Highly Stable Persistent Photoconductivity with Suspended Graphene Nanoribbons

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Graphene nanoribbon (GNR), also known as 1-dimensional graphene, with a non-zero band gap has a huge potential for various electrical and optoelectrical applications because of its high transparency, flexibility, controllable band gap, and unique edge states. Recent advances in the synthesis of GNR enable us to show the possibility of GNRs as future high performance electrical devices. However, the applicability of GNRs to optoelectrical devices is unclear. Here we report that suspended GNR devices can show persistent photoconductivity (PPC) with long decay time (over 72 h) and adequate environmental stability. Repeated non-volatile memory operation is also demonstrated with an integrated PPC device using GNRs. This very stable PPC device can be applied to a wide variety of fields such as ultra-low-power non-volatile memory, nanoscale imaging, and biological sensors. Our results have opened the door to advance the study of GNRs in novel directions such as optoelectrical applications.

Graphene nanoribbon (GNR), a strip of graphene with nanometer width, can possess a non-zero band gap, thereby changing the electrical property of graphene from metallic to semiconducting behavior. GNR is expected to be utilized in various electrical and optoelectrical applications because of its high optical transparency, mechanical flexibility, controllable band gap, and unique spin-polarized edge states. Progress has been made in the synthesis of GNR such as bottom-up chemical synthesis of edge-controlled GNR and epitaxial synthesis of armchair GNR on a Ge substrate. Although these advances enable us to demonstrate the possibilities of GNR in future high performance electrical devices, it is still a challenge to clarify the suitability of GNRs for optoelectrical applications because of the difficulty of GNR manipulation. Recently we have realized integrated synthesis of suspended GNRs with nanobar catalyst at the wafer scale, which can offer a novel platform for GNR study to measure various optoelectrical properties of GNRs.

The current levels associated with persistent photoconductivity (PPC) can be modulated by photo-irradiation, and the modulated current can be maintained even after the photo-irradiation has ceased; this is a well known phenomena for bulk 3-dimensional (3D) materials. PPC can be expected to be utilized for non-volatile memories, imaging sensors, and various chemical sensors. Decreasing the size of the PPC device is important to increase the capacity of non-volatile memory, improve imaging resolution down to nanoscale, and develop other novel applications. Recently, it has been reported that PPC can be made to appear even in 2D materials by modifying the contact resistance between graphene and Au electrodes or by employing a heterojunction of graphene and molybdenum disulfide (MoS2). The modification of Au electrodes can show stable PPC in air but the decay time is very short (~5 s). Although the graphene/MoS2 heterojunctions can show a longer decay time, their environmental stability is relatively low because of the high chemical reactivity of MoS2, with various molecules such as oxygen, hydrogen, and water. Fabrication of low-dimensional (low-D) PPC devices with high environmental stability is a crucial subject from the industrial point of view.

Here we report the successful fabrication of a highly stable PPC device with a suspended GNR array, which is grown by nanobar-catalyzed plasma CVD. The functionalized GNR device shows pronounced PPC with a long decay time (over 72 h) and excellent room temperature stability in air. The PPC operation can be also realized even in water, indicating that GNR-based PPC devices can possess excellent environmental stability. By using these highly stable GNR-PPC devices, non-volatile memory operation was also demonstrated. The 1-bit device size can be decreased to ~0.25 μm², and over 3960 GNR devices were integrated. The origin of the PPC was also investigated.
systematically investigated. The very stable GNR-based PPC device has a huge potential for the realization of ultra-low-power non-volatile memory and various biological applications such as DNA sequencers in micro fluid devices\textsuperscript{19}, \textit{in vivo} imaging\textsuperscript{20}, and microchip implants\textsuperscript{21}. Our results demonstrate a novel direction of GNR study for optoelectrical applications.

The GNR devices used in this studies were fabricated by plasma CVD with a Ni nanobar catalyst (Fig. 1a–d)\textsuperscript{11,12}. The GNRs are about 10–30 nm in width, which was controlled by the initial Ni nanobar width. An image of a typical device is shown in Fig. 1a, where the field-effect transistor (FET) configuration of the suspended GNR device is formed with the source and drain electrodes made of Ni. The gate bias voltage (\(V_G\)) was applied to the highly doped silicon substrate that had a 300 nm thick SiO\(_2\) layer as the gate insulator. Photo-irradiation was applied to this FET device by a solar simulator with light power (\(P\)) of 200 mW/cm\(^2\) unless otherwise specified. Typical spectrum of our light source (solar simulator) is shown in Fig. S1. A typical drain-source current (\(I_{DS}\)) curve of this GNR-array device shows highly conductive ambipolar properties (Fig. 1b). Although the accurate layer number of GNR used in this study is not sure, it can be around 5 to 10 layers by judging from our previous studies.

