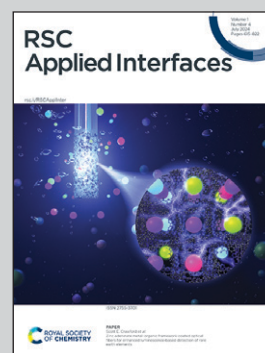


Showcasing research from Dr. Alpana Nayak's laboratory, Department of Physics, Indian Institute of Technology Patna, Bihar, India.

Artificial nociceptor using an Ag/Ag₂S/Pt atomic switch

Pain is a crucial sensation for organisms to adapt to dynamic environments, hence a key capability for realistic machine intelligence. We present an Ag/Ag₂S/Pt atomic switch as an artificial nociceptor, where voltage-dependent conductance modulation mirrors stimulus-dependent pain sensation. The voltage-dependent switching time, governed by different rate-limiting processes, parallels the varying response times of nociceptors. Quantized conductance levels encode distinct pain intensities, analogous to NMDAR channel openings in nociceptor networks. This work highlights signal-dependent conductance modulation in the Ag/Ag₂S/Pt atomic switch, mimicking biological pain pathways and enhancing neuromorphic defense mechanisms.

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Artificial nociceptor using an Ag/Ag₂S/Pt atomic switch†

Anwasha Mahapatra and Alpana Nayak *

Pain is an essential sensation for any living organism to thrive in the everchanging environment, and hence an important aspect to be added to the capabilities of realistic machine intelligence. Here we present an Ag/Ag₂S/Pt atomic switch as an artificial nociceptor whose voltage dependent conductance modulation is similar to that of a stimuli dependent pain sensation. Different rate limiting processes for the voltage dependent switching time illustrate its analogy to varying response times for different types of nociceptors. Distinct pain levels have been encoded in terms of quantized conductance, which is associated with opened NMDAR channels of the nociceptor network. This work emphasizes the signal dependent conductance modulation of the Ag/Ag₂S/Pt atomic switch in order to mimic the biological pathway for pain sensation that will advance the defence mechanism in neuromorphic devices.

A. Introduction

The integration of artificial intelligence and robotics has paved the way for machines to interact with and navigate the physical world with increasing sophistication.¹ However, the absence of a mechanism to perceive and respond to potentially harmful stimuli limits their ability to operate in a dynamic and unpredictable environment. Recently, artificial nociceptors utilizing diffusive memristors² with maladaptive³ responses have garnered a lot of attention in the field of neuromorphic devices. In response to potentially or actually harmful stimuli (like harmful chemicals, and high temperatures²/pressure⁴ or light⁵), specialized sensory neurons called nociceptors generate electrical impulses that are transmitted to the central nervous system. The activation of N-methyl-D-aspartate receptors (NMDAR) has been linked to elevated and elongated pain sensitivity in peripheral and central neural systems, according to recent biological research.⁶ When the noxious signal is strong enough to cause NMDAR channel activation in the post synaptic terminal, the network becomes more sensitive with less threshold value for activation, responding more potently than it would in a normal state. Similar activation modulation occurs in atomic switch devices when the conductance value becomes greater than the first quantization of conductance⁷ *i.e.*, $1G_0 = \frac{2e^2}{h}$.⁸ An atomic switch is a nano-ionic device that shows

a resistive switching memory effect^{9–11} at the atomic scale. Silver sulphide (Ag₂S), a silver chalcogenide compound, is an ionic crystal that exhibits mixed ion-electron conductivity due to the combined contribution of electrons and interstitial silver ions in the conduction mechanism.^{12,13} With sulphur atoms rigidly sitting in a fixed place, the silver atoms are mobile in the solid electrolyte participating in the filament formation, eliminating the problem of parasitic reaction, as occurs in oxygen-based systems. Ag₂S possesses its acanthite state (α -Ag₂S) with a band gap around 1 eV, and behaves like an n-type semiconductor at the ambient temperature.¹⁴ α -Ag₂S is a polymorph with a monoclinic crystal structure. On the other hand, the octahedral and tetrahedral interstitial lattice points in the high-temperature argentite (β -Ag₂S) are taken by Ag⁺ ions, while the S atoms are organized in a rigid, slightly deformed BCC structure.¹⁵ β -Ag₂S is superionic due to the liquid-like behaviour of the silver sub-lattice, with a high ion mobility and electron conductivity (band gap 0.4 eV).¹⁶ In addition to temperature, an electric field can lead to the α -Ag₂S to β -Ag₂S phase transition.¹⁷ The heterostructure of an Ag/Ag₂S nano-crystal makes it a good candidate for resistive switching-based memory devices with temperature sensitivity due to the reversible process of an α -Ag₂S to β -Ag₂S transformation along with a conducting channel formation¹⁸ under an electric field. The Ag/Ag₂S/Pt structure is well established and well known for atomic switch applications with room temperature quantized conductance and neuromorphic hardware.¹⁹ In recent works, Ag/Ag₂S-based flexible memristors have shown excellent endurance (10⁵ switching cycles) and retention (10⁴ s) proving its suitability for device applications.²⁰ Ag₂S is also known for its temperature-dependent conduction modulation¹⁸ and optoelectronic

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properties.²¹ Along with ease of fabrication and good reproducibility, these properties make the Ag/Ag₂S/Pt atomic switch suitable for bio-inspired electronics.

