Atomic scale memristive photon source

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Abstract

Memristive devices are an emerging new type of devices operating at the scale of a few or even single atoms. They are currently used as storage elements and are investigated for performing in-memory and neuromorphic computing. Amongst these devices, Ag/amorphous-SiO\textsubscript{x}/Pt memristors are among the most studied systems, with the electrically induced filament growth and dynamics being thoroughly investigated both theoretically and experimentally. In this paper, we report the observation of a novel feature in these devices: The appearance of new photoluminescent centers in SiO\textsubscript{x} upon memristive switching, and photon emission correlated with the conductance changes. This observation might pave the way towards an intrinsically memristive atomic scale light source with applications in neural networks, optical interconnects, and quantum communication.

Introduction

Compact on-chip photon sources are of great interest to the scientific community. Ideally, such light sources offer a compact footprint, low power consumption, are operated electrically and are compatible with the standard CMOS fabrication process, leading to high integration densities and energy-efficient operation at reduced cost\textsuperscript{1}. Such compact electrically-driven photon sources would be much needed within integrated circuits. For instance, for optically interconnecting processor and memories\textsuperscript{2}; or to optically communicate a sensing event\textsuperscript{3}. In quantum communications, they could act as on-chip single photon sources. In the field of neuromorphic computing, they could be used to communicate a memristive state\textsuperscript{4}.

Research in compact electrically-driven photon sources has led to quite a few innovative solutions over the past years. For example, quantum dot based light sources already deliver excellent emission efficiency with controlled spectra\textsuperscript{5,7} but require complex integration processes for fabrication. Electrically-driven light emitting tunnel junctions can be extremely compact and versatile\textsuperscript{8–11}. However, the vertically stacked architectures still require a large injection area\textsuperscript{8} and fine control over the fabrication of a thin oxide barrier, whereas the in-plane architectures require advanced nanofabrication\textsuperscript{9} or stochastic arrangements\textsuperscript{10} that are not scalable.

In parallel, innovative atomic scale electronic devices have emerged notably with the advent of memristors\textsuperscript{12}. Memristive devices are attractive for downscaling, as the operation only relies on the movement of a few atoms\textsuperscript{13–15}. The low energy, high-speed operation\textsuperscript{16} makes memristors suitable for high-density storage\textsuperscript{17}, in-memory computing, and neuromorphic computing\textsuperscript{18}. Interestingly, these devices may be advantageously merged with optical functions: memristively controlled optical switches\textsuperscript{4} and photodetectors\textsuperscript{19} have been introduced. Yet, so far, the photonic operation of a memristor relies on external or co-integrated photon sources\textsuperscript{20}.

In this paper, we introduce an atomic scale memristive device capable of emitting photons during resistive switching, superseding thus the need for an external optical source. Our device features the compact footprint of transistors and compatibility with the emerging
memristive technology. The device is based on transient-mode electroluminescence (EL) triggered by memristive switching. More precisely, near-infrared emission occurs within the gap of an in-plane Ag/amorphous-SiO\textsubscript{x}/Pt memristive junction when the resistance state changes. To enhance the emission, the apex of the Ag and Pt triangular electrodes are engineered to form a plasmonic nanoantenna. Our demonstration could potentially trigger a new conceptual paradigm for devices operating at the atomic level as electrical and optical functionalities may be embedded on the same nanoscale component. The paper thus addresses the challenge of downscaling photon sources similar to what we currently witness within electronics. Moreover, the new emitters offer not just light emission but are functional electrical devices on their own.

**Results**

**Sample description and working principle**

The atomic scale memristor photon source presented here, referred to as atomic photon source (APS), consists of an Ag/a-SiO\textsubscript{x}/Pt memristive switching in-plane junction fabricated on a glass coverslip. For more information on the fabrication process, the readers are referred to Supplementary Section I. An APS’s conceptual illustration and a scanning electron microscope (SEM) image are shown in Fig. 1a and b, respectively. By applying a voltage between the Ag and Pt electrode, resistive switching is achieved by the formation and dissolution of a conductive Ag filament\textsuperscript{13,15}. We discover that light is emitted within the gap between the Ag and Pt electrode during these critical forming and dissolution phases. To enhance the radiation and collection efficiencies, a plasmonic
nanoantenna is connected to the apex of the two quasi-triangular metal contact electrodes. The number of photons emitted by a single device is sufficient for detection with a standard CCD sensor, as shown in Fig. 1c (the setup is described in Fig. S1 of the Supplementary). Here an optical transmission image of one APS is overlaid with the photons detected during its operation (see Fig. S2 in the Supplementary for the two separate optical images).

