High-Performance Stable Field Emission with Ultralow Turn on Voltage from rGO Conformal Coated TiO$_2$ Nanotubes 3D Arrays

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A facile method to produce conformal coated reduced graphene oxide (rGO) on vertically aligned titanium oxide (TiO$_2$) nanotubes three dimensional (3D) arrays (NTAs) is demonstrated for enhanced field emission display applications. These engineered nano arrays exhibit efficient electron field emission properties such as high field emission current density (80 mA/cm$^2$), low turn-on field (1.0 V/μm) and field enhancement factor (6000) with high emission current stability. Moreover, these enhancements observed in nano arrays attribute to the contribution of low work function with non-rectifying barriers, which allow an easy injection of electrons from the conduction band of TiO$_2$ into the Fermi level of reduced graphene oxide under external electric field. The obtained results are extremely advantageous for its potential application in field emission devices.

Recently, high quality field emitters have gained importance owing to their potential for reliable integration into optoelectronic devices. Electron sources are essential elements in a variety of applications that include microwave amplifiers, parallel electron beam microscopes, x-ray sources and flat panel display technology$^1$–$^3$. One dimensional (1D) vertically-organized nanostructures (e.g. nanowires$^4$–$^6$, nanotubes$^7$–$^9$, nanobelts$^{10,11}$ and nanoneedles$^{12}$) are considered to be excellent field emission (FE) based electron emitters for delivering high current density at a low applied potential due to their high aspect ratios, high-field enhancement factor and low work functions. It is well established that the nanostructures having sharp tips can reduce the strength of turn-on electric fields by several orders of magnitude and decreases the barrier width due to the enhance local electric field at these tips. As a result, these nanostructure materials exhibited the excellent electron emission characteristics$^{13}$. Among these 1D nanostructure materials, diamond based and carbon nanotubes (CNTs) have exhibited good FE performances owing to relatively low emission threshold field and high thermal and electrical conductivity$^8,14,15$. However, further development of FE emitters depends critically on the challenging task of growing CNTs with specific properties as well as in current optoelectronic devices. In addition, higher work function, lack of adequate long-term or high-temperature FE stabilities and unsatisfactory mechanical properties have hindered the development of these materials for practical applications.

Wide band-gap semiconductors including TiO$_2$, MoO$_3$, SiC$^{17}$, ZnO$^{12}$, WO$_3$ and similar materials have also attracted much interest for their favorable FE properties because of their low electron affinity as well as better chemical stability. The band-bending effect of wide band gap semiconductors allows the field emission by lowering the surface barrier and bringing more electrons to the bottom of the conduction band. Among these materials, titanium oxide (TiO$_2$), as a wide-band-gap (∼ 3.1 – 3.2 eV) semiconductor, has been extensively studied because of its long-term thermodynamic stability, low cost,

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The morphology, dimensions and structural parameters of TiO$_2$ NTAs are easily controlled by anodic oxidation process such as anodic voltage, oxidation time and electrolyte composition. An anodic oxidation technique with conformal coating of rGO on highly oriented annealed TiO$_2$ 3D NTAs samples (40 V, 2 hours) was used to form hybrid materials for enhanced FE property.

The graphene hybrid nanostructures have been extensively explored since the past decade for its highly-efficient field emission performance. But in particular, the conformal coating of reduced graphene oxide (rGO) on vertically aligned TiO$_2$ NTAs hybrid structure for FE studies has not been demonstrated till date. The combination of vertically aligned TiO$_2$ NTAs with rGO is expected to extend the development of various flexible devices. This also extends the application scopes and reinforces the properties of graphene and TiO$_2$ materials. The conformal coating of rGO on TiO$_2$ NTAs greatly facilitates large surface area and charge carrier dynamics and improves the FE performance compared to other nanostructures such as commercial TiO$_2$ nanoparticles (NPs), as-synthesized TiO$_2$ NTAs and annealed TiO$_2$ NTAs samples, which is not reported so far. The schematic presentation of synthesis process for highly oriented architecture of TiO$_2$ NTAs via anodization technique with conformal coating of rGO on highly oriented annealed TiO$_2$ 3D NTAs samples (40 V, 4 hours, 500°C) is shown in Fig. 1.