In this work, we have mimicked the properties of the NMDAR pathway linked to a nociceptor using an Ag/Ag₂S/Pt atomic switch by applying different electrical signals similar to nocuous stimuli.²² A flexible top electrode, *i.e.*, a conductive atomic force microscope (C-AFM) tip has been used to realize the gap-type atomic switch structure.²³ This allows real-time and real-space observations of the process arising between the tip and the sample.²⁴ Synaptic delay in the NMDAR pathway²⁵ of the nociceptor network and its modulation with pain intensity has been related to the voltage dependency of the switching time of the atomic switch.

B. Methods

The sample Ag₂S was prepared by exposing a freshly polished silver plate to sulphur vapor at 130 °C for 15 min.²⁶ Field emission scanning electron microscopy (FESEM) and energy dispersive X-ray spectroscopy (EDS) were carried out to examine the surface morphology. For conductive atomic force

microscopy (C-AFM) based characterization, Bruker multimode 8 has been used in the contact mode with a feedback set point of 10 nN, and a current limit of 1 μA. An SCM-PIT-V2 tip is used as the Pt electrode. As the precipitation occurred on the surface and the tip moved back, the simultaneous change in the z-position of the tip was recorded with time. The morphology of the surface was imaged before and after every switching event. For the pulse measurements, an Agilent 5500 AFM and a Keithley 4200 parameter analyzer were used, which allowed higher current ranges (100 μA and 1 mA).

C. Results and discussions

The atomic switch is realized as the Pt coated AFM tip approached the Ag₂S thin film grown on the Ag substrate, as described in Fig. 1a. The AFM tip acts as a flexible top electrode, which allows precipitation on the surface, beneath the tip. One such observation has been illustrated in the Supplementary Information in Fig. S1.† Fig. 1b is the I-V response to bias sweeps of 0 V → 0.5 V → 0 V taken on 100 different positions on an Ag₂S surface. The histogram of the switching voltages obtained in these measurements is shown