### Results and Discussion

First, the photoresponse property of \(I_{DS}\) was measured for the GNR device immediately after the synthesis of the GNR; this will be referred to hereafter as a “fresh GNR device”. The deviation of \(I_{DS}\) (\(\Delta I_{DS}\)) between \(I_{DS}\) with light irradiation and without light irradiation (\(I_{dark}\)) is defined as \(\Delta I_{DS} = I_{DS} - I_{dark}\). The current levels of \(I_{DS}\) just before and after stopping photo irradiation in the steady state are defined as \(I_{on}\) and \(I_{off}\), respectively. In this study, we use two of important photo induced currents named persistent photoconductivity and temporal photoconductivity, which are defined as \(I_{per} = I_{dark} - I_{off}\) and \(I_{temp} = I_{off} - I_{on}\), respectively (Fig. 1e). The fresh device shows the usual photoresponse, where current suddenly decreases upon irradiating the device with light, and then the current level goes back to the original value after stopping the light irradiation (black curve in Fig. 1f). This means the \(I_{per}\) is close to 0. Next, a similar measurement was carried out for an “old” GNR device that had been stored in air for several months. The old device also shows a similar photoresponse which is temporal depression of \(I_{DS}\) during the light irradiation (Fig. 1e and purple curve in f). This is characteristic of PPC behavior\textsuperscript{13–16}. Similar phenomena have been reported for other 2D materials such as suspended graphene with oxidized Au electrodes\textsuperscript{17} and heterojunctions of graphene and MoS\(_2\)\textsuperscript{18}.

To elucidate the difference between fresh and old GNR devices, we introduced an oxidation process to the fresh device. Oxidation was carried out with mild oxygen (O\(_2\)) plasma treatment. It was found that \(I_{per}\) gradually increases with an increase in the plasma irradiation time and obvious PPC can be observed that is similar to that of the old GNR device (Fig. 2a–d). Similar results can be also observed after simple annealing in air at 350 °C (Fig. S2). This indicates that the PPC that appeared in the old GNR device should be due to the oxidation of Ni or related reactions; this is similar to the previously reported graphene and oxidized-Au results\textsuperscript{17}. It is noteworthy that the \(I_{per}\) for the GNR with oxidized-Ni electrode can maintain its current level more than 3 days (72 h), which is about 25,000 times longer than that of graphene with oxidized-Au electrodes (Fig. 2e).
Using this very stable PPC device, similar measurements were also carried out in water. The GNR device was covered with a droplet of water, and $I_{DS}$ was measured with and without photo-irradiation (Fig. 3a and b). Surprisingly, obvious $I_{per}$ can be also observed even in water with excellent stability (Fig. 3c). This indicates that...
the GNR-based PPC device has very high environmental stability and can be used for various biological applications such as DNA sequencers in micro fluid devices, in vivo imaging, and microchip implants.