**Results**

The crystallinity and phase of samples were analyzed by X-ray diffraction technique. Fig. 2(a–c), shows the XRD pattern of as-synthesized TiO$_2$ NTAs (Fig. 2a), annealed TiO$_2$ NTAs (Fig. 2b) and conformal coated rGO on annealed TiO$_2$ NTAs hybrid structures (Fig. 2c). The quantitative analysis of Fig. 2(a,b) shows that all diffraction peaks correspond to TiO$_2$ and Ti substrate and represents the tetragonal crystal structure with space group S.G. 14 1/amd (141). The typical diffraction peak (101) centred at 25.1° indicates the TiO$_2$ anatase phase (JCPDS No. 21–1272), which is formed after annealing at 500°C for 2 hours. The peaks observed at (101), (103), (004), (112), (200), (105), (211), (213), (116) and (220) correspond to TiO$_2$ anatase phase. It can be noticed that overall crystal structure becomes greatly refined after annealing. The lattice parameters were calculated from the observed d-values through a least square fitting method using computer program 'unit cell refinement software'. These lattice parameter values are $a = b = (3.7852 \pm 0.0029) \text{Å}$, $c = (9.5139 \pm 0.0067) \text{Å}$ for as-synthesized TiO$_2$ NTAs and $a = b = (3.79632 \pm 0.0091) \text{Å}$, $c = (9.5832 \pm 0.0097) \text{Å}$ for annealed TiO$_2$ NTAs samples. The average domain non-toxicity, strong oxidizing power and its optical as well as electrical properties. Hence, the vertically aligned TiO$_2$ nanotube 3D arrays (NTAs) have created significant interest for good FE properties because of their sharp tips, low work function (4.4 eV), high aspect ratios, vertical orientation, tunable mesopore size, large internal surface area, convenient recycling and direct path for electron transport are considered having important FE properties. However, the studies carried out of FE properties of TiO$_2$ nanostuctures have still been rather inadequate due to the limited success in reliable synthesis of conductive arrays. For better performance and economical cost of field emitter materials, it is important to look for carbon based nanostructure materials having high surface area. In this exploration, especially, the 2D (two dimensional) nanomaterials (e.g. graphene) can be a better alternative, which can be easily integrated with TiO$_2$ on nanoscale to form hybrid materials for enhanced FE property.

Graphene with unique 2D $\pi-\pi$ conjugated structure and a super strong form of carbon, has been regarded as a component of devices in recent years owing to its high electrical and thermal conductivity (~5,000 Wm$^{-1}$K$^{-1}$), good mobility of charge carriers (~200,000 cm$^2$V$^{-1}$s$^{-1}$), superior chemical stability, high specific surface area (~2,630 m$^2$ g$^{-1}$) and sharp edges as well as its potential applications in field emission, solar cells, gas sensors, transparent conducting electrodes and photocatalysts. Graphene oxide (GO) is an excellent system with oxygen containing functional groups attached to the basal plane and edges, which makes it insulating and hydrophilic. These functional groups reduce the interaction energy between the graphene layers and thus make it dispersible in aqueous media. Graphene exhibits higher emission currents with lower external electric field and provides the large injection carriers as well as electron-transport material than other carbon based materials.

The graphene hybrid nanostructures have still been rather inadequate due to the limited success in reliable synthesis of conductive arrays. For better performance and economical cost of field emitter materials, it is important to look for carbon based nanostructure materials having high surface area. In this exploration, especially, the 2D (two dimensional) nanomaterials (e.g. graphene) can be a better alternative, which can be easily integrated with TiO$_2$ on nanoscale to form hybrid materials for enhanced FE property.

Herein, we report an approach to develop rGO-TiO$_2$ NTAs hybrid nanostructures as efficient field emitters. In the investigations, vertically aligned TiO$_2$ NTAs were grown on titanium (Ti) substrates via anodic oxidation method and then conformal coating of rGO was transfer onto TiO$_2$ NTAs. It shows much improved FE properties than those obtained from pure TiO$_2$ NTAs and rGO nanostructures. The morphology, dimensions and structural parameters of TiO$_2$ NTAs are easily controlled by anodic oxidation parameters such as anodic voltage, oxidation time and electrolyte composition. An anodic oxidation process has been used extensively for the rapid production of aligned TiO$_2$ nanotubes because of their sharp tips, low work function (4.4 eV), high aspect ratios, vertical orientation, tunable mesopore size, large internal surface area, convenient recycling and direct path for electron transport.