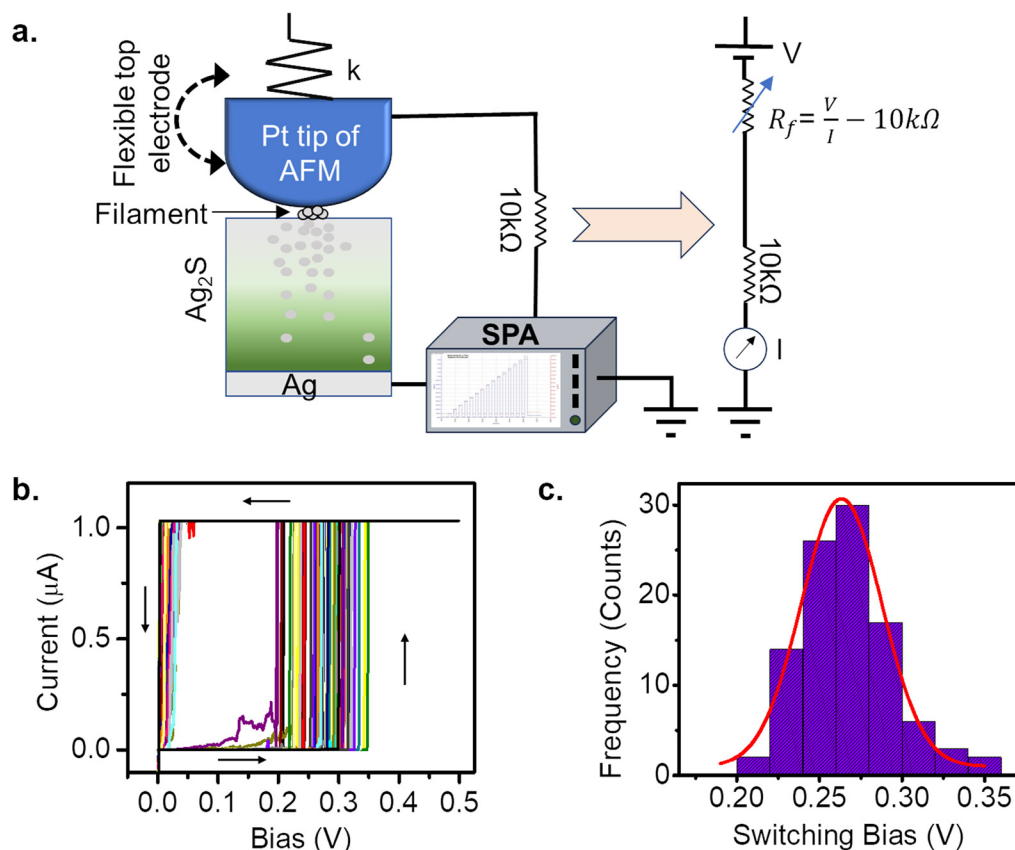


Fig. 1 (a) The Ag/Ag₂S/Pt atomic switch structure mimicked using a C-AFM tip (of spring constant $k = 2.5 \text{ N m}^{-1}$) gently approaching the Ag₂S thin film grown on an Ag substrate and the equivalent circuit diagram (b) switching spectra taken under same conditions for a 0 → 0.5 V → 0 V sweep at 100 different places on the Ag₂S thin film. (c) The histogram showing different switching voltages for the 100 sweeps, fitted with a Gaussian function (red) centered at 0.25 V with a standard deviation of 0.02 V.



in Fig. 1c. A Gaussian fit (red) to the data shows a peak at 0.25 V with a standard deviation of 0.02 V, which is expected due to the roughness of the Ag_2S surface.^{26,27} The FESEM and AFM images of the Ag_2S surface revealed crystal sizes of sub-micrometre order (Fig. S2†). The switching voltage (or the threshold V_{th}) is defined as a voltage value at which the atomic switch reaches a minimum of $1G_0$ (*i.e.*, $2e^2/h$). It is to be noted that the threshold value depends on the sweep rate in the case of voltage sweep applications and the pulse amplitude in the case of voltage pulse applications. In this measurement, a switching voltage of 0.25 V is reproducible for a given sweep rate as long as the conditions of the silver substrate (purity and roughness), sulphurization conditions (temperature and time/amount of exposure to the sulphur vapour) remain the same. Such an atomic switch can be regarded as an artificial nociceptor whose response to the amplitude and time of the stimuli, quantized levels of conductance, and desensitization with reverse stimuli are presented here. The conductance of the atomic switch has been used to quantify the strength of the pain signal generated by a nociceptor.

To draw the analogy of the atomic switch with a stimuli-dependent pain response of biological nociceptors, a pinched hysteresis for an applied sweep of 0 V \rightarrow 0.5 V \rightarrow 0 V \rightarrow -0.5 V \rightarrow 0 V has been obtained (Fig. 2a). The corresponding device structure is shown in the inset. The current value

jumped from sub-nano Amperes to tens of micro-Amperes at 0.25 V. The equivalent pain perception is expressed in terms of the pain intensity (*i.e.*, the system conductance, $2e^2/h$) as a function of stimuli intensity (*i.e.*, bias strength, V) in Fig. 2b. The navy-blue line indicates the response of a normal state (forward sweep), whereas the orange line corresponds to an injured state (backward sweep). In the normal state, the conductance values are low for sub-threshold voltages, and increase above $1G_0$ (in this case, $10G_0$) for higher voltages. In the injured state, the high conductance is retained even for sub-threshold values. This resembles the elevated pain response for non-nocuous stimuli (sub-threshold voltages) by an activated nociceptor. For example, gentle tapping on a burnt skin seems more painful than a normal skin sensation.

Another essential feature of a nociceptor is its maladaptive behaviour. Maladaptivity yields a heightened response to a stronger pain stimulus, which is important for defense mechanism to survive in an unpredictable environment. This behaviour is demonstrated in our system by measuring the current responses for increased amplitudes of the input pulses with a given width. One such measurement is shown in Fig. 2c, where we applied voltage pulses with amplitudes (V_p) ranging from 0.2 V to 1.6 V, for a pulse width of 2 ms. As the pulse amplitude increased linearly in steps of 0.2 V, the current response grew from 7.03 to 139.25 μA . The total conductance G_{total} has been calculated and plotted with

