In-flight gas phase growth of metal/multi layer graphene core shell nanoparticles with controllable sizes

Saurabh K. Sengar¹, B. R. Mehta¹, Rakesh Kumar¹ & Vinod Singh¹,²

¹Thin Film Laboratory, Indian Institute of Technology New Delhi, India-110016, ²Department of Applied Physics, Delhi Technological University, Delhi, India-110042.

In this report, we present a general method for a continuous gas-phase synthesis of size-selected metal/multi layer graphene (MLG) core shell nanoparticles having a narrow size distribution of metal core and MLG shell for direct deposition onto any desired substrate kept under clean vacuum conditions. Evolution of MLG signature is clearly observed as the metal-carbon agglomerates get transformed to well defined metal/MLG core shell nanoparticles during their flight through the sintering zone. The growth takes place via an intermediate state of alloy nanoparticle (Pd-carbon) or composite nanoparticle (Cu-carbon), depending upon the carbon solubility in the metal and relative surface energy values. It has been also shown that metal/MLG nanoparticles can be converted to graphene shells. This study will have a large impact on how graphene or graphene based composite nanostructures can be grown and deposited in applications requiring controllable dimensions, varied substrate choice, large area and large scale depositions.

The observation of unique combination of properties, such as, mechanical strength, good thermal and electronic conductivity, zero band gap and ballistic electron transport in the ‘ideal two dimensional’ graphene structure have resulted in an unprecedented research interest¹–⁶. In the last few years, graphene-metal nanoparticle composites have resulted in an improved, new and novel applications⁷–¹⁸. State of the art chemical vapor deposition allows the growth of graphene sheets on metal foils¹⁹. The restrictions of graphene growth only on few selected metal substrates and the requirement of transfer are the main nagging issues for using the graphene-metal nanostructures prepared by the above method in applications requiring large scale and large area deposition such as catalysis, Li ion batteries and solar cells⁷–⁹,¹⁴,²⁰. The multistep chemical processes, used for growing graphene-metal composites involve simultaneous or separate reduction of graphene oxide with metal precursors yielding undefined configurations, high dispersibility and wide distribution of nanoparticle sizes and graphene thickness²¹. Chemical contamination incorporated into metal nanoparticles and graphene layers during reduction process is known to affect optical, electronic and catalytic properties²²–²⁴. The average nanoparticle size and distribution of Au nanoparticles grown by physical vapor deposition methods is linked to the number of graphene layers²⁵,²⁶. In physical methods of metal nanoparticles deposition, both particle density and particle size depend upon deposition time, and thus, nanoparticle size cannot be controlled independently. Further, in earlier reports on metal-carbon core shell nanoparticles prepared by arc discharge, metal carbide (in place of metal) core was found to be encapsulated in polyhedron carbon shells²⁷. In another report, the graphite layer surrounding the metal nanoparticle was found to appear throughout the sample²⁸. Broad size distribution and phase inhomogeneity are the other issues in these metal/graphite core shell nanoparticles prepared by arc discharge method²⁷,²⁸. Metal/graphite nanoparticles prepared by chemical methods have been reported to have low crystallinity, chemical contamination, and broad size distribution²⁹,³⁰. Therefore, these methods are not suitable for applications requiring controllable size, narrow size distribution and well defined graphene-metal interface such as biomedical and opto-electronic applications. In this report, a gas phase synthesis methodology for growing size selected metal/multilayer graphene (M/MLG) core shell nanoparticles with controllable size, narrow size distribution, well defined metal core and graphene shell is described.

Results

M/MLG core shell nanoparticles were fabricated using an integrated gas phase synthesis setup of which schematic diagram is shown in Fig. 1. The structural properties of the nanoparticles sintered at different temperatures during
in-flight sintering were carried out using glancing angle x-ray diffraction (GAXRD). GAXRD diffractograms of the Pd-C nanoparticles sintered at different temperatures are shown in Fig. 2a. The unsintered Pd-C agglomerates show high intensity peak at 2θ (d-value) of 39.36° (2.287 Å) corresponding to Pd-C solid solution in face centered cubic (FCC) phase having carbon atoms at interstitial positions. In addition, a relatively low intensity peak at 40.04° (2.250 Å) corresponding to Pd FCC phase is also evident. A lattice parameter of 2.309 Å is reported in Pd-C nanoparticles due to the incorporation of carbon at the interstitial positions. This behavior is similar to that observed in case of Pd-H system where hydrogen incorporation at the interstitial positions results in lattice expansion due to formation of α-palladium hydride phase. In nanoparticles sintered at 300°C and deposited at RT, intensity of the Pd peak increases, accompanied by a decrease in Pd-C peak intensity and shift towards higher 2θ value. On further increasing the sintering temperature to 500°C, intensity of the Pd peak becomes higher than that of the Pd-C peak indicating that more and more of the carbon phase segregates out as the sintering temperature is increased. Finally, in nanoparticles sintered at 700°C, only Pd peak is observed with nearly complete segregation of carbon phase out of the Pd lattice. Unlike, Pd-C system, XRD peaks corresponding to the FCC Cu phase remains unchanged in all the samples sintered at different temperatures and stay at 43.275° (2.090 Å), as shown in Fig. 2b. This indicates that small carbon solubility has negligible effect on Cu lattice.