The field emission properties of TiO$_2$ NTAs were investigated before and after being modified with rGO conformal coating and it was found that rGO conformal coated TiO$_2$ NTAs have low turn-on field, high current density and uniform emission with better stability over a large area as compared to as-synthesized TiO$_2$ NTAs. It is being demonstrated here that the incorporation of rGO through conformal coating on TiO$_2$ NTAs greatly facilitates large surface area and charge carrier dynamics and improves the FE performance compared to other nanostructures such as commercial TiO$_2$ nanoparticles (NPs), as-synthesized TiO$_2$ NTAs and annealed TiO$_2$ NTAs samples, which is not reported so far. The schematic presentation of synthesis process for highly oriented architecture of TiO$_2$ NTAs via anodization technique with conformal coating of rGO on highly oriented annealed TiO$_2$ 3D NTAs samples (40 V, 4 hours, 500°C) is shown in Fig. 1.
size for as-synthesized and annealed TiO$_2$ NTAs samples are 105 nm and 121 nm with respect to (101) plane, which are estimated by using Scherer’s formula. The TiO$_2$ NTA domains increase in size from 105 to 121 nm after the annealing process. The XRD pattern of as-synthesized rGO nanosheets is shown in Figure S1 (see Supplementary Information), where (002) and (101) planes confirm the graphitic nature with hexagonal phase (JCPDS No. 75–1621). There is no obvious difference in the TiO$_2$ NTAs phase in annealed TiO$_2$ NTAs (Fig. 2b) and conformal coated rGO on annealed TiO$_2$ NTAs hybrid structure (Fig. 2c), indicating that the crystalline structure of TiO$_2$ NTAs was not influence by the deposition of rGO nanosheets on TiO$_2$ NTAs. However, no diffraction peak corresponding to rGO is observed in the hybrid structure, which may be because of the ultrathin nature of the rGO nanosheets layer coated on TiO$_2$ NTAs. Raman spectroscopic measurement was used to characterize the reduction of graphene oxide (GO) as this process is very sensitive to crystallinity and microstructure of the materials. It was employed to confirm the formation of TiO$_2$ anatase phase and the existence of rGO nanosheets on the surface of conformal coated rGO on annealed TiO$_2$ NTAs hybrid structure. Figure 2d shows the Raman spectra of annealed TiO$_2$ NTAs (Fig. 2d (i)) and conformal coated rGO on annealed TiO$_2$ NTAs hybrid structure (Fig. 2d (ii)). Fig. 2d (i) shows the Raman peaks at 395, 516 and 640 cm$^{-1}$ corresponding to the B$_{1g}$, B$_{1g}$ and A$_{1g}$ modes of anatase TiO$_2$, respectively, which are consistent with the results of XRD. The Raman spectrum of as-synthesized TiO$_2$ NTAs is shown in Figure S2 (a) (see Supplementary Information) in which all the three peaks (at 395, 516 and 640 cm$^{-1}$) are observed. It confirms the formation of anatase phase. It can be easily observed that the peaks intensity is enhanced after annealing treatment (Fig. 2d (i)). The Raman spectrum of conformal coated rGO on annealed TiO$_2$ NTAs hybrid structure is shown in Fig. 2d (ii). In addition to different TiO$_2$ modes, the other new vibration mode D-band at 1354 cm$^{-1}$ and G band at 1600 cm$^{-1}$ are also observed. These peaks correspond to the D and G band of rGO, which are attributed to the breathing mode of the k point photons of A$_{1g}$ symmetry and first order scattering of E$_{2g}$ phonon of the sp$^2$ C atoms, respectively$^{42}$. This confirmed the presence of graphene oxide in this conformal coated rGO on TiO$_2$ NTAs hybrid structure. All other peaks (Fig. 2d (ii)) are due to the TiO$_2$ NTAs. The intensity ratio of D to G band ($I_D/I_G$) has been proposed

Figure 1. The synthesis process for highly oriented architecture of TiO$_2$ NTAs via anodization technique with conformal coating of rGO on highly oriented annealed TiO$_2$ 3D NTAs samples (40 V, 4 hours, 500°C).
to be an indication of disorder in the graphene or rGO nanosheets and a low ratio indicates a greater disorder arising from structural defects. The conformal coated rGO on TiO$_2$ NTAs has an intensity ratio (I$_D$/I$_G$) near to 0.940, suggesting that increased defects are brought by the reduction of graphene oxide$^{43}$. The raman spectra of as-synthesized rGO nanosheets is shown in Figure S2 (b) (see Supplementary Information). Such characteristics demonstrate that the rGO nanosheets have direct evidence of conformal coating of rGO on the TiO$_2$ NTAs.

The scanning electron microscope (SEM) and transmission electron microscope (TEM) were utilized to characterize the morphologies of as-synthesized, annealed TiO$_2$ NTAs and conformal coated rGO on TiO$_2$ NTAs hybrid structure. The typical SEM micrographs of as-synthesized and annealed TiO$_2$ NTAs samples at anodization voltage 40 V for 4 hours are shown in Figure S3 (see Supplementary information). Figures S3 (a-b) exhibits the lateral and top view of as-synthesized TiO$_2$ NTAs. Figures S3 (c-d) represents the lateral and top view of annealed TiO$_2$ NTAs. It can be seen that both the samples consist of uniform open nanotubes of TiO$_2$ with an average outer diameter of $\sim$110 nm and maximum length of $\sim$2$\mu$m. The alignment is not disturbed after annealing; rather it helps to improve the crystallinity of sample, which is confirmed by XRD results. The typical SEM micrographs of as-synthesized TiO$_2$ NTAs at different anodization voltage of 30, 40 and 50 V for 4h anodization time are shown in Figure S4 (see Supplementary Information). The lateral view of as-synthesized TiO$_2$ nanotube arrays sample for different anodization time intervals 1.5 and 2.5 h and the top view of TiO$_2$ nanotube arrays sample at different anodization voltage of 30 and 50 V are shown in Figure S5 (see Supplementary Information). The bottom views of highly dense TiO$_2$ NTAs after annealing at 500$^\circ$C for 2 hours is shown in Figure S6 (see Supplementary Information).