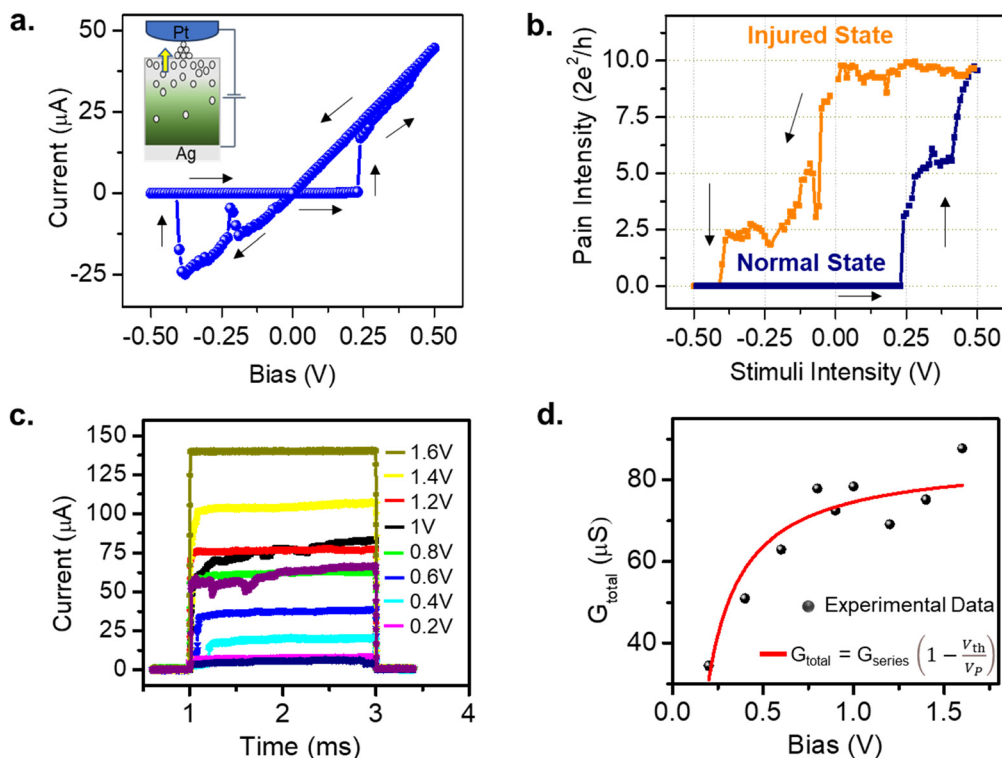


Fig. 2 (a) Pinched I-V hysteresis of the Ag/ Ag_2S /Pt system (inset) (b) conductance value representing the pain intensity as a function of the stimuli intensity *i.e.*, bias, where a heightened response is observed for the injured state. (c) The current response of the Ag/ Ag_2S /Pt system for square pulses of 2 ms width and varying amplitude (d) Total conductance of the circuit for the highest current values during each pulse, extracted from Fig. 2c fitted to the equation of a voltage divider (red).



respect to voltage amplitude in Fig. 2d. From the equivalent circuit diagram shown in Fig. 1a, we can write

$$\frac{1}{G_{\text{total}}} = \frac{1}{G_{\text{series}}} + \frac{1}{G_{\text{filament}}} \quad (1)$$

where, G_{series} and G_{filament} correspond to the conductance of the 10 k Ω resistance and the atomic switch, respectively. Since G_{series} is fixed, the increase in G_{total} is solely due to the formation of the Ag filament (G_{filament}); hence, we relate $G_{\text{filament}} \propto V_p$. As the filament forms, the conductance of the sample increases, and the voltage drop across the sample decreases.^{16,17} Once this voltage drops below V_{th} , further formation or growth of the filament stops, saturating the current value for a constant amplitude, as seen in Fig. 2c. Following the voltage divider rule, we write,

$$G_{\text{total}} = G_{\text{series}} \left(1 - \frac{V_{\text{th}}}{V_p} \right) \quad (2)$$

By fitting G_{total} vs. V_p data with eqn (2), we extracted the parameter $V_{\text{th}} = 0.127 \pm 0.015$ V. Hence, the Ag/Ag₂S/Pt nociceptor enters its injured state above a voltage 0.127 ± 0.015 V for a given pulse width of 2 ms.

Nociceptors are known to signal faster for stronger pain stimuli. The delay plasticity in pain sensation has been studied by measuring the variation of switching time with pulse amplitude. Fig. 3a shows a typical switching time measurement for a pulse amplitude of 1 V. The switching time is defined as the time required to reach $1G_0$ after the pulse initiation. This can be compared with the synaptic/response delay^{28,29} of a nociceptor. At the end of the pulse, the current decays exponentially (inset of Fig. 3a), similar to the NMDAR pathway's calcium current tail²⁵ that plays an important role to modulate the next signal response. Variation of the switching time (t_{sw}) with different pulse amplitudes (V_p) has been plotted in Fig. 3b, which shows an exponential decrease in switching time with increasing pulse amplitude. This is similar to the faster pain sensation with a heightened stimuli strength, as observed in a biological nociceptor. Further, the t_{sw} vs. V_p plot shows two distinct slopes indicative of two different rate-limiting mechanisms.^{30–32} Region 1 is found to be the nucleation-limited process^{30,33,34} and region 2 is limited by the hopping rate. Similarly, in biology, different ranges of response speeds are found in thermal, mechanical, chemical and silent nociceptors due to the heterogeneity of the constituent ion-channels in the nociceptor network.^{35,36}

