Electrochemically and Optically Readable Light Emitting Memories

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Electrochemical metallization memories based on redox-induced resistance switching have been considered as the next-generation electronic storage devices. However, the electronic signals suffer from the interconnect delay and the limited reading speed, which are the major obstacles for memory performance. To solve this problem, here we demonstrate the first attempt of light-emitting memory (LEM) that uses SiO2 as the resistive switching material in tandem with graphene-insulator-semiconductor (GIS) light-emitting diode (LED). By utilizing the excellent properties of graphene, such as high conductivity, high robustness and high transparency, our proposed LEM enables data communication via electronic and optical signals simultaneously. Both the bistable light-emission state and the resistance switching properties can be attributed to the conducting filament mechanism. Moreover, on the analysis of current-voltage characteristics, we further confirm that the electroluminescence signal originates from the carrier tunneling, which is quite different from the standard p-n junction model. We stress here that the newly developed LEM device possesses a simple structure with mature fabrication processes, which integrates advantages of all composed materials and can be extended to many other material systems. It should be able to attract academic interest as well as stimulate industrial application.

Numerous progresses have been achieved in the last several decades on discrete semiconductor decices, such as memories and light emitting diodes. Nowadays, both the performance and reliability of these devices have already become sufficient for use in practical applications. Future work may involve the research of devices that possess wider functionality and compatibility in a variety of data storage and processing. Notably, the demand of high-speed inter-chip and intra-chip link has dictated the developments of communication. The conventional electronic devices gradually approach their limit due to the increasing difficulties in controlling the carriers at shrinking sizes. Among several candidates for next-generation memory cells, resistive random access memory (RRAM) based on a simple two-terminal electrical switch has the potential to serve as a replacement for conventional memory structures due to its good switching properties, low power consumption and especially, three-dimensional multilayer stacking to achieve high density memory. The typical current-voltage (I-V) characteristics of the RRAM cells exhibit an extreme change in resistance between high resistance state (HRS) and low resistance state (LRS). One of the underlying mechanisms responsible for the RRAMs arises from the formation of conducting filament networks under the application of a positive voltage across the cell. Upon application of a negative bias, the conducting filaments are dissolved and the memory cell switches to the HRS. However, the reading process is still in a serial sequence, that is, the data is transmitted by scanning one bit after another. In application, there is much interest in developing the reading process in parallel, which will boost the data transmission rate dramatically. One of the promising ways in realizing parallelizing reading process is the optical interconnect. For example, when the light-emitting memories (LEMs) are turned on, data are received by optical sensor (e.g., CCD), i.e., regions without light emission are served as logic “0”, and regions with light emission are served as logic “1”. By monitoring the whole LEM array, optical sensor can sense and distinguish each light emission generated by LEMs at the same time. Therefore, the maximum data throughput can be greatly increased, which depends on the density of LEM array and the capability of optical sensor. To achieve the above described functionality, the integration of light emitter and memory device is definitely one of the feasible approaches. In this work, we made the first attempt of an inorganic LEM, which enables data communication via optical and electrical detections and provides both parallel reading process and high data bandwidth simultaneously.
The group III-nitride based semiconductors have been the focus of promising application in optoelectronics devices with its mature fabrication technologies and the composition dependent variable bandgap. The InGaN based light-emitting diodes (LEDs) and laser diodes (LDs) can cover the light spectrum from UV to visible regions. Although the InGaN based technology is well developed, it requires complex process involving multiple-quantum-wells (MQWs). In this work, the LEM was achieved by developing the graphene-insulator-semiconductor (GIS) light-emitting diode (LED) in tandem with a SiO2 based memory cell. The GIS-LEDs do not rely on p-n junctions, which enables to expand the list of usable semiconductors to materials that cannot be easily doped and can be economically fabricated without difficulties. We constructed the GIS-LEDs by using graphene layer as a metal contact on p-GaN substrate with an insulating SiO2 interlayer.