Figure 3(a) represents the TEM image of as-synthesized TiO$_2$ NTAs and Fig. 3(b) exhibits the single TiO$_2$ nanotube with outer diameter of $\sim$110 nm, which is in good agreement with the SEM results. Figure 3(b) is the magnified version of Fig. 3(a), where microstructure of a single TiO$_2$ nanotube can be easily seen in details. Fig. 3(c,d) shows the TEM/HRTEM images of vertically aligned TiO$_2$ NTAs annealed at 500$^\circ$C for 2hours. Figure 3(c) shows the TEM image of an isolated TiO$_2$ nanotube with $\sim$112 nm in diameter. It can be observed that after annealing, there is a slight variation in the outer diameter of TiO$_2$ nanotube. The SAED pattern (inset of Fig. 3c) confirms the crystalline nature of TiO$_2$ NTAs. The SAED ring pattern corresponding to (101), (200) and (004) lattice planes reveals the presence of TiO$_2$ anatase phase. All the indexed planes are also observed in the XRD pattern of annealed TiO$_2$ NTAs.
Figure 3. TEM images of (a) as-synthesized TiO$_2$ NTAs, (b) magnified version of (a), (c) annealed TiO$_2$ NTAs at 500 °C for 2 hour and inset shows the SAED pattern of annealed TiO$_2$ NTAs and (d) HRTEM image of annealed TiO$_2$ NTAs, (e) SEM, (f) TEM, (g) HRTEM images of conformal coated rGO on annealed TiO$_2$ NTAs; where micrographs clearly evidence the conformal coating of rGO on annealed TiO$_2$ NTAs and (h) EDAX pattern of conformal coated rGO on annealed TiO$_2$ NTAs.
Figure 3(d) demonstrates the HRTEM image of TiO2 nanotube having high quality lattice fringes without any distortion, which clearly demonstrates clear lattice fringes of TiO2 nanotubes after annealing process. The estimated interplanar spacing of adjacent lattice fringes is about ~0.35 nm, which corresponds to the (101) plane of anatase TiO2. The TEM and HRTEM images of TiO2 NTAs annealed at 500°C for 2 hours, is shown in Figure S7 (see Supplementary Information). The TEM and SEM images of as-synthesized rGO nanosheets are shown in Figure S8 (see Supplementary Information), which reveals that the rGO nanosheets are composed of few layers of graphene.

In order to explore the surface morphology of rGO coated TiO2 NTAs, SEM study is carried out and results are shown in Fig. 3(e). A thin shadow of graphene around the TiO2 nanotube can be seen as marked by arrow. Additionally, the XPS studies on conformal coated rGO on annealed TiO2 NTAs were also conducted to find out the purity and chemical composition of TiO2 nanotubes. The XPS spectrum of conformal coated rGO on annealed TiO2 NTAs hybrid structure is shown in Figure S9 (see Supplementary Information) and inset clearly shows the core level spectrum of Ti. In the XPS spectrum, signals corresponding to titanium, oxygen and carbon are observed. No other signals are detected, which shows the high purity of as-synthesized conformal coated rGO on annealed TiO2 NTAs. The typical TEM image of conformal coated rGO on annealed TiO2 has been shown in Fig. 3(f). The TEM micrograph reveals clear microstructural information about the conformal coating of rGO on TiO2 nanotubes in the hybrid structure. The estimated number of rGO layers is simply calculated by the difference between the diameter of TiO2 nanotubes before and after coating of rGO on the TiO2 nanotubes arrays by using TEM image (from Fig. 3(c,f)). The obtained thickness of few layers of rGO conformally coated on TiO2 nanotubes is around 6 nm, which indicates that 16–17 layers of rGO are coated on TiO2 nanotubes. Further, the HRTEM image was taken to study the interface between the rGO and TiO2 and results are shown in Fig. 3(g). The HRTEM image has been taken from yellow marked region of Fig. 3(f). In Fig. 3(g), the yellow dashed line shows the interface between rGO and TiO2 lattices, from where we have estimated the lattice spacing. The graphene is well established for its binding capabilities with metal oxide particles as well as metals, such as TiO2 and Eu through covalent bonding or complexation without any aggregation44,45. In the present investigations, rGO nanosheets conformal coated on TiO2 NTAs appear to have strong interactions between them, which should lead to development of advanced hybrid materials to be used for various potential applications such as in field emission devices. Furthermore, the TiO2 nanotubes as well as elemental composition were evaluated by EDAX analysis. The spot EDAX measurement was performed with reduced beam spot size to enhance the signal to noise ratio. The EDAX spectrum was recorded on rGO conformal coated TiO2 NTAs area as shown in Fig. 3(f). The EDAX study reveals the presence of titanium, oxygen, copper and carbon element for rGO conformal coated TiO2 NTAs sample, as shown in Fig. 3(h). The small content of copper is from copper grid, which is used in TEM analysis. The atomic % ratio of titanium to oxygen is almost 1:2 as expected in the TiO2 molecule.