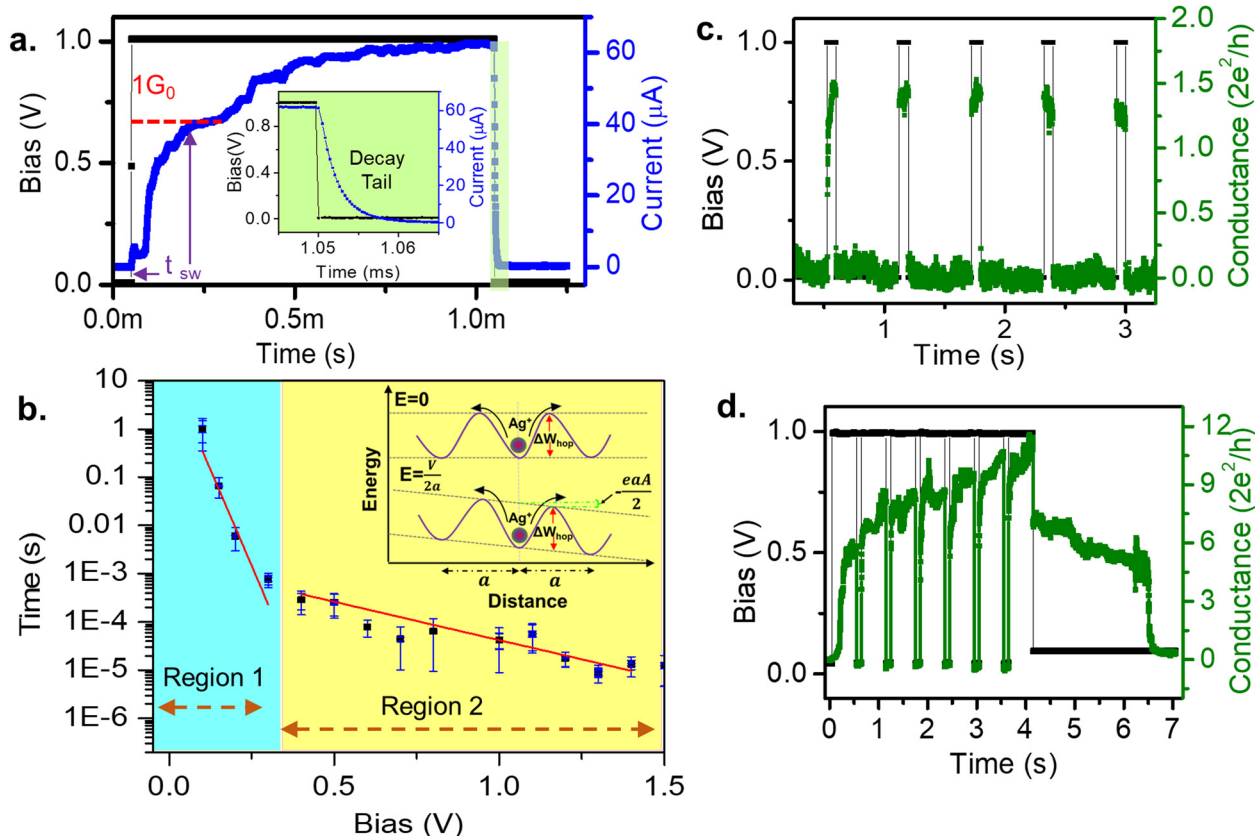


Fig. 3 (a) An example of switching time measurement using a square pulse of 1 V amplitude with a 1 ms width; the current response takes switching time t_{sw} to reach the $1G_0$ value. The decay tail at the pulse end is similar to that of the calcium tail current of NMDAR (inset). (b) Switching time is plotted as a function of the pulse amplitude yielding two distinct regions corresponding to two different rate limiting processes. The decrease in the hopping barrier due to the applied voltage is shown in the inset. (c) Saturation of low-valued conductance for small signals applied with a long interval, replicating the relaxation dynamics of a nociceptor. (d) Increased conductance of the device for wide pulses applied in short intervals, mimicking elevated and prolonged pain sensation for the nociceptor exposed to longer stimuli before relaxation.