Since very stable PPC can be observed with our GNR device, we attempted to demonstrate memory operation. For the practical application of any memory device, it is necessary to realize three main operations: writing, reading, and erasing. The creation of stable \( I_{\text{per}} \) as demonstrated above shows that the writing operation is possible by irradiating the device with light. The writing information can be read by the deviation of current through the GNRs. Since the origin of the \( I_{\text{per}} \) can be attributed to the trapped charge created by light irradiation, we attempted to apply a pulsed gate voltage to the GNR device to release the trapped charge (the detailed mechanism of \( I_{\text{per}} \) will be discussed later). Here, the erasing rate is defined as \( r_e = 100 \times \frac{I_{\text{dark}} - I_{\text{per}}}{I_{\text{per}}} \), and \( I_{\text{eras}} = F_{\text{dark}} - I_{\text{dark}} \). The pulse width for erasing \( t_{\text{eras}} \) was set to 3 sec. When \( V_G \) is switched from \( V_G = -60 \text{ V} \) to positive \( V_G \), \( \Delta I_{\text{DS}} \) temporally increases then decreases and stabilizes after the \( V_G \) is back to original value (\(-60 \text{ V}\)). The \( \Delta I_{\text{DS}} \) of the stable state clearly decreases with an increase in \( V_G \) and reaches zero when \( V_G = 50 \text{ V} \), where \( r_e \) reaches around 90% (Fig. 4b and c). The detailed erasing mechanism will also be discussed later. This indicates that the erasing operation can be accomplished by applying a relatively high pulsed gate bias voltage. Repeated operations of writing, reading, and erasing were also demonstrated (Fig. 4d). These periodic changes of \( I_{\text{per}} \) can be produced repeatedly, showing that our GNR device can operate as a non-volatile memory with high stability under normal atmosphere and room temperature. For a practical optical memory device, integration of the memory device is very important to improve the storage ability. With this in mind, an integrated GNR
memory has been fabricated as shown in Fig. 4e–h, where 9 memory cells were fabricated, and each memory cell contains 440 GNR arrays. We measured the differential resistance \( \Delta R = R_{\text{after}} - R_{\text{before}} \); here \( R_{\text{before}} \) and \( R_{\text{after}} \) are the resistances of the GNR arrays before and after photo-irradiation for all 9 cells. A pronounced change of \( \Delta R \) was observed in all the cells after a mild O₂ plasma treatment (Fig. 4f). This shows that all of the memory cells can function as a non-volatile memory. In one memory cell, 440 GNRs are integrated with high integration density (0.25 \( \mu m^2 \)) (Fig. 4e). It is also confirmed that the PPC can appear not only for the GNR array, but also for a single GNR device (Figs 4g,h and S3). This indicates that each GNR device can work as a single-bit memory. The memory size for 1 bit of storage can be given by a simple calculation, which shows that the 1-bit memory size can be decreased down to 0.25 \( \mu m^2 \) with our single GNR device (Fig. 4g and h). This scale advantage implies that 0.4 Gbits of memory can be integrated within 1 cm². The GNR memory also has the advantage of low energy operation. Figure 4i summarizes the writing energy vs. writing time for various memories. The NAND memory that is widely used in practical devices needs a writing energy of \( 10^{-9} \) J and its speed is about \( 10^{-3} \) sec\(^2\). Recent advances in memory technology can lower the writing energy and increase the speed using novel operational principles such as PCRAM, ReRAM, STT-MRAM, FeRAM, and SRAM\(^2\). When we plot the specifications of our GNR memory in Fig. 4i, the potential abilities of GNR memory can be elucidated. Because of the limitations of the measurement system, the writing speed of our GNR memory was set as 0.1 sec (Fig. S4). If we had used a writing speed of 10 ns, which is similar to that of FeRAM, the writing energy of GNR memory could be decreased down to \( 5 \times 10^{-21} \) J, which is about \( 10^{-7} \) times lower than that of FeRAM\(^2\). It has been reported that the photoresponse time can be decreased to 5.5 ps with graphene/TMD devices\(^2\), indicating that the assumption of 10 ns operating time for the GNR memory should be reasonable.