The electron field emission involves extraction of electrons from the NTAs by quantum tunneling through the surface potential barrier46. The field emission characteristics (field emission current density (J) as a function of applied electric field (E)) at a sample to cathode distance of 100 μm for conformal coated rGO on annealed TiO2 NTAs hybrid structure, annealed TiO2 NTAs, as-synthesized TiO2 NTAs, rGO-commercial TiO2 NPs, commercial TiO2 NPs, rGO-Ti sheet, rGO and Ti sheet samples are shown in Fig. 4(a). It is found that the emission current density exponentially increases with increase in the applied field for all the samples. An emission current density of 80 mA/cm2 at 230 V is obtained for conformal coated rGO on annealed TiO2 NTAs sample, which is the highest value compared to the other rGO-commercial TiO2 NPs, commercial TiO2 nanoparticles (NPs), as-synthesized TiO2 NTAs, annealed TiO2 NTAs samples, pure Ti sheet, rGO nanosheets and rGO on pure Ti sheet substrate samples. The obtained results suggest that rGO conformal coated on TiO2 NTAs hybrid structure is ultimate choice for better field emission characteristics. It may be due to the presence of large no. of delocalized π electrons on the surface of rGO which act as electron injection carriers33,34. Field emission current also depends on the aspect ratio of the TiO2 nanotubes, which is very high in the present case. The field emission characteristics of conformal coated rGO on annealed TiO2 NTAs, annealed TiO2 NTAs, as-synthesized TiO2 NTAs and commercial TiO2 NPs samples are shown in Figure S10a (see Supplementary Information). It can be noticed that the turn-on field (E<sub>to</sub>) for conformal coated rGO on annealed TiO2 NTAs, annealed TiO2 NTAs, as-synthesized TiO2 NTAs and commercial TiO2 NPs are 1.0, 1.4, 3.7 and 4.8 V/μm, respectively (from Fig. 4a and S10a). The turn-on field E<sub>to</sub> values follows the sequences E<sub>to</sub> (conformal coated rGO on annealed TiO2 NTAs) < E<sub>to</sub> (annealed TiO2 NTAs) < E<sub>to</sub> (as-synthesized TiO2 NTAs) < E<sub>to</sub> (commercial TiO2 NPs). Furthermore, the field emission characteristics of different as-synthesized samples of conformal coated rGO on annealed TiO2 NTAs (sample 1, sample 2, sample 3 and sample 4) are also examined to explore reproducibility and the results are shown in Figure S10b (see Supplementary Information). It can be noticed that all the samples show similar and consistent behaviour. In addition to the above, the field emission behaviour of rGO, rGO-Ti sheet, rGO-commercial TiO2 nanoparticles and conformal coated rGO on annealed TiO2 NTAs samples from 1<sup>st</sup> to 4<sup>th</sup> cycle run are shown in the Figure S11a-d (see Supplementary Information). All tested samples show better emission uniformity and a good reproducibility of field emission behaviour during the initial 4 cycle run. The conformal coated rGO on annealed TiO2 NTAs clearly demonstrate the higher current density at low turn-on field (80 mA/cm<sup>2</sup>, 1.0 V/μm) in compared to all other samples (annealed TiO2 NTAs, as-synthesized TiO2 NTAs, rGO-commercial TiO2 NPs, commercial TiO2 NPs, rGO-Ti sheet, rGO and Ti sheet; Fig. 4a).
The stability of emission current is also evaluated at 230 V for conformal coated rGO on annealed TiO$_2$ NTAs, as shown in Fig. 4b and it is found to be very stable and no significant change is observed over a time period of 120 min at a current density 80 mA/cm$^2$. Thus, the conformal coated rGO on annealed TiO$_2$ NTAs, as-synthesized TiO$_2$ NTAs, rGO-commercial TiO$_2$ NPs, commercial TiO$_2$ NPs, rGO-Ti sheet, rGO and Ti sheet samples, (b) stability of field emission currents from a typical field emission device (conformal coated rGO on annealed TiO$_2$ NTAs hybrid structure) at 230 voltages and (c) Fowler-Nordheim characteristics curves for different field emission devices (conformal coated rGO on annealed TiO$_2$ NTAs, annealed TiO$_2$ NTAs, as-synthesized TiO$_2$ NTAs and commercial TiO$_2$ NPs).

Figure 4. Field emission characteristics of typical field emission devices based on rGO-TiO$_2$ NTAs hybrid nanostructures (a) Field emission characteristics of different field emission devices (conformal coated rGO on annealed TiO$_2$ NTAs hybrid structure, annealed TiO$_2$ NTAs, as-synthesized TiO$_2$ NTAs, rGO-commercial TiO$_2$ NPs, commercial TiO$_2$ NPs, rGO-Ti sheet, rGO and Ti sheet samples), (b) stability of field emission currents from a typical field emission device (conformal coated rGO on annealed TiO$_2$ NTAs hybrid structure) at 230 voltages and (c) Fowler-Nordheim characteristics curves for different field emission devices (conformal coated rGO on annealed TiO$_2$ NTAs, annealed TiO$_2$ NTAs, as-synthesized TiO$_2$ NTAs and commercial TiO$_2$ NPs).
TiO$_2$ NTAs shows a good electrical contact between the TiO$_2$ nanotubes and rGO as well as it provides a long term stability of field emission currents.

Field emission is generally analyzed using the Fowler–Nordheim (F–N) theory, which describes the tunneling of electrons through a potential barrier formed at the interface between a metal surface and vacuum[46]. According to F–N theory, the field emission current ($I$) or current density ($J$) is related to work function ($\Phi$) of the material and external electric field ($E$) through the relation,