A switching cycle of the APS is shown in Fig. 1d. It is characteristic for a memristive threshold switching device. Once the switching voltage of ~2.6 V is reached during the voltage ramping up, the device rapidly switches from a high resistance state to a low resistance state and reaches a set compliance current of $I_{CC} = 100$ nA. The low resistance state is volatile; as a result, the device will switch back to the initial high resistance state below a certain voltage threshold, which is around 0.3 V in this example. The release of photons is observed during the abrupt current change (highlighted by the black arrows), as illustrated in Fig. 1e and f. We attribute the transient-mode photon emission mechanism to luminescent defects—oxygen vacancy clusters—generated during resistive switching and thus filament growth and dissolution, which will be discussed below in more detail.

**Investigation of electroluminescence**

The transient-mode photon emission and its relations to the applied voltage and current are investigated by time-resolved electroluminescence (EL) measurements. The sample containing the device under investigation (APS) is placed on top of an inverted microscope and is imaged by using an oil immersion objective. An avalanche photodiode records the optical activity of the device. A voltage bias is applied to the device under test (APS) by an arbitrary waveform generator (AWG). The current and the emitted light are measured synchronously by an oscilloscope.

**Fig. 2 APS Electroluminescence Measurement.** a) The setup of the measurement. The APS sample is placed on top of an inverted microscope and is imaged by using an oil immersion objective. An avalanche photodiode records the optical activity of the device. A voltage bias is applied to the device under test (APS) by an arbitrary waveform generator (AWG). The current and the emitted light are measured synchronously by an oscilloscope. b) Upper panel: current evolution upon filament formation. The applied voltage is gradually ramped up to 7 V, shown in the black curve. At ~280 ms, the current increases abruptly from the noise floor of ~10 nA to several µA, shown in the blue curve. Lower panel: corresponding photon activity. A burst of photons can be seen upon the abrupt increase of current. Both the photon rate and accumulated photon counts are plotted. Red curve: binned photon counts with a size of 10 µs. Green curve: accumulated photon counts. Most photons are generated within the transient switching time at around 280 ms. c) Emission from filament dissolution, a burst of photons can be seen upon the abrupt current drop at around 12 ms. Photon counts are binned every 10 µs.
state (LRS) after the 5 V pulses. The upper panel displays voltage and current, and the lower panel in Fig. 3a shows the simultaneously acquired photon activity displayed for two binning times. Clearly, the device emits bursts of photons whenever an electrical stress is applied. It should be noted that unlike the previous experiments in Fig. 1d and
Fig. 2, no compliance current has been applied here. This mode of operation with a relatively high voltage (5 V) allows for repeated changes of the filament’s structure caused by electrical stress. More specifically, the relatively high voltage applied across the gap enables repeated Ag filament formation. Once the filament is formed, the transport of charges destabilizes the filament structure via local Joule heating\(^1\), and a renewed growth of the filament can take place. These repeating cycles driven by the motion of a few atoms during a voltage pulse lead to a current fluctuation. This current fluctuation can be seen more clearly in the zoom-in plot of a single voltage pulse in Fig. 3b and c. To correlate the optical activity with the change of current, we define the ratio of the current variation as \( R = \frac{I_t}{I_0 - \Delta I_0} \), which represents the rate of filament structural changes. A large \( |R| \) signals the morphological atomic reorganization of the conductive filament within the switching layer. The ratio is displayed in the bottom frame of Fig. 3b and Fig. 3c (green curve), together with the number of photons detected during the pulse (red curve). A correlation between the photon counts detected and \( R \) can be readily observed: the photon counts are high whenever \(|R|\) is large. There is barely any photon emission when \( R \) is close to 0. This is even clearer in the zoomed-in plot of the pulse shown in Fig. 3c, as highlighted by the grey dashed box. From the correlation of \( R \) and the photon counts, we conclude that such an electrically-induced structural change is systematically correlated with the emission and ultimately the cause for releasing the photons.