To further improve device performance, it is necessary to fully understand the operating mechanism. For this reason, detailed PPC measurements for the array of GNRs were carried out with under different values of \( V_G = -60 \) V or 0 V (Fig. 5a and b). As already discussed above, \( I_{\text{temp}} \) can produce a negative value of \( \Delta I_{\text{DS}} \) during photo-irradiation. This is observed with high reproducibility even under high vacuum conditions, indicating that the adsorption or desorption of impurities on the surface of the GNR caused by photo-irradiation can be ignored in our GNR devices. The origin of this \( I_{\text{temp}} \) can be explained by considering the conductance change during photo-irradiation. It has been confirmed that the conductance of our GNR device decreases with increasing temperature, denoting a negative bolometric coefficient around 300 K (Fig. S5)\(^2\). Thus, the temperature...
increase of GNR caused by photo-irradiation should be the critical origin of $I_{\text{temp}}$. For the $I_{\text{temp}}$ component, an obvious change of the polarity can be observed for different values of $V_G$. The negative and positive $I_{\text{temp}}$ can be observed with $V_G = -60 \text{ V}$ (Fig. S5a) and $V_G = 0 \text{ V}$ (Fig. S5b), respectively. The dependencies of $I_{\text{temp}}$ and $I_{\text{ppc}}$ on $V_G$ are plotted in Fig. S5c. We can see that $I_{\text{temp}}$ strongly depends on $V_G$, and can be tuned between negative and positive levels by changing $V_G$ from $-60 \text{ V}$ to $+60 \text{ V}$, whereas $I_{\text{ppc}}$ hardly depends on $V_G$. The weak $V_G$ dependence of $I_{\text{temp}}$ is consistent with the explanation based on the bolometric effect. To elucidate the origin of the $V_G$ dependence of $I_{\text{temp}}$, the $I_{\text{DS}} - V_G$ curve was measured with and without light irradiation. A clear shift of the charge neutral point ($V_{\text{cnp}}$) in the negative $V_G$ direction can be observed during photo-irradiation. The value of the shift increases with the light power ($P_{\text{in}}$) (Fig. S5d). The transconductance of electrons ($g_{\text{m\,p}}$) decreases (blue triangle in Fig. S5e) while that of holes ($g_{\text{m\,n}}$) hardly changes with increasing $P_{\text{in}}$ (red square in Fig. S5e). These results imply that GNRs are n-doped by photo-irradiation, and the doped carriers may act as scattering centers for electrons, resulting in the decrease of $g_{\text{m\,n}}$. Therefore, the negative and positive persistent current can be observed with $V_G = -60 \text{ V}$ and 0 V, respectively (Fig. S5f). Carrier doping effects caused by photo-irradiation have been reported in graphene devices, and the mechanism is the photo-gating effect that originates in charging of the SiO$_2$ surface during photo-irradiation$^{24,25}$; consequently, it can be assumed that hot carriers excited by light irradiation can be deeply trapped at specific sites, causing a very stable $I_{\text{ppc}}$ in our GNR devices. Note that the minimum power of solar simulator for the appearance of PPC was $\sim 25 \text{ mW/cm}^2$, which is several orders higher than that of other previous results$^{52,53}$. This is because of the effect of wavelength dependence. Only ultraviolet (UV) light region is effectively used for PPC generation (discussed later).

As discussed above, if $I_{\text{ppc}}$ originates from the trapped charge, stable trap sites should be formed in our GNR device by mild O$_2$ plasma treatment. To identify the effects of mild O$_2$ plasma treatment on the formation of trapping site, following systematic experiments were carried out.

**Effects of mild O$_2$ plasma treatment on the GNR structure.** First, we try to identify the effect of mild O$_2$ plasma treatment to GNR structures. Raman scattering spectroscopy was used to identify the introduction of defects to GNR. Note that because it is very difficult to consider the edge effect of GNR, we used mechanically-exfoliated graphene instead of GNR for this Raman experiment. For the pristine graphene transferred on the Ni (50 nm)/SiO$_2$/Si substrate, repeatable mild O$_2$ plasma treatments were carried out. It is found that D-band peak relating with the disorder of graphitic structure can be observed even after 1 min plasma treatment (Fig. S6). The ratio of D-band to G-band ($I_D/I_G$) gradually increased with plasma treatment time. The introduction of defects was more significant for the thinner graphene. This indicates that the defects should be introduced to GNR by our mild O$_2$ plasma treatment. It should be noted that even after long time plasma treatment (5 min), over all film structure did not change (Fig. S6), denoting not the etching reaction but introduction of local defects should be dominant in our mild O$_2$ plasma treatment. This is because of the low ion energy in mild plasma reaction, which had been already developed by our previous study$^{50}$.

**Effects of mild O$_2$ plasma treatment on the Ni electrode.** To identify the effects of plasma treatment on the Ni electrode, detailed analysis was carried out. The candidates for the functionalized structures of Ni are NiO, Ni$_2$O$_3$, or Ni(OH)$_2$, where the band gap has been reported to be about 4.3 eV$^{27}$, 3.38 eV$^{28}$, and 3.6–3.9 eV$^{29}$ respectively. The wavelength ($\lambda$) dependence of the photoresponsivity $R_p = \Delta I_{\text{ppc}}/P_{\text{in}}$ was measured for the old GNR device, where $I_{\text{ppc}}$, $P_{\text{in}}$, and $R_p$ denote current change before and after light irradiation, incident light power, and the area of the GNR, respectively. The $\lambda$-selective irradiation was carried out by splitting the light source from a Xe lamp with a conventional spectrometer. A clear $\lambda$ dependence of $R_p$ was observed, and only irradiation at UV wavelength would be capable of producing such a high value of $R_p$ (Fig. S7). With the light of 360 nm wavelength, PPC can be observed even with the low power of 41 $\mu$W/cm$^2$. The optical adsorption spectra were measured for Ni thin films deposited on a quartz substrate without graphene layer before and after the mild O$_2$ plasma treatment. Mild O$_2$ plasma treatment is carried out for 10 min with homemade plasma CVD system (See method). The adsorption spectra show obvious adsorption in UV region, but this appears only for mild O$_2$-plasma-treated Ni films (Fig. S8). This is consistent with the $\lambda$ dependence of $R_p$, suggesting that the dominant trapping sites are not oxidized GNR (band gap: 0.02~2 eV$^{30}$), but rather, oxidized or hydroxidized Ni.