Morphological changes in Pd-C and Cu-C nanoparticles on sintering were studied using high resolution electron microscopy (HRTEM). HRTEM micrographs of the Pd-C nanoparticles sintered at different temperatures are shown in Figs. 3a–d (also see supplementary Fig. S1). It is clear from Fig. 3a that the initial agglomerates consist of 2–5 nm sized primary nanoparticles as revealed by lattice fringes (2.390 Å) corresponding to Pd-C alloy. On increasing the sintering temperature to 300°C, particles start merging as compaction process is initiated (Fig. 3b). Thereafter, at a sintering temperature of 500°C, compact, quasi-spherical and crystalline nanoparticles having carbon layers in the shell around the inner core are formed (Fig. 3c). Finally, sintering at 700°C results in the formation of nearly spherical, Pd/C core shell nanoparticles with carbon shell clearly visible in the HRTEM image (Fig. 3d). In the Pd/C core shell nanoparticle sintered...
at 700 °C a d-value of 2.340 Å is obtained from the lattice fringes of the Pd nanoparticle-core. The d-values of Pd-C and Pd phases using TEM are larger in comparison to those calculated from XRD. Slightly larger value of d-spacing as calculated by TEM in comparison to XRD may be due to the localized measurement in electron diffraction technique and a possibility of lattice expansion at smaller sizes. The d-value obtained from XRD is the average value calculated from a larger sample having widely varying sizes of clusters especially in unsintered samples while the d-value obtained from HRTEM measurement corresponds to a specific nanoparticle. A large change in the lattice constant has been reported in a number of material systems. Similar trends in the morphological changes of Cu-C nanoparticle system during the conversion of Cu-C agglomerates to Cu/C core shell nanoparticles are observed. In the HRTEM images (Figs. 3e–h) lattice fringes with inter-planar spacing of 3.95 Å and 2.50 Å are observed at different locations in case of agglomerated Cu-C particles. These values differ with the corresponding bulk values of 3.40 Å and 2.09 Å for graphite and Cu, respectively. These differences may be due to lattice expansion at smaller sizes of metal and carbon clusters in the agglomerates at RT. Again the difference between the d-value of Cu calculated from XRD and HRTEM may be due to the localized measurement in HRTEM and widely varying cluster sizes in unsintered samples. On sintering at a temperature of 300 °C, the compaction process gets initiated as shown in Fig. 3f. At 500 °C sintering, formation of the Cu nanoparticle core and carbon shells is initiated (Fig. 3g). On further increasing the sintering temperature to 700 °C quasi spherical metal nanoparticle core enveloped by well formed carbon shell around them is clearly observed in the HRTEM micrograph (Fig. 3h).