Among the group of III-nitride based semiconductors, GaN serves as the pillar of its family. Compared with the ternary InGaN alloys, the growth of good quality of GaN film is relatively easy, which can produce a strong light emission. Moreover, the combination of blue light with phosphor powders or quantum dots with suitable concentrations can generate the desired color of light. Based on this advantage, the device with blue-light illumination enables controllable color generation. In addition, the choice of p-type GaN film is because the barrier height of SiO2/p-GaN device is much lower than that of SiO2/n-GaN. Therefore, one of the benefits for using p-GaN as substrates is that it can generate an inversion layer to accumulate electrons near the SiO2/p-GaN interface. The accumulated electrons can easily recombine with the holes tunneling from graphene layer and create light emission. Notably, in order to generate the accumulation of electrons near the silicon oxide and semiconductor interface, the relative work functions between semiconductor and graphene is very crucial. This criterion is very important for the extension of our current work to other material systems. Moreover, the resistivity of a compensated GaN film is much higher than that of doped GaN, which is not suitable for the deployment in our light emitting device. Certainly, the LEM devices developed here can be extended to many other materials, including III–V or II–VI semiconductors, as long as the band alignment and work functions between the constituent elements fulfill the requirement for producing light emission.

Furthermore, in contrast to other frequently used conducting materials, graphene has been considered as a promising conducting material offering low cost and high transparency. Notably, the SiO2 based memory structure has been widely used in RRAM cells due to its insulating property and has been well developed in recent years. Because of the uniqueness of the LEM device shown here, our work could pave a milestone for the development of integrated optoelectronic devices and open up a new route for the application of optical communication, digital memories and recordable display panels.

Results

The graphene was synthesized by chemical vapor deposition (CVD) as described previously. According to published reports, the absorption of graphene is strongly correlated to the number of graphene layers. Therefore, we characterized the graphene layer by transmission spectrum, Raman spectrum as well as atomic force microscope (AFM). The obtained thickness and quality of the transferred graphene is evaluated from Raman spectrum, as shown in Figure 1a. The intensity ratio between 2D and G bands clearly reveals that our prepared graphene was single layer or bilayer. The AFM image shown here to give an overview of the surface morphology after the deposition of graphene onto the underlying p-GaN substrate, as shown in the inset of Figure 1a. Figure 1b shows the transmittance of the graphene in the wavelength range from 400 nm to 1100 nm. It is clear that the graphene can be used as a substitute for transparent electrodes. Moreover, to measure the mobility of graphene and show how the quality of graphene is, a field-effect device made with graphene/SiO2/p-Si was constructed by a back-gate structure. Based on the field effect data, we are able to characterize the quality of the graphene layer and to obtain the carrier mobility. Figure 1c shows the back-gate transfer characteristics with a low source-drain voltage (10 mV). The field-effect carrier mobility can be extracted from the linear regime as follows:

$$\mu = \frac{Lg_m}{W C_{ox} V_{th}}$$

where L and W are the channel length and the channel width, respectively, $g_m$ is the transconductance, $C_{ox}$ is the oxide capacitance and $V_{th}$ is the channel bias. It is found that the field-effect carrier mobility of the graphene is about 1200 cm$^2$V$^{-1}$s$^{-1}$.

The configuration of the LEM is demonstrated in Figure 2a. The SiO2 layer was deposited by a radio-frequency sputtering system. The
large-area CVD graphene was used as current spreading electrode, the Ni/Au contact was used as current collecting electrode (cathode) and the Ag contact was used as control electrode (anode). In addition to serve as the transparent conducting layer, graphene also works as a stable interlayer without affecting the redox reaction in the formation of metal filament networks. Moreover, because the tunneling SiO₂ layer only has a thickness of about 3 nm, the transferred graphene on top of the SiO₂ layer does not need the addition process for coating a metal film. It can avoid the damage of the thin SiO₂ layer and prevent leakage current. In our device geometry, the top Ag electrode works as a reflective layer and the emitted light was detected from the p-GaN side. If the graphene layer were replaced by coating other metal films with high reflectivity, it may affect the redox reaction in the formation of metal filament networks of the memory cell and induce leakage current. The inset of Figure 2a shows the photograph of the LEM. Figure 2b shows the photoluminescence (PL) spectrum of p-GaN under light excitation of 325 nm at room temperature. The PL spectrum is dominated by a blue emission peak at ~415 nm and a broadband yellow emission at ~550 nm. The emission peak at ~415 nm could be mainly attributed to the energy level below the conduction band at 170 meV (due to the interstitial Mg atoms) to the energy level above the valence band at 250 meV (due to the acceptor level)20. These doping-related energy levels strongly depend on the carrier density, that is, increase of the doping concentration results in the dominance of blue emission in the PL spectra21. Moreover, the annealing process for ohmic contact formation on p-GaN can also induce red shift in PL spectrum3,22. Our result obtained here is consistent with previous reports that for moderated Mg-doped GaN with the hole concentration approaching 10¹⁷ cm⁻³, the PL spectra are dominated by the blue emission.