The APS presented here has been tested for \( \sim 100 \) cycles during which photon emission (electroluminescence) has been observed. It is worth noting that in each cycle the device is going through many filament formation and dissolution cycles even within a single electrical pulse excitation, therefore the photonic-emission (optical) endurance cannot be directly compared with the electrical endurance. Still, electrical endurance is one of the most critical measures of memristive devices\(^2\), and electrical endurance tests have previously been performed on devices with a similar material stack\(^3\). Those devices performed 50,000 consecutive cycles without a single failure.

The spectrum of the emitted photons taken during a series of 5 V, 5 ms pulses is shown in Fig. 3d. The APS features a broad emission spectrum spanning the near-infrared region with an emission peak at \( \sim 745 \) nm. The simulated quantum efficiency \( \eta_z \) as a function of the filament-antenna gap \( d \). Inset, the structure of the Ag and Pt nanoantenna used in the simulation. The filament is modeled as a cylindrical Ag rod with a radius of 0.5 nm.

Emission efficiency optimization

To optimize the quantum efficiency of the APS, a plasmonic nanoantenna at the tip of the APS electrode has been designed to engineer the electromagnetic environment. To that aim, we constructed a geometry tuned to the emission peak shown in Fig. 3d. The antenna dimensions have been optimized to maximize the number of photons collected by the objective beneath the sample. This is done by maximizing the radiation efficiency \( \eta_z \), the fraction of photons emitted towards the \( z \)-half-space. The simulated \( \eta_z \) of the fabricated APS is depicted in Fig. 4a. The inset shows the geometrical structure model. The simulation confirms a resonance peak at \( \sim 710 \) nm with a high radiation efficiency maximum of \( \sim 36\% \) as well as an efficiency above \( 20\% \) for a broad spectral range of 630 to 860 nm.
In addition, simulations with an Ag filament between the two electrodes are performed to estimate its effect on the optical quantum efficiency of the APS (details see Supplementary Section IV). The quantum efficiency of the APS is approximately proportional to the product of $\eta_z$ and the local density of optical states (LDOS) enhancement $L$ (details see Supplementary). In Fig. 4b, this factor $L \cdot \eta_z$ is investigated for different gap sizes between the filament and the silver antenna. As shown by the simulations, $L \cdot \eta_z$ only drops slightly with the presence of a filament. This point is discussed in greater detail in the Supplementary Section IV. We conclude that engineering the extremity of the electrodes is a necessary route to further improve the efficiency of the APS.

Investigation of the luminescence mechanism

To investigate the origin of the photon emission, a two-step analysis is conducted. Our hypothesis is based upon the creation of optically-active defect centers induced by the structural changes from resistive switching. To test this assumption, we perform photoluminescent (PL) measurements to investigate the presence of new species located in the gap after electrically stressing the APS. Second, various PL and EL measurements are performed in the presence and absence of silver, indicating that the luminescent centers likely originate from defects in the SiO$_x$ matrix rather than from the silver.

Investigation with photoluminescence

To locate the origin of the luminescence and to show that it stems from the gap, PL scans are performed on an APS before and after electrically activating the device. The APS is optically excited with a 515 nm diode laser. The details of the optical setup are given in Supplementary Section II. First, a PL confocal scan is carried out on a pristine device that has never been switched before. The image displayed in Fig. 5a features two dark triangular regions that are the metal contacts to the antenna structure.
bright photoluminescence background observed when the laser excites the matrix originates from native luminescent defects in the as-sputtered a-SiOx cladding layer, which is naturally oxygen-rich (see the XPS measurement in Supplementary Section V). After a few cycles of resistive switching, the PL confocal scan is repeated, and the result is shown in Fig. 5b. Compared to the as-sputtered SiOx, a much brighter PL emission center appears between the metal contacts, indicating that new luminescent sites are created in the gap after resistive switching. Please note that a lower laser power is applied for PL excitation in Fig. 5b to avoid excessive photobleaching. Also note that the PL spectrum does not perfectly match the EL spectrum since the contributing population and nature of the defects may differ between electron injection and photo-excitation. Similar EL and PL differences have been reported in the literature.