$$J = (A\beta^2E^2/\Phi) \exp (-B\Phi^{3/2}/\beta E)$$

or

$$\ln (I/E^2) = \ln (A\beta^2/\Phi) - B\Phi^{3/2}/\beta E$$

Where $J$ is the current density, $E$ is the applied field, $\Phi$ is the work function of the emitting materials (~4.4 eV for TiO$_2$), $\beta$ is field enhancement factor and $A$ and $B$ are constants with values of 1.56 \times 10^{-6} (A V^{-2} eV)$ and 6.83 \times 10^{5} (V \mu m^{-1} eV^{-3/2}) respectively. The value of $\beta$ is related to the emitter geometry, crystal structure, vacuum gap and spatial distribution of the emitter centres. The F–N plots of log($I/E^2$) versus 1/$E$ for different samples are shown in the Fig. 4c and different slopes for the TiO$_2$ NTAs before and after being modified with rGO conformal coating are observed. Good linearity within the measurement range suggests that electron emission by samples follows the F–N plots and the emission is indeed due to a vacuum tunnelling process. Moreover, the work function ($\Phi$) of as-synthesized samples was calculated using photoelectron emission (PEE) technique. The PEE spectra for conformal coated rGO due to a vacuum tunnelling process. Moreover, the work function ($\Phi$) of as-synthesized samples was calculated using photoelectron emission (PEE) technique. The PEE spectra for conformal coated rGO on annealed TiO$_2$ NTAs, annealed TiO$_2$ NTAs, as-synthesized TiO$_2$ NTAs and commercial TiO$_2$ NPs samples are shown in Figure S12a (see Supplementary Information). The plausible schematic model of edge states and corresponding energy-band diagrams of field emission from conformal coated rGO on annealed TiO$_2$ NTAs, annealed TiO$_2$ NTAs, as-synthesized TiO$_2$ NTAs and commercial TiO$_2$ NPs samples is shown in Figure S12b (see Supplementary Information). It reveals that conformal coated rGO on annealed TiO$_2$ NTAs consist of higher ratios of C–O–C ether chain edge states, which causes the potential barrier of electrons have to overcome in vacuum to be diminished, resulting in a lower work function of conformal coated rGO on annealed TiO$_2$ NTAs hybrid structure. So electrons tunnel through near the top of the barrier and can easily pass across the full barrier width. The experimentally obtained work function value for conformal coated rGO on annealed TiO$_2$ NTAs, annealed TiO$_2$ NTAs, as-synthesized TiO$_2$ NTAs and commercial TiO$_2$ NPs are ~3.1 eV, ~3.3 eV, ~3.9 eV and ~4.4 eV respectively. Thus, the slope of F–N plots and calculated values of work function, we can easily estimate the field enhancement factors $\beta$ from equation 2. It is ~6000, ~5000, ~700 and ~600 for conformal coated rGO on annealed TiO$_2$ NTAs, annealed TiO$_2$ NTAs, as-synthesized TiO$_2$ NTAs and commercial TiO$_2$ NPs, respectively. Thus, from the above results, we can observe that the field enhancement factor value is higher for conformal coated rGO on annealed TiO$_2$ NTAs samples compared to the other previous reported TiO$_2$ nanostructures[4,16,37,47,48]. This can be due to the combined effect of rGO and TiO$_2$. The rGO provides an additional interface to the large curvature of TiO$_2$ NTAs because 2D rGO nanosheets have a sufficient no. of the delocalized $\pi$ electrons available[49,50], which act as electron injection carriers and, as a result the difference between the Fermi levels of rGO and conduction band of TiO$_2$ in rGO- TiO$_2$ hybrid structure is reduced, consequently the work function also get reduced. The experimentally obtained work function of the rGO-TiO$_2$ hybrid structure is less than that of the TiO$_2$, which can be seen from Figure S12a. According to the results presented here, TiO$_2$ NTAs possess moderate high performance field emission property, which is enhanced remarkably after being modified with conformal coating of rGO. It is mainly attributed to low work function and high aspect ratio. Thus, the introduction of rGO on the surface of TiO$_2$ NTAs can increase the number of emitters and tunneling probability, which leads to higher field emission for the hybrid emitters. These results showed that the field emission properties of TiO$_2$ NTAs can be tailored by conformal coating of rGO on its surface. The improved field emission characteristics in conformal coated rGO-TiO$_2$ NTAs hybrid structures are attributed to the contribution of low work function of the metal and the field free vacuum ($E_f$)) (Figure S12b). The ohmic contact with non-rectifying barriers allows electrons to be easily injected from the conduction band of TiO$_2$ to Fermi level of rGO under external electric field. Then, the electrons go from TiO$_2$ to rGO, then to vacuum through subsequent F–N tunneling under the low turn-on field. These results indicate the great shift of Fermi level towards higher energy, as shown in schematic diagram (Figure S12b). Therefore, compared to TiO$_2$, a Fermi level with higher energy is observed for conformal coated rGO-TiO$_2$ NTAs hybrid structures. This shifting improves both the conductivity and field emission properties of conformal coated rGO-TiO$_2$ NTAs sample. This work demonstrates the approach to convert TiO$_2$ nanotube arrays into conformal coated rGO-TiO$_2$ NTAs hybrid structure. As a result, it creates more acceptor and donor states (both) above the valence band maximum and below the conduction band minimum in the band gap of TiO$_2$ nanotube, which helps to reduce work function of hybrid structure (clearly shown in Figure S12a and S12b). Therefore, the conformal coated rGO-TiO$_2$ NTAs is better hybrid structure for obtaining the high-performance field emission applications.
Discussion

A highly-efficient method to produce hybrid structure of rGO nanosheets conformal coated on vertically aligned TiO2 nanotubes 3D arrays for enhanced field emission display applications has been successfully demonstrated. The structural characterization of rGO conformal coated annealed TiO2 NTAs exhibits the formation of a highly ordered 3D NTAs with a pure anatase phase and good crystallinity. SEM and TEM results indicated that the average diameter and length of TiO2 NTAs are about ~110 nm and ~2 μm respectively, at optimum anodization condition (anodization at 40 V for 4 h). The HRTEM image of TiO2 NTAs shows high quality lattice fringes without lattice distortion, which clearly demonstrates the improvement of crystal line quality of TiO2 NTAs after annealing.