Bistable switching performance of the memory cell is important to control the luminescence arising from of the LEM. Therefore, we investigated the performance of the memory cell first. Figure 3a shows the I-V characteristics of the Ag/SiO₂/graphene memory cell at room temperature and the inset of Figure 3a shows the schematics of Ag/SiO₂/graphene memory cell. As can be seen, the Ag/SiO₂/graphene memory cell was at the HRS initially and showed a low-current characteristics in the low-voltage range. When the applied voltage exceeded a certain value, ~3 V in Figure 3a, the injection current increased dramatically followed by an abrupt increase in the current flow and was switched from the HRS to the LRS, where the ON/OFF current ratio is about 10³. To prevent damage during the I-V measurements, we set a compliance current at 3 mA. The state transition is equivalent to the “writing” command in the digital storage devices and the ON/OFF current ratio promises a misreading probability in data access. According to previous reports, the conducting filament model with an electrochemical reaction is responsible for the resistive switching behavior3–8. Hence, the state of Ag/SiO₂/graphene memory cells can be switched between the HRS and the LRS using dc voltages. Switching performances between HRS and LRS were evaluated by performing the operation 100 times, as shown in Figure 3b. The current fluctuation at the high resistance state (HRS) may come from the incomplete dissolution of metal filament network24. It is clear that there was a little fluctuation of the HRS and LRS current levels and the ON/OFF ratio was quite stable. It is true that the on/off ratio is limited by the current compliance and is not representative of the device. The setting of the current compliance is used to prevent the device from damage. Indeed, a large on/off current ratio of more than 10⁴ can be obtained in our device. However, under such a high current operation, the reproducibility of the I-V curve becomes worse than the operation under the compliance current of 3 mA.

Figure 4a shows the I-V characteristic of the LEM at room temperature, which is similar to the I-V characteristics shown in Figure 3a. The current flow sharply increased by 2 orders of magnitude at a critical voltage (~8 V) and reflected the fact that the LEM view. As can be seen, the Ag/SiO₂/graphene memory cell was at the HRS initially and showed a low-current characteristics in the low-voltage range. When the applied voltage exceeded a certain value, ~3 V in Figure 3a, the injection current increased dramatically followed by an abrupt increase in the current flow and was switched from the HRS to the LRS, where the ON/OFF current ratio is about 10³. To prevent damage during the I-V measurements, we set a compliance current at 3 mA. The state transition is equivalent to the “writing” command in the digital storage devices and the ON/OFF current ratio promises a misreading probability in data access. According to previous reports, the conducting filament model with an electrochemical reaction is responsible for the resistive switching behavior3–8. Hence, the state of Ag/SiO₂/graphene memory cells can be switched between the HRS and the LRS using dc voltages. Switching performances between HRS and LRS were evaluated by performing the operation 100 times, as shown in Figure 3b. The current fluctuation at the high resistance state (HRS) may come from the incomplete dissolution of metal filament network24. It is clear that there was a little fluctuation of the HRS and LRS current levels and the ON/OFF ratio was quite stable. It is true that the on/off ratio is limited by the current compliance and is not representative of the device. The setting of the current compliance is used to prevent the device from damage. Indeed, a large on/off current ratio of more than 10⁴ can be obtained in our device. However, under such a high current operation, the reproducibility of the I-V curve becomes worse than the operation under the compliance current of 3 mA.

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was switched from the HRS to the LRS. The bistable switching performance of LEM is due to the bistability of Ag/SiO₂/graphene memory cell. Noting that the LEM was achieved by developing a tandem structure, in which the writing voltage was expended since the voltage was applied to the graphene/SiO₂/p-GaN GIS-LED and the Ag/SiO₂/graphene memory cell. Because the I-V characteristics depend on the corresponding HRS or LRS, it is expected that the electroluminescence (EL) intensity of LEM should be different when the memory cell was switched from HRS to LRS. When the LEM is at the HRS, there is no EL signal until ~8 V bias. However, when the LEM is at the LRS, the EL signal is detectable when the bias exceeds ~6 V. The emission state of LEM can be switched between the HRS and the LRS based on the I-V characteristics of the Ag/SiO₂/graphene memory cell, as shown in the inset of Figure 4a. Switching performances between the HRS and LRS were evaluated by performing the operation 100 times, as shown in Figure 4b. It is clear that the switching characteristics of EL signal is also quite stable.