To further identify the detailed structure of functionalized Ni serving as the trapping site, X-ray photoelectron spectroscopy (XPS) measurements were carried out. After mild O$_2$ plasma treatment (5 min), the peak of pure Ni vanished and only the chemical-shift peak was observed at 855.69 eV which is obviously different from that of NiO (854.0 ± 0.5 eV) and very close to that of Ni$_2$O$_3$ (856.6 ± 0.8) and Ni(OH)$_2$ (855.7 ± 0.4) (Fig. S9a)$^{31}$. The shapes of the peaks over a wide range of spectrum were also taken into consideration to judge the origin of the peak at 855.69 eV. The observed spectrum after mild O$_2$ plasma treatment is well matched with that of Ni(OH)$_2$ (Fig. S9b)$^{32}$. Atomic force microscopy (AFM) measurements were also performed to obtain structural information about the functionalized Ni after mild O$_2$ plasma treatment (5 min). Before mild O$_2$ plasma treatment, the surface of the Ni is relatively smooth, and its average of roughness height ($R_h$) is 0.055 nm (Fig. S10a and b). On the other hand, after mild O$_2$ plasma treatment, surface becomes rough ($R_h = 0.59$ nm) and a honeycomb-like nanos-structure is formed (Fig. S10c and d) with ~1 nm height and ~10 nm width (Fig. S10d). A similar layered structure has been reported for Ni(OH)$_2$ ($\beta$-Ni(OH)$_2$)$^{33}$. Judging from this systematic analysis, it can be concluded that $\beta$-Ni(OH)$_2$ nanostructures are probably formed on the surface of Ni by mild O$_2$ plasma treatment.

**Interlayer structure between GNR and Ni after mild O$_2$ plasma treatment.** Because it is revealed that mild O$_2$ plasma treatment modify GNR and Ni surface to disordered GNR and $\beta$-Ni(OH)$_2$, respectively, we have to think about the effects of those modified structures on the appearance of PPC. If the disorder site of GNR itself is the dominant trapping site of carrier, PPC should be observed independent from electrode materials. Then, we fabricated similar suspended-graphene device with various electrode materials such as Ni, Au, and Cu.
by transferring mechanically-exfoliated few-layer graphene to electrodes. Interestingly, PPC can be observed only for Ni electrodes after mild O₂ plasma treatment (Fig. S11). This indicates that kinds of electrode materials possess significant contribution to determining the appearance of PPC, i.e. forming defects to channel region of GNR is not enough to cause PPC.

Then, we attempt to confirm the effects of β-Ni(OH)₂. Similar suspended-graphene device was fabricated (Fig. S12). Before the transfer of graphene, Ni electrode was pre-oxidized by mild O₂ plasma treatment. Interestingly, current can not be through in this pre-oxidized Ni/GNR device, which should because of the poor contact between pre-oxidized Ni and GNR. By considering this point, it can be conjectured that surface of Ni underneath of GNR is not completely oxidized but partial oxidation can happen for the device where PPC can be observed (Fig. S13). Such local structures of disordered GNR/β-Ni(OH)₂ can dominantly work for the appearance of PPC. Very stable trapping site may be formed at such modified structures. As a control experiment, we also carried out similar experiments with not suspended but supported graphene device with Ni electrode. Noticeably, however, neither PPC nor temporal response can not be observed (Fig. S14). Because of the bottom contact structure of graphene and Ni electrode, it should be difficult to form local structures (disordered graphene and Ni(OH)₂) at the contact region between graphene and Ni for supported graphene device, which can be a possible explanation for the lack of PPC features in the supported graphene device. This indicates that the top contact of Ni and GNR should be important to cause PPC and suspended GNR devices can effectively provide such ideal contact structures for PPC.