To confirm the presence of new species formed upon repetitive switching, PL spectra of the as-sputtered SiOx outside of the gap and in the gap after resistive switching are compared in Fig. 5c and d, respectively. The PL spectra reveal that there are two peaks centered at ~625 and 745 nm in both spectra. Furthermore, by comparing the two spectra, it is evident that the amplitude of the peak at 745 nm revealed by the decomposition is increased by a factor of 3 after switching compared to the spectrum of the SiOx cladding. This suggests that the PL emitted by the activated gap results from the additive contribution of new luminescence centers created in the oxide during the resistive switching process. The formation and aggregation of the oxygen vacancy clusters in forward-switching can be understood as follows: Intrinsic defects in deposited amorphous SiOx (such as wide O–Si–O bond angles) or strain introduced defects during filament growth and dissolution could act as precursors for oxygen vacancy clusters. These precursors lower the energy barrier to create oxygen vacancies, making the resistive switching more controllable with a reduced switching voltage.

Investigation under reverse-bias switching

To determine what type of luminescent sites cause the light emission in the APS, additional EL and PL measurements with a reverse-bias are performed on a pristine device (see Supplementary Section VII). Unlike forward-bias switching, Ag oxidation and Ag filament formation do not occur in reverse-bias switching. Hence the matrix remains free of Ag under this biasing condition. By conducting a reverse-bias switching measurement, the luminescence from metal species such as silver nanoclusters can be excluded.

Reverse biasing required much higher voltages and showed weaker photon emission (see Supplementary Section VII). Nevertheless, very similar EL and PL spectra compared to forward-bias switching are observed. Notably, a similar switching–induced PL spectral change is observed under both reverse- and forward biasing, which indicates the creation of luminescent sites in either operation mode by the same processes. We conclude that the origin of photons is the same for both forward and reverse bias. Correspondingly, as no silver is present in the gap during reverse switching, we argue that the light emission does not originate from silver nanoclusters but from defects in the SiOx matrix.

Discussion

Following the previous two sections, it is now understood that the luminescence stems from a structural change of the SiOx matrix within the gap. The role of the silver in the APS is to lower the switching voltage and to make the luminescence more controllable and intense. Here we discuss what luminescent sites are the most likely origin of the observed light emission and propose a photon emission mechanism.

In the case of reverse-biasing, the APS switches by a controlled dielectric breakdown in the SiOx matrix. Such mechanisms are known from OxRAM-type memristors, where operation is attributed to the creation and aggregation of oxygen vacancies. Whereas high voltages are required in the reverse-bias case, the creation and aggregation of oxygen vacancies into oxygen vacancy clusters also occur in the case of forward-bias experiments at lower voltage in the presence of the Ag filament. It has been reported that the introduction of metal species (Ag in our device) lowers the energy barrier to create oxygen vacancies, making the resistive switching more controllable with a reduced switching voltage.

The formation and aggregation of the oxygen vacancy clusters in forward-switching can be understood as follows: Intrinsic defects in deposited amorphous SiOx (such as wide O–Si–O bond angles) or strain introduced defects during filament growth and dissolution could act as precursors. These precursors lower the energy barrier for the creation of oxygen vacancies. Once created, the barrier to create additional oxygen vacancies closeby is further reduced, which leads to an aggregation of oxygen vacancies into oxygen vacancy clusters. Oxygen vacancies are
known to be luminescent upon electrical or optical excitations\textsuperscript{35}. However, the reported emission spectra typically feature signatures located in the UV to visible spectral regions and do not match our observations. Nevertheless, the creation and aggregation of oxygen vacancies into oxygen vacancy clusters generate locally silicon-rich regions within the SiO\textsubscript{x} matrix. Luminescence spectra of silicon-rich oxide matching our observed EL and PL peak wavelengths have been reported in various references\textsuperscript{36–39}. Thus, we attribute the photon emission of the APS to defects in locally silicon-rich regions\textsuperscript{39,40} formed by the creation and aggregation of oxygen vacancies into oxygen vacancy clusters upon electrical activation of the device.