The rGO conformal coated TiO2 NTAs exhibited high emission current and excellent field emission stability with a low turn on field compared to commercial TiO2 nanoparticles (NPs), as-synthesized TiO2 NTAs, annealed TiO2 NTAs samples, pure Ti sheet, rGO nanosheets and rGO on pure Ti sheet substrate samples. The linearity of the F-N plots confirms that the process is governed by the Fowler-Nordheim equation, based on tunneling electron emission. Thus, this simple, effective and robust approach provides new prospects to develop highly-efficient electron sources for stable and ultra low turn on field FE devices based on the rGO conformal coated TiO2 NTAs hybrid nanostructures.

Methods

Materials. The titanium (Ti) sheet (99.8% purity, size ~0.5 mm × ~20 mm × ~15 mm) was purchased from Sigma-Aldrich. Graphite flakes (SP-1 graphite, ~150 μm size) was purchased from Bay Carbon Corporation. Ammonium fluoride (NH4F), ethylene glycol (C2H4O2), hydrogen peroxide solution (H2O2), potassium permanganate (KMnO4) and all other reagents were of analytical grade (AR) and used as received without further purification. Double distilled water was used throughout the experiments.

Synthesis of vertically aligned TiO2 3D NTAs. Vertical aligned TiO2 3D NTAs were fabricated by anodic oxidation of 0.5 mm thick Ti sheet. Prior to anodization, titanium sheets were first mechanically polished with different grades of emery papers and final finishing was done with zero grade paper. Then, Ti sheets were ultrasonically (frequency; 25 kHz) cleaned in acetone and ethyl alcohol for 10 minutes in each solution. This process was repeated three times to get nearly clean Ti-sheet and then dried in air at room temperature. The synthesis process for highly oriented architecture of 3D TiO2 nanotubes arrays using anodization technique is illustrated in Fig. 1. The electrochemical anodization of Ti sheet was carried out in a two-electrode cell, with a platinum foil as counter electrode, which was immersed in electrolytic solution in a beaker. The distance between the two electrodes was 2 cm. Both electrodes were placed parallel to keep constant flux lines or the uniform current between the electrodes. Two Cu-wires were used for making the connection through the electrodes. These electrodes were mounted on glass rods using rapid repair material (self polymerizing powder and liquid). Electrolyte was comprised of 0.3 wt% NH4F and 5.0 vol% deionized water in an ethylene glycol solution. The anodization procedure was carried out at different applied potential of ~30 V, ~40 V and ~50 V as well as for various time intervals of 1.5, 2.5, 3.5 and 4.0 hours to optimize the anodization process. The electrochemical experiments were carried out at room temperature under the assistance of magnetic stirring. After anodization, the samples were rinsed with deionized water to remove any unwanted ions on the surface of the TiO2 NTAs samples and dried in air. The optical images of as-synthesized TiO2 3D NTAs samples at different anodization voltage as well as for various time intervals are shown in Figure S13 (see Supplementary Information). The detailed electrochemical conditions with calculated length and diameter of tubes are listed in Table TS1 and TS2 (see Supplementary Information). The surface morphology of as anodized samples was observed by scanning electron microscope and found that the optimum condition for anodization process with large outer diameter of tube is occurred at 40 V for 4 hours. After optimizing the condition, the obtained amorphous nanotube arrays samples were annealed at 500°C for 2 hours with heating and cooling rates of 2°C/min to obtain pure anatase phase. The high-resolution optical micrograph image of TiO2 NTAs at 4 V anodization voltages and 4 hours time intervals at different scale is shown in Figure S14 (see Supplementary Information) at different scale.

Synthesis of reduced graphene oxide nanosheets. Reduced graphene oxide nanosheets were as-synthesized by the oxidation of graphite flakes using improved method proposed and established by James Tour et al.34, 3.0 g graphite flakes was added in solution of H2SO4/H3PO4 (360:40 mL) and 18.0 g KMnO4 producing a slight exotherm at 35–40°C temperature. The mixture was continuously stirred at 50°C for 12 h and allowed to cool to room temperature. It was poured onto ice (400 mL) and treated with hydrogen peroxide solution (H2O2, 30 wt%, 3 mL). The mixture was sifted through a metal U.S. Standard testing sieve (WS. Tyler, 300 μm) and filtered through polyester fiber (Carpenter Co.). Then, it was centrifuged (4000 rpm, 4 h) and the supernatant was decanted away. The remaining solid material was washed in succession with 200 mL of 30% HCl, water and ethanol. For each wash, the mixture was sifted through the U.S. Standard testing sieve and then filtered through polyester fiber with the filtrate being centrifuged and the supernatant decanted away. The remaining material after this extended, the multiple-wash process was coagulated with 200 mL of ether. The resulted suspension was filtered over a PTFE membrane with 0.45 μm pore size. The powder obtained on the filter was dried overnight at room temperature under vacuum and 5.8 g graphene oxide product was obtained. Finally, 0.1 g graphene oxide
was dispersed into 100 mL distilled water via ultrasonication and then NaBH₄ was added to reduce the graphene oxide nanosheets to graphene nanosheets at 80°C.