Figure 5a shows the EL spectra centered at ~400 nm with different injection currents at room temperature. The emission peak is similar to that of the PL spectrum shown in Figure 2b. In order to clearly illustrate the underlying mechanism of the EL spectra, let us examine the band diagram of the graphene/SiO₂/p-GaN GIS-LED in the LEM. Figure 5b shows the band diagram of graphene/SiO₂/p-GaN GIS-LED at thermal equilibrium according to the data reported previously. Since the SiO₂ layer acts as an energy barrier, the carrier injected from graphene responsible for the radiative recombination of the light emission must tunnel through the SiO₂ layer, which is quite different from that of the common p-n junctions. When applying a positive bias on graphene, most of the voltage drops across the SiO₂ layer because graphene and p-GaN have relatively low resistance in comparison with SiO₂, as shown in Figure 5c. The applied electric-field will induce the band bending of GaN at the SiO₂/p-GaN interface and introduce an inversion layer with the accumulated electrons. Meanwhile, the Fermi level of graphene moves downward, which enables the holes in graphene to tunnel into the valence band of p-GaN and cause the EL signal. The origin of UV emission can be attributed to the recombination of the trapped holes in deep acceptor levels with electrons in the conduction band.

Figure 4 | (a) I-V characteristics of light-emitting memory (LEM). The inset shows electroluminescence (EL) intensity of LEM as a function of voltage at high resistance state (HRS) and low resistance state (LRS). (b) Switching behavior of LEM over 100 cycles.

Figure 5 | (a) Electroluminescence (EL) spectra of light-emitting memory (LEM) with different injection current. (b) Energy band diagram of graphene/SiO₂/p-GaN metal/insulator-semiconductor light-emitting diode (GIS-LED) under thermal equilibrium. (c) Energy band diagram of graphene/SiO₂/p-GaN GIS-LED under bias.
Discussion

To further understand the conduction mechanisms of LEM, we qualitatively analyze the relationship between the injected current \( I \) and the applied bias \( V \) of the Ag/SiO\(_2\)/graphene memory cell first. Figure 6 shows the logarithmic plot and linear fitting of HRS curve in Figure 3a. The I-V curve of HRS exhibits a linear relationship between injection current and bias with a slope of 1.15, which is very close to 1 and matches the Ohmic conduction\(^{27}\). The Ohmic conduction can be expressed as,

\[
I \propto V \exp\left(\frac{-E_a}{kT}\right),
\]

where \( E_a \) is the activation energy of electron. At the LRS, the injection current increased dramatically and saturated, which is dominated by the conducting filament model, as described in previous reports\(^{3-8}\).

Similarly to the case of Ag/SiO\(_2\)/graphene memory cell, we can deduce the conduction mechanisms of LEM from the typical plots of the I-V characteristics. As shown in Figure 7, the conduction mechanism of LEM is much more complicated and each HRS and LRS I-V curve can be divided into two portions. From 0 V to 7 V at the HRS, the LEM exhibits a logarithmic relationship between injection current and bias with a slope of 1.4. It is a consequence of the combination of the I-V characteristics of Ag/SiO\(_2\)/graphene memory cell and graphene/SiO\(_2\)/p-GaN GIS-LED, but the Ag/SiO\(_2\)/graphene memory cell dominates the conduction mechanism at HRS. When bias is above 8 V at the HRS, an abrupt increase in current occurs, and the I-V characteristic exhibits a typical Fowler–Nordheim (FN) tunneling curve with a negative slope as shown in Figure 7b. According to the previous reports\(^{28}\), within the Simon’s approximation, in the limit of a large applied bias, the FN tunneling can be written as,

\[
I \propto V^2 \exp\left(-\frac{4d \sqrt{2m^* \phi}}{3\hbar V}\right),
\]

where \( d \) is the barrier width, \( m^* \) is the electron effective mass and \( \phi \) is the tunnel barrier height. However, for the LEM at LRS, the I-V characteristics can be well fitted by direct tunneling and FN tunneling at low applied bias (\(<4 \text{ V}\)) and high applied bias (\(>4 \text{ V}\)), respectively. The tunneling relationship for direct tunneling can be expressed by the following equations\(^{28}\)

\[
I \propto V \exp\left(-\frac{2d \sqrt{2m^* \phi}}{\hbar}\right),
\]