Photon emission in these Si-rich clusters matching the EL and PL wavelength of the APS shown in Fig. 3d and Fig. 5d can be further divided into radiative recombination in Si nanocores and Si–O compound clusters\textsuperscript{37,41,42}. The latter appear either as clusters formed at the interface layer of Si nanocores\textsuperscript{37,38,41,43} or as distinct clusters in silicon-rich regions\textsuperscript{37,44,45}. The photon emission in the APS is more likely attributed to Si–O compound clusters for two reasons. First, the reported PL of Si nanoclusters matching our PL wavelength is generally attributed to the Si–O interface layer, whereas PL from the Si nanocore is only reported in the blue\textsuperscript{37,41,46,47}. Second, we conducted PL lifetime measurements (plots are given in Supplementary Section VIII), revealing that the switching-induced PL peak at ~875 nm features a lifetime of around 8 ns. As this measured lifetime is around ten times longer than the reported lifetimes from Si nanocore transitions\textsuperscript{41,42}, they can likely be excluded as the origin of the APS light emission. However, the lifetime of Si–O compound interfaces is reported to be \(10^4\) to \(10^5\) times longer than our measured decay\textsuperscript{37,41,47}. Such a discrepancy can be reasonably explained by the drastic shortening of the lifetime due to the Purcell effect, which is investigated by our LDOS simulation shown in Supplementary Section IV. Once created, this type of Si-rich defects can then be excited by tunneling electrons transported in the gap and can be radiatively recombined\textsuperscript{46} and thus represent the origin of light in the memristive atomic photon source.

The photon emission by memristive switching is more generic and not limited to the Ag/SiO\textsubscript{x}/Pt material system. We have already observed a much stronger photon emission from memristors with Ag/PMMA/Ag layer stacks. We believe the emission is also from the memristive switching-induced luminescent centers and atomic rearrangements of the Ag filament. However, the luminescence mechanism in this material system needs further investigation to confirm our hypothesis. In any case, the new memristive light emitters have the potential to enable many new applications e.g. related to neurons, which upon activation, fire and emit.

Conclusion

In conclusion, photon emission is reported during the resistive switching process of Ag/amorphous SiO\textsubscript{x}/Pt atomic scale memristors. Our investigations suggest that the emission stems from electroluminescent Si-rich defects generated during resistive switching and atomic rearrangements of the conductive filament. The findings are supported by electroluminescence and confocal photoluminescence measurements. The optical response of the atomic switch can be optimized for a high radiation efficiency by introducing an asymmetrical Ag-Pt antenna as well as by maximizing the LDOS enhancement with the help of Ag filament formation. The engineered photon source discussed here features an atomic-sized footprint and a straightforward and scalable fabrication process. As the emitted photons are associated with a resistive state change, our findings can be exploited in optical memristive neural networks to identify weight changes from the corresponding memristor.

Materials and methods

The details of sample fabrication, experimental setup and numerical simulations are provided in Supplementary Information section I, II, and IV, respectively.

Acknowledgements

This work has been funded by the Werner Siemens Foundation, by the European Research Council under the European Community’s Seventh Framework Program FP7/2007–2013 Grant Agreement 306772, the European Union through the PO FEDER-FSE Bourgogne 2014/2020 programs, the Conseil Régional de Bourgogne Franche-Comté and has been supported by the EPHI Graduate School (contract ANR-17-EURE-0002). We thank the cleanroom operations team of the Binnig and Rohrer Nanotechnology Center (BRNC) for their help and support in sample fabrication. We thank Michael Doderer, Killian Keller, and Fabian Ducry for joining the luminescence theory discussion and Aymeric Leray for his help at analyzing the PL lifetimes. We also thank Anna Krystanik and Oliver Heitz for the XPS characterization. It was carried out at the technological platform ARCCN Carnot with the support of the Région de Bourgogne Franche-Comté, the Délégation Régionale à la Recherche et à la Technologie (DRRT), and the CNRS.

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Author contributions

B.C., A.B. and J.L. conceived the concept and supervised the project. B.C., T.Z. and K.M. carried out the measurements and processed the data. B.C., X.Z. and A.E. fabricated the sample. M.L., E.P. and A.E. assisted in the fabrication and experiment. X.Z., T.Z. and U.K. performed the simulation. B.C., T.Z., K.M., M.L., A.B. and J.L. wrote the manuscript. All authors discussed and commented on the manuscript.

Conflict of interest

The authors declare no competing interests.

Supplementary information

The online version contains supplementary material available at https://doi.org/10.1038/s41377-022-00766-z.
