the linker and cadaverine molecules. According to the phenomenological model provided by Zimbovskaya⁴⁰, capture of cadaverine molecules leads to a decrease in the average potential barrier of the tunneling junction, which allows for an augmented electron flow across it. The tunneling current is an exponential function of the potential barrier encountered by electrons⁴⁵. Therefore, even a slight decrease in the potential barrier leads to a significant increase in the tunneling current, which is observed as a strong signal upon capturing cadaverine.

Theoretical parameter extraction demonstrated that the exposure of the fabricated device to ~80 ppm cadaverine gas reduced the average potential barrier from ~5 eV to ~0.9 eV, resulting in exponential tunneling current variation with the square root of the barrier potential change, and that the fitting parameters β and ΓED in Eq. 3 were 21.685/ppm and 0.3688 A/V⁹/₂, respectively. It must be noted that the model-extracted values of the reduced average potential barrier were not a quantitative measurement of the HOMO-LUMO levels of individual target/linker molecules but a cumulative indication of the average potential barrier faced by tunneling electrons. In fact, Zimbovskaya⁴⁰ mentioned that the inherent

**Experimental and fitted model I–V characteristics at different gas concentrations**

Figure 7a shows plots of experimentally obtained I–V data curve-fitted to the tunneling current model, as described in Eq. 3, at various analyte concentrations and their tight agreement. The electrical measurements were taken after the sensor signal had reached a steady state. As evident from the plot, exposure to ~80 ppm cadaverine led to a reduction in the junction resistance by ~9 orders of magnitude. The capture of cadaverine molecules is facilitated by the electrostatic attraction between the end benzoic acid group of the linker molecule and the highly polar NH₂ end group of the cadaverine molecule. Additionally, the nano-air-gap dimension is designed to be ~6 nm. The linker molecule is ~2.5 nm in length. Therefore, after proper functionalization, the nano-air-gap dimension reduces to ~1.0 nm, making it ideal for capture of cadaverine since its length is ~1.0 nm. We believe that the electrostatic attraction combined with careful fabrication of the nanogap is the reason cadaverine is captured and thereby detected strongly. Successful capture is accompanied by the formation of hydrogen bonds between the linker and cadaverine molecules. According to the phenomenological model provided by Zimbovskaya⁴⁰, capture of cadaverine molecules leads to a decrease in the average potential barrier of the tunneling junction, which allows for an augmented electron flow across it. The tunneling current is an exponential function of the potential barrier encountered by electrons⁴⁵. Therefore, even a slight decrease in the potential barrier leads to a significant increase in the tunneling current, which is observed as a strong signal upon capturing cadaverine.

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disadvantage of this mathematical model is that although one can accurately describe $I-V$ characteristics using Eq. 3, detailed information, including the electrostatic potential profile of the transport channel, cannot be obtained. This is because of the WKB approximation for electron transmission functions used to derive the model in the first place.

**Transient response characteristics: adsorption-desorption dynamics**

Measurement results indicated that the dynamics of a junction resistance drop were governed by a quasi-irreversible process of cadaverine molecules forming hydrogen bonds with the linker end group and producing a hybrid molecular bridge. Such a typical chemisorption process had been previously described mathematically by Elovich’s equations, and the desorption process can be described by the Polanyi–Wigner equation, which describes the adsorbed gas molecule being released back into a gaseous form, ideally consisting of complete breaking of hydrogen bonds between the target gas molecules and the linker end group. Figure 7b shows the normalized conductance response of the sensor to one period of exposure of the nanogap device to cadaverine and its subsequent removal, curve-fitted to adsorption-desorption models.

**Selectivity of the gas response**

To investigate the cross-sensitivity of the sensor, we exposed the device to analytes with different molecular end groups, commonly found VOCs and gases such as helium, hydrogen, and CO$_2$. We define the sensor response as the normalized junction resistance drop in the steady-state after exposure to the analyte. Figure 8a shows the sensor response when exposed to these analytes compared to that when exposed to the intended target gas, cadaverine. Figure 8b shows the quantitative response of the nanogap sensor when exposed to analytes with different molecular end groups. Measurements revealed an $R_{\text{OFF}}/R_{\text{ON}}$ ratio of more than 3 orders of magnitude when exposed to only 40 ppm cadaverine, whereas a maximum $R_{\text{OFF}}/R_{\text{ON}}$ ratio of ~1 order of magnitude was observed when exposed to much greater concentrations of the other analytes. The concentration of the VOCs and other gases was maintained at greater than 1000 ppm. The response to moisture was ~2 orders of magnitude; however, the relative humidity was maintained at ~80% (~20,000 ppm). To measure the device response in the presence of other gases, we flooded the test chamber with the specific test gas and then monitored the resistance drop of the sensor. After observing the steady-state response to a certain target gas, the sensor was removed from the test chamber, and the analyte molecules were allowed to naturally escape. This unassisted recovery was monitored electrically by continuously probing the junction resistance of the sensor, and complete recovery was assumed to have been achieved when the junction resistance was measured to be at its original value (as measured before exposure to the gas). Following this, the sensor was subsequently exposed to the next target gas. These results suggest highly selective sensor action against the most commonly found VOCs. The limited sensitivity to other gases may be due to smaller molecules or even moisture reducing the air gap between the electrodes due to chemisorption.

**Conclusions**

We present a new class of chemiresistors based on the capture of gas molecules within a nanoscale gap. We...
fabricated gas-sensing devices with gold electrodes separated by an ~6 nm nanogap functionalized with a fully conjugated terphenyl linker molecule (4-((4-mercapto phenyl)ethynyl)phenyl)ethynyl)benzoic acid) for electrostatic capture of cadaverine gas. We demonstrated ultralow-power resistance switching in batch-fabricated nanogap junctions upon detection of the target analyte—cadaverine. The stand-by power consumption was measured to be less than 15.0 pW, and the $R_{OFF}/R_{ON}$ ratio was more than eight orders of magnitude when exposed to ~80 ppm cadaverine. A phenomenological electrical model of the device is also presented, in good agreement with experimental observations. The cross-sensitivity of the gas sensor was tested by exposing the device to some of the commonly found VOCs and other atmospheric gases. The experiments revealed highly selective sensor action against most of these analytes. These batch-fabricated sensors consume ultralow power and demonstrate high selectivity; therefore, they are potential candidates for sensor application in power-critical IoT applications and can fulfill other low-power sensor needs. Given the dynamic response of the sensor, we believe that it can be used for quantitative measurements proportional to the target gas concentration. This enhances the possible impact of this technology toward the development of selective sensors based on this detection principle.

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A.B. was responsible for device design, fabrication, testing and characterization, modeling of sensor action, manuscript writing and figure preparation. S.H.K. helped in device fabrication, testing and manuscript preparation. A.Bu. helped in GC-MS of the target gas and associated calculations. S.N. helped in fluorescence imaging. S.B. and R.L. helped in the synthesis of chemicals.
required for sensor action, K.H.K. helped in sensor testing, and K.H. procured funding, helped in the experimental setup, preparation and proofreading of the manuscript. C.H.M. helped throughout the testing, fabrication, modeling of sensor action, analysis of results, and preparation and proofreading of the manuscript.

Conflict of Interest

The authors declare no competing interests.

Supplementary information

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