Possible model for appearance of PPC in GNR device. Based on these results, we can propose the following mechanisms of PPC in our GNR device (Fig. 6). As discussed above, local defects are introduced to GNR by mild O₂ plasma treatment. The local defect can enhance the formation of nanoscale layered (β-Ni(OH)₂) structures around the defects site. Then, locally modified structure of disordered GNR/β-Ni(OH)₂ should be formed at the interface between the GNRs and the Ni electrodes (Fig. 6a). When photo-irradiation is directed at this system under an applied \( V_G < V_{\text{CNP}} \) (Fig. 6b), electrons can be optically excited in disordered GNR/β-Ni(OH)₂ and relaxed by going through Ni or GNR resulting in the creation of a localized hole. If this localized hole is trapped at a stable site at the interface of disordered GNR and Ni electrode, it should be difficult to form local structures (disordered graphene and Ni(OH)₂) at the contact region between graphene and Ni for supported graphene device, which can be a possible explanation for the lack of PPC features in the supported graphene device. This indicates that the top contact of Ni and GNR should be important to cause PPC and suspended GNR devices can effectively provide such ideal contact structures for PPC.

![Figure 6.](image-url)
We mainly discussed the stable and long-decay time of PPC as a performance of our GNR device. As a one of the important performance of PPC, the conductivity-switching effect is also reported. Since the GNRs used in our study is relatively thicker and wider geometry, only weak conductivity switching was observed. By using further thinner and narrower GNRs including high on/off current ratio, large conductivity-switching effects can be also expected with GNR device, which can be significant difference compared with graphene-based PPC device.

In summary, we have fabricated very stable PPC devices with suspended GNRs. The stable PPC can be obtained not only in air but also in a solution phase. Optically-driven non-volatile memory operation has been also demonstrated with these stable PPC devices with suspended GNRs. The GNR memory device can be densely integrated, and minimum memory size for 1 bit of storage can be decreased to 0.25 μm². The energy for writing operation can be as low as 0.1 pJ/bit, indicating that the GNR-based non-volatile photo memory has a huge potential for practical use in future low power electronics. The detailed operation mechanism was also investigated, showing that the heterojunction between locally-disordered GNR and nanoscale β-Ni(OH)₂, structures can behave as a stable trapping site for optically generated holes, causing the excellent stability of the PPC.

**Methods**

**Plasma CVD.** A homemade plasma CVD system was used for the rapid-heating plasma CVD (RH-PCVD). Before the plasma CVD growth, an electric furnace was heated to the desired temperature (typically 800–900 °C) under flowing hydrogen (50 Pa). A substrate was immediately transferred to the center area, and rapid heating was performed. CH₄ and H₂ gases at a 9:1 ratio (250 Pa) were inletted immediately after a pre-set temperature was reached. Next, radio frequency power (100 W, 13.56 MHz) was supplied to the coils outside of the quartz tube.

The plasma was typically maintained for 5–30 s. Following the plasma CVD, the substrate was moved from the center to the outside of the electrical furnace so that its temperature would rapidly decrease.

**Mild O₂ plasma treatment.** A homemade plasma treatment system was used. An mild O₂ plasma can be generated by supplying radio frequency power (26 W, 13.56 MHz) to the coils outside of the quartz tube under flowing O₂ gas (150 sccm). A grid mesh was set 15 cm ahead of the center of the coil. The sample was 40 cm away from the center of the coil.

**Characterizations.** The structure of the GNR array sample was characterized by scanning electron microscopy (SEM; Elionix, ELS-7500EXTK and Hitachi, SU1510, Japan). The electrical measurements of the GNR devices were performed using a vacuum probe station with a semiconductor parameter analyzer (HP 4155C). The elemental analysis of the Ni film was characterized by X-ray Photoelectron Spectroscopy (XPS; Ulvac-phi, ESCA1600, Japan). The structure of the Ni surface was analyzed by atomic force microscopy (AFM; JEOL, JSFM-5400, Japan).

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Author Contributions
T. Kato conceived the experiments and supervised the project. H.S. fabricated the devices and performed the measurements. H.S., N.O. and T. Kato analyzed the data and H.S., T. Kaneko and T. Kato, prepared the manuscript. All authors discussed the results and commented on the manuscript.

Additional Information
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