**Fabrication of reduced graphene oxide conformal coated TiO₂ 3D nanotubes arrays.** The as-synthesized rGO 2D nanosheets were ultrasonically dispersed into ethanol, followed by ultrasonication at 25 kHz frequency for 1 hour to form a homogeneous suspension with a concentration of 0.05 mg/mL. The resulted solution was drop-casted on the samples perpendicular to the orientation of the NTAs (40 V, 4 hours, 500 °C) and dried in air, as shown in Fig. 1. It formed Ti-O-C bonding between TiO₂ and rGO, which is further confirmed by XPS. The dilution of rGO played a critical role to obtain enhanced FE properties as well as easy to coat conformally around the nanotube walls. Similarly, the rGO conformal coated TiO₂ NTAs samples are prepared four times for reproducibility test and are labeled as sample 1, sample 2, sample 3 and sample 4.

**Characterization.** For phase identification and gross structural analysis, the structural characterization was performed using X-ray diffractometer (XRD, Rigaku: MiniFlex, Cu Kα₁, λ = 1.5406 Å). The surface morphology, length and diameter of TiO₂ nanotubes were determined by scanning electron microscopy (SEM, Model No. EVO-MA 10 VPSEM). The microstructural studies were carried out using high-resolution transmission electron microscopy (HRTEM, Model No. Technai G20-twin, 300 kv) with super twin lenses having point and line resolution of 0.144 nm and 0.232 nm, respectively equipped with energy dispersive x-ray analysis (EDAX) facility. Raman spectra were obtained using Renishaw InVia Raman spectrometer, UK with an excitation source of 514.5 nm. The XPS analysis was carried out in an ultra-high vacuum (UHV) chamber equipped with a hemispherical electron energy analyzer (Perkin Elmer, PHI1257) using non-monochromatized Al Kα source (excitation energy of 1486.7 eV) with a base pressure of 4 × 10⁻¹⁰ torr at room temperature. The work function has been evaluated through open-counter photoelectron emission (PEE) spectroscopy system.

**Field emission measurements.** The field emission measurements were carried out at room temperature under a vacuum of ~10⁻⁹ torr. A rod like copper probe with a cross section of about 0.6 mm² was served as an anode and all samples; conformal coated rGO on annealed TiO₂ NTAs, annealed TiO₂ NTAs, as-synthesized TiO₂ NTAs and commercial TiO₂ NPs on the Ti substrate were fixed onto ITO as the cathode under same condition. Field emission measurements were performed in high vacuum to prevent the rGO from absorbing oxygen. The spacing between the electrodes was maintained at 100–500 μm, 100 μm was kept as an optimum distance. A dc voltage sweep from 300 to 1100 V was applied to the samples in steps of 20 V to generate the electric field (E). The emission current was monitored by an electrometer (Keithley 6514) with picocammere sensitivity.

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B.K.G. wrote the manuscript and analyzed the data. B.K.G. conceived and supervised the project. All ray diffraction and Raman measurements.

B.K.G. conceived the concepts of the research. B.K.G., Y.A. and P.K. designed and fabricated the samples, for financial assistance to carry out this work. P.K. and J.D. performed the X

The authors wish to thank Prof. R.C. Budhani, Ex-Director, CSIR-NPL., New Delhi for his keen interest in the work. The authors are thankful to Prof. O.N. Srivastava (Banaras Hindu University, Varanasi) and Mr. H.K. Maini (Ex Scientist NPL, New Delhi), for his encouragement. The authors gratefully acknowledged the University Grant Commission (UGC) and Council of Scientific and Industrial Research (CSIR), Govt. of India for financial assistance to carry out this work.

Acknowledgments
The authors wish to thank Prof. R.C. Budhani, Ex-Director, CSIR-NPL., New Delhi for his keen interest in the work. The authors are thankful to Prof. O.N. Srivastava (Banaras Hindu University, Varanasi) and Mr. H.K. Maini (Ex Scientist NPL, New Delhi), for his encouragement. The authors gratefully acknowledged University Grant Commission (UGC) and Council of Scientific and Industrial Research (CSIR), Govt. of India for financial assistance to carry out this work.

Author Contributions
B.K.G. conceived the concepts of the research. B.K.G., Y.A. and P.K. designed and fabricated the samples, and set up models. V.N.S. performed the scanning electron microscopy and transmission electron microscopy measurements. P.K. and J.D. performed the X-ray diffraction and Raman measurements. Y.K. and P.K. carried out the field emission measurements. G.K. prepared the figures. G.K., R.K.G. and B.K.G. wrote the manuscript and analyzed the data. B.K.G. conceived and supervised the project. All authors contributed to revising the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Agrawal, Y. et al. High-Performance Stable Field Emission with Ultralow Turn on Voltage from rGO Conformal Coated TiO₂ Nanotubes 3D Arrays. Sci. Rep. 5, 11612; doi: 10.1038/srep11612 (2015).

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