The fitting I-V characteristics of LEM at LRS is shown in Figure 7c, which is in good agreement with the theoretical prediction. Linear decrease at low bias region and linear increase at high bias voltage indicate that the conduction mechanisms in LEM at LRS are based on direct tunneling and FN tunneling at low applied biases and high applied biases, respectively. This result is reasonable because FN tunneling is a conduction mechanism, where the carriers tunneling through a triangular barrier requires a high electric-field. The above analysis of the conduction mechanism provides a firm evidence to support the fact that the tunneling process plays a very important role for the EL in our LEM devices.

Finally, let us provide a discussion for the optimization of the on/off current ratio and the reduction of the threshold voltage for the LEM device. The thickness of the SiO\(_2\) layer above the graphene layer is 50 nm and is about 3 nm below the graphene layer. Clearly, the threshold voltage and the HRS current will increase and decrease with respect to the increase of the thickness of the upper and lower SiO\(_2\) layers, respectively. For instance, in the LHS, the current is
dominated by the tunneling through the lower SiO₂ layer. As shown in equations (3) and (4), the relationship between the tunneling current and the SiO₂ thickness follows an exponential function. A small increase in the SiO₂ thickness can drastically reduce the magnitude of the tunneling current. According to the above description, the key factor to have a higher ON/OFF ratio and lower threshold voltage for our LEM device does strongly depend on the thickness of the upper and lower SiO₂ layers. The decrease of the thickness of the lower SiO₂ layer will result in a lower threshold voltage and a higher current in the LRS. Consequently, a smaller thickness of the lower SiO₂ layer without leakage current is the best choice. On the other hand, the decrease of the thickness of the upper SiO₂ layer will increase the current in the HRS and decrease the threshold voltage. Therefore, the thickness of the upper SiO₂ layer has to be optimized in order to have a higher on/off ratio and a lower threshold voltage. Another excellent alternative to have a higher on/off ratio and lower threshold voltage is to replace the GIS-LED developed here by the well mature and commercially available III-V compound LEDs.

In summary, we have developed a new LEM with bistable luminiscence control. The success of LEM mainly arises from the integration of light emitters and memories with a graphene interlayer, which combines several advantages of the composed materials. The LEM was achieved by developing the GIS-LED in tandem with a SiO₂ based memory cell. The underlying mechanism of the observed EL spectra can be interpreted quite well by the carrier tunneling through the insulating layer, which is very different from the standard p-n junction model. The feasibility of the new GIS-LED is made possible by the excellent properties of conductivity and transparency possessed by graphene. It is emphasized that the new LEM enjoys a simple structure and can be fabricated by using mature facilities. It is ready to be extended into many other material systems for practical applications. In view of the unique features demonstrated by the integration of light emitters and memories, the approach shown here may open up a new route for the development of integrated optoelectronic devices and pave a milestone for optical communication, digital memories and recordable display panels.

Methods
We grew graphene on copper foils at temperature up to 1000°C by CVD process and the mixture of methane and hydrogen was used as carbon source. At the temperature around 1000°C, the carbon atoms from gaseous reactants are deposited onto the metal substrate via chemical adsorption. Poly(methyl methacrylate) (PMMA) was spun-coated on the graphene sheet. The layer, which was transferred onto p-GaN, was annealed at 250°C for 10 min. The device structure of LEM consists of Ag/SiO₂/graphene/SiO₂/p-GaN. The p-GaN was doped by Mg with the hole concentration in the range of 6 × 10¹⁸ cm⁻³. Initially, p-GaN wafer was rinsed in acetone, isopropanol and DI water. Ni/Au was firstly deposited on p-GaN with annealing to obtain a ohmic contact. The first SiO₂ layer with a thickness of 3 nm was grown on p-GaN via RF sputtering after the formation of Ni/Au ohmic contact. The graphene layer was then assembled by a transfer process on p-GaN. The second SiO₂ layer with a thickness of 50 nm was sequentially deposited on top of graphene. The LEM was completed by the deposition of Ag electrode to serve as anode.

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Author contributions
C.W.C. and Y.F.C. designed the concept. W.C.T. synthesized the material. M.L.L. and Y.F.C. supervised the project and finalized the manuscript.

Additional information
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