First, the nucleation rate (J_n) in the Ag/Ag₂S/Pt nociceptor has been determined by using an atomistic model for the electro-crystallization under a positive applied potential to the Ag substrate, given by

$$J_n = N_{\text{Ag}^+} Z_0 \Gamma \mathbb{N} \frac{k_B T}{h} \exp\left(-\frac{\Delta G + \phi(N_c)}{k_B T}\right) \exp\left(\frac{(N_c + \alpha)eV}{k_B T}\right) \quad (3)$$

where N_{Ag^+} , Z_0 , Γ , and \mathbb{N} are the number of Ag⁺ ions, the number of active sites, the Zeldovich factor, and the transmission probability, respectively. $k_B T$, ΔG , $\phi(N_c)$, and eV represent the thermal energy, free activation enthalpy of the charge transfer reaction at $V = 0$, specific surface energy of the critical nucleus, and applied voltage potential, respectively. N_c denotes the number of atoms in the critical nucleus and α represents the charge transfer coefficient. h and e are Planck's constant and the elementary charge, respectively. For the measurements carried out under the same thermodynamic conditions, we can assume $J_{n0} = N_{\text{Ag}^+} Z_0 \Gamma \mathbb{N} \frac{k_B T}{h} \exp\left(-\frac{\Delta G + \phi(N_c)}{k_B T}\right)$ to be constant. Hence eqn (3) reduces to

$$J_n = J_{n0} \exp\left(\frac{(N_c + \alpha)eV}{k_B T}\right) \quad (4)$$

yielding the relation between the switching time governed by nucleation and pulse voltage as

$$t_n = t_{n0} \exp\left(-\frac{(N_c + \alpha)eV}{k_B T}\right) \quad (5)$$

where $t_{n0} = 1/J_{n0}$. The slope of region 1 is calculated as

$$m_{\text{neuc}} = -\frac{e(\alpha + N_c)}{k_B T} = -27.4 \pm 2.7$$

From this slope, we calculated values of $\alpha = 0.7$ and $N_c = 0$, attributing the superionic β -phase of the Ag₂S formed under the high electric field during the switching process.^{13,37}

Next, for estimating the Ag⁺ ionic hopping current (J_h) we have used the Mott–Gurney law³⁰ given by

$$J_h = 2zeca f \exp\left(-\frac{\Delta W_{\text{hop}}^{(0)}}{k_B T}\right) \sinh\left(\frac{aze}{2k_B T} E\right) \quad (6)$$

Here, z is the charge number (For Ag⁺, $z = 1$), c is the ion concentration, and f is the attempt frequency. $\Delta W_{\text{hop}}^{(0)}$ is the barrier height of the resistive layer with no applied bias, a is the hopping distance and w denotes the dimension over which the voltage drops. The average electric field that results from the voltage V dropping over a specific distance w , $E = V/w$, is what propels the motion of the ions. Since the electric field experienced by the moving ion spatially fluctuates, the use of the average field is more justified than the Lorentz field approach.³⁸ Hence, under an applied electric field, the hopping barrier is $\Delta W_{\text{hop}} = \Delta W_{\text{hop}}^{(0)} \pm \frac{1}{2} zeaE$ (Fig. 3b, inset). This leads to a hyperbolic sine term in eqn (6).

Under constant thermodynamic conditions, we can consider that $J_{h0} = 2zeca f \exp\left(-\frac{\Delta W_{\text{hop}}^{(0)}}{k_B T}\right)$ is a constant value. Thus eqn (6) reduces to

$$J_h = J_{h0} \sinh\left(\frac{azeV}{2k_B T w}\right) \quad (7)$$

resulting in a voltage dependency of the switching time given by

$$t_h = t_{h0} \sinh^{-1}\left(\frac{azeV}{2k_B T w}\right) \quad (8)$$

Region 2 of Fig. 3b matches the corresponding slope for hopping as calculated by

$$m_{\text{hop}} = -\frac{aze}{2k_B T w} = -2.5 \pm 0.7$$

in our experiment. Typically the average value of a is 0.3–0.5 nm and w is 1.5–2.5 nm.³⁰ Hence, for hopping being solely the rate-limiting process, the condition is $\left|m_{\text{hop}} \times \frac{2k_B T}{e}\right| < 0.2$, which is of value 0.12 in our case. Overall, we can draw the analogy that the decrease in the switching time with increased pulse amplitude is a fingerprint for the nociceptor-like behaviour of the atomic switch. This indicates the greater the strength of the nocuous stimuli is, the faster the nociceptor indicates the pain sensation.

Further, an interesting analogy can be drawn from the relaxation dynamics of a nociceptor. If a stimulus arrives before the system retrieves from the previous injury, the pain sensation is found to increase. Moreover, a prolonged stimuli heightens the pain sensation. Both these features can be seen in our system (Fig. 3c and d). In Fig. 3c we have plotted the response for a pulse train consisting of five pulses with a 1 V amplitude, 100 ms width, and 750 ms interval. The resulting conductance value was 1.5 G_0 for each of the pulses. After every pulse end, a sufficient time interval of 750 ms allowed the system to relax back to its initial low conductance state, thus not affecting the response for the following pulse. On the other hand, Fig. 3d depicts the response for seven pulses of amplitude 1V and width 500 ms applied at 100 ms intervals. Note that for the same amplitude, the conductance value reached up to 6 G_0 for the first pulse, in contrast to the 1.5 G_0 value in Fig. 3c, due to the increased width of the signal. Again, due to the short interval of 100 ms between the consecutive pulses, the next pulse arrives before the system relaxes back to its initial conductance value. Thus, for the consecutive pulses, the conductance values increased from 8 to 10.3 G_0 . This successfully mimics the heightened pain sensation (*i.e.*, conductance) with frequent and prolonged stimuli (*i.e.*, frequency and width of the pulse) of the biological nociceptor.^{4,5}

Room temperature quantization of conductance⁷ is one of the most intriguing aspects of the atomic switch,³⁹ as successful observations of 1 to 10 G_0 have been reported.⁴⁰



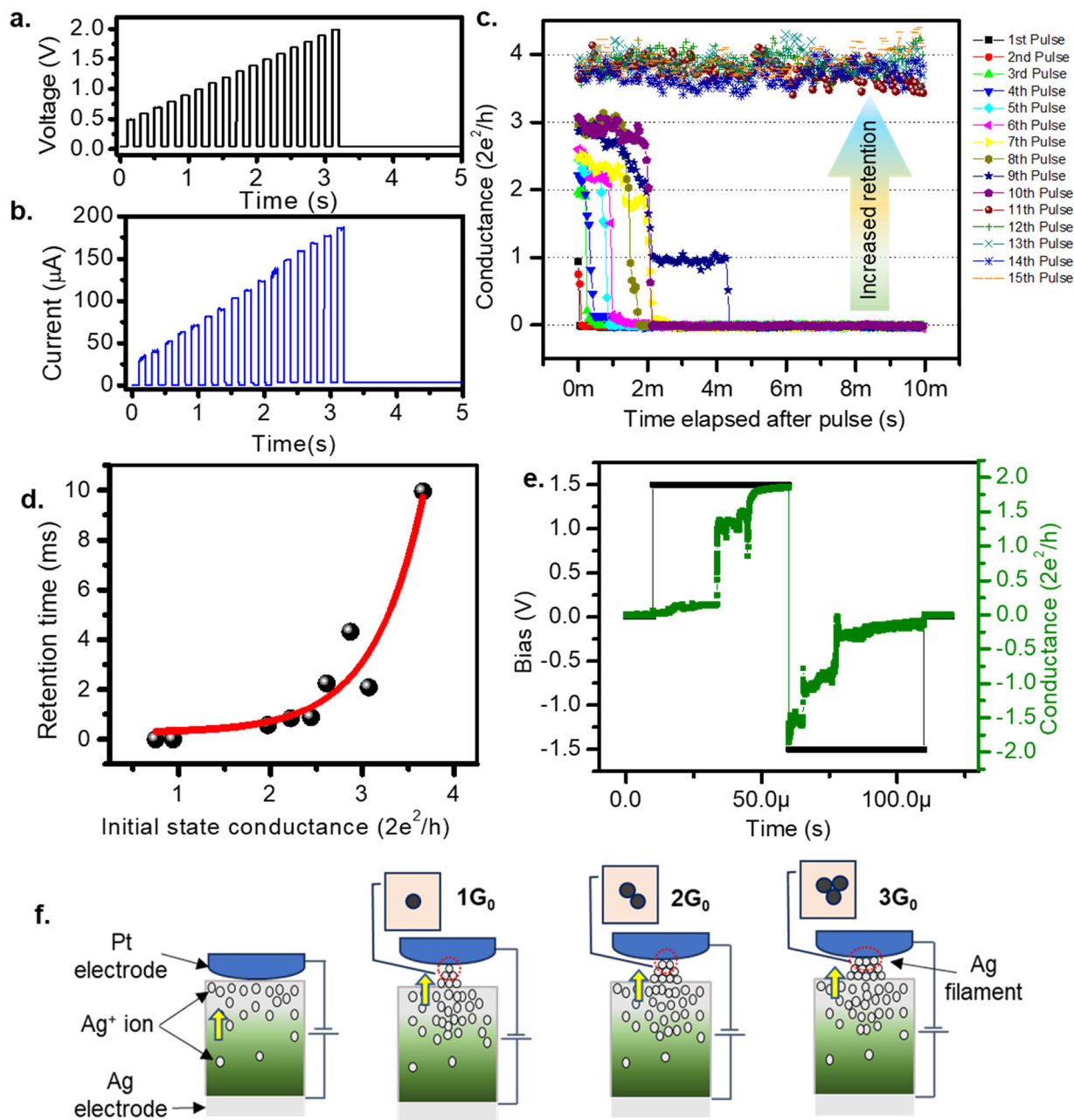


Fig. 4 (a) Pulse train applied to the system and (b) its response in terms of the current value with (c) the measured conductance values during the interval between each pulse. (d) Retention time as a function of the conductance of the state fitted to an exponentially growing function (red). (e) The retained conductance of the system for an achieved state with $2G_0$ that dissolves upon negative bias application. (f) Different conductance states arising due to different numbers of 1D opened conduction channels connecting the electrode.

Similarly, in biology, the quantity of NMDAR channels in an active region is documented up to ten units.⁴¹ Therefore, we suggest that discrete pain levels, represented by opened NMDAR channels in biology, can be analogous to quantized conductance states in an atomic switch. Fig. 4a shows the input voltage of fifteen pulses with successive increments in amplitude from 0.5 V to 2 V, in steps of 0.1 V, with an interval of 10 ms. The corresponding current response is shown in Fig. 4b. The current reaches a high value during the pulse and decays to a low value in the interval after each pulse. The

peak value of the current, during the pulse, increased from 30 to 185 μA . The low current observed after each pulse was not the initial value, instead it grew from 0.003 to 3.589 μA for the tenth pulse, which was retained for an observation time of 2 s after the end of the fifteenth pulse. These retained current values when expressed in terms of conductance (Fig. 4c) display clearly how the conductance decayed during the interval between two successive pulses. The initial conductance ($1G_0$) after the first two pulses spontaneously decayed at the end of the pulse. This indicates the system



only sensed the stimuli, without retaining any memory. When the system was raised to a conductance state of $2G_0$ or higher, the decay rate was slower and eventually leveled off to a permanent retention value of $4G_0$ at the end of the tenth pulse, which was retained for 2 s of observation time even after the fifteenth pulse. Since the conductance level is analogous to the pain level, retention of a higher conductance means a prolonged pain sensation for heightened pain stimuli. These conductance values and their corresponding retention times are plotted in Fig. 4d which is fitted to an exponential growth function (shown by the red line). This plot indicates that the states with conductance values $\leq 1G_0$ tend to decay instantaneously. Achieving a conductance level $\geq 2G_0$ is necessary for the system to enter a prolonged pain sensation (*i.e.*, a retained conductance value). The stability of a higher G_0 state is further verified by observing the conductance values for input of two successive pulses of opposite polarity with an amplitude of 1.5 V and width of 50 μ s (Fig. 4e). For the positive polarity, the conductance value rose to $2G_0$, which was retained until a negative polarity diminished the value back to initial one.

We hereby reasonably argue that the opening of one NMDAR channel in a nociceptor can be represented by $1G_0$ of our system. Flexibility with a greater number of pain levels is achieved by opening more than one NMDAR channels corresponding to $G > 1G_0$. In the atomic switch, the narrowest region of the conducting filament connecting the two electrodes resembles a quantum point contact.⁴⁰ For ballistic transport in this region, the Landauer theory gives $G = N_{\text{ch}} \beta G_0$, where N_{ch} is the number of 1D opened conduction channels connecting the electrode and β takes a value between 0 and 1 ($\beta = 1$ in our case). $N_{\text{ch}} = 1$ for a single atom contact and it takes values 2, 3, 4, *etc.*, for the corresponding number of atoms contacted as illustrated in Fig. 4f. Thus, the room temperature quantization of conductance in atomic switch provides the advantage of coding distinct pain levels depending on the signal strength, mimicking the NMDAR associated nociceptor of a mammalian nervous system.

D. Conclusions

In this work, we have proficiently demonstrated that an atomic switch can mimic the properties of a biological nociceptor. The conductance levels of the atomic switch are considered to be equivalent to the pain levels of a nociceptor. The high conductance ON-state of the atomic switch has been compared to the injured nociceptor, which signals elevated pain for sub-threshold values. The threshold is an important concept for implementing the maladaptive behaviour of the nociceptor, below which it signals no threats from the stimuli. Above the threshold value, the conductance, *i.e.*, the pain sensation of the system, increases with the increasing stimuli strength. We have estimated the threshold value in our system. The possibility of more than one rate-limiting process in the switching time indicates the resemblance to varying response times for

different nociceptors. The relaxation dynamics of our system mimic the heightened pain sensation (*i.e.*, conductance) with frequent and prolonged stimuli (*i.e.*, frequency and width of the pulse) of the biological nociceptor. To encode versatile high precision pain levels in the artificial nociceptor, quantized conductance states have been associated with opened NMDAR-channels of a biological network.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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