To investigate the nature of carbon, micro-Raman measurements of Cu-C and Pd-C nanoparticles sintered at different temperatures were carried out and are shown in Figs. 4a and b, respectively. In the initial Cu-C and Pd-C agglomerates without any sintering, no Raman peak in the entire measurement range of 1100–3200 cm⁻¹ is observed. In the unsintered samples, carbon is present in the form of amorphous and nanosized carbon clusters or Pd-C clusters which are loosely connected to form agglomerates. A significant portion of carbon is present in the form of Pd-C and is expected not to show any D and G peak. In an earlier report, similar results on the absence of Raman D and G peaks in unsintered Ti-B-C films have been reported. G peak is due to the relative motion of sp² carbon atoms while D peak is linked to the breathing mode of carbon rings (defects). As shown in Fig. 4a, G peak lie at 1584 cm⁻¹ and I_D/I_G ratio is found to be 0.73 in the Cu-C nanoparticles sintered at 300 °C. These values of G peak and I_D/I_G ratio correspond to either amorphous carbon or nanocrystalline graphite. The absence of second order 2D peak which is the second order overtone of D peak in Cu-C samples sintered at 300 °C indicates that it is amorphous.
carbon and not nanocrystalline graphite. The possibility of diamond like carbon instead of amorphous carbon has been ruled out as the I_D/I_G ratio is >0.2. On increasing the sintering temperature to 500°C, G peak appears at 1594 cm\(^{-1}\) while the I_D/I_G ratio increases to 1.08 in Cu-C nanoparticles. Finally on sintering at 700°C, G peak remains constant and I_D/I_G ratio increases to 1.15 in Cu-C nanoparticles. Further, in Cu-C nanoparticles sintered at 700°C, 2D peak appears at 2685 cm\(^{-1}\). According to the 3-stage theory, G peak position near to 1600 cm\(^{-1}\) and a continuous increase in I_D/I_G ratio indicates that amorphous carbon gets converted to nanocrystalline graphite on increasing the sintering temperature. Further, in Cu-C nanoparticles sintered at 700°C, I_D/I_G ratio increases to 1.08 while the ID/IG ratio remains nearly same at 0.91. Finally at 700°C, I_D/I_G ratio remains nearly same at 0.91. It is observed from the HRTEM images in Figs. 5a and 5b that nanoparticle core is spherical and has a very well defined size (D_g) of 29.9 ± 2.9, 21.9 ± 1.7 and 18.3 ± 1.4 nm in the sintered Cu/MLG core shell nanoparticles (see supplementary Figs. S2 and S3). In Cu/MLG core shell nanoparticles having D_m = 35 nm, the average size of the Cu core and the thickness of the MLG shell are 14.5 ± 0.9 nm and 2.3 ± 0.6 nm, respectively, as shown in the size histograms in Fig. 5c. In case of Pd/MLG samples too, nearly same values of sizes are observed (supplementary Figures S2 and S3). From the comparison of HRTEM micrographs of Pd/MLG and Pd (without any carbon) nanoparticles it is clear that unlike Pd/MLG, Pd nanoparticles don’t have any MLG like shell structure (supplementary Fig. S2). With increase in the size of the nanoparticle core, the width of the graphene shell is observed to decrease. This may be due to the difference in solubility of carbon in metal nanoparticle of different sizes. The Raman spectra of size selected core shell Cu/MLG nanoparticles having D_m values of 35, 52 and 68 nm is shown in Fig. 5d. The I_D/I_G ratio for the Cu/MLG nanoparticles having D_m values of 35, 52 and 68 nm comes out to be 1.66, 1.49 and 0.79, respectively. As discussed earlier, this indicates that properly sintered spherical shaped Cu/C nanoparticles have a shell comprising of multi

![Figure 5](https://www.nature.com/scientificreports/)

**Figure 5 | Size selected Cu/MLG nanoparticles.** [(a), (b)] HRTEM micrographs of Cu/MLG nanoparticles with mobility equivalent diameter (D_m) of 68 and 52 nm, respectively. (c) Histogram showing size distribution of shell thickness (D_s), Core diameter (D_c) and nanoparticle size (D_g) for Cu/MLG nanoparticle having D_m = 35 nm. Narrow size distribution, control of nanoparticle size by controlling electrical mobility diameter and well formed nanoparticle core and graphene shell are clearly visible. (d) Raman Spectra of size selected Cu/MLG nanoparticles for D_m = 35, 52 and 68 nm showing decrease in graphene layers with increase in nanoparticle size.
layer graphene. The continuously decreasing $I_{\text{G}}/I_{\text{2D}}$ ratio indicates that the number of graphene nanolayers decrease on increasing the particle size. Decrease in number of graphene layers on increasing the $D_m$ value is consistent with HRTEM of Cu/MLG core shell nanoparticles.

Wet chemical etching was used to remove the metal core in order to obtain the MLG shells (Fig. 6a) from the Cu/MLG core shell nanoparticles using FeCl$_3$ solution. After chemical etching MLG shell structures consisting of about 3–4 graphene layers can be easily seen in the HRTEM image (Fig. 6b). The selected area electron diffraction (SAED) patterns (Fig. 6b) with six fold symmetry confirm the MLG shell structure after removing the metal core. The Raman spectrum (Fig. 6c) shows an increase in $I_{\text{D}}/I_{\text{G}}$ peak ratio from 1.59 to 2.18 after etching the Cu from the nanoparticle core which may be due to defects created during etching. To show the universality of the present method to spray coat the M/MLG nanoparticles onto any desired substrate, Cu/MLG core shell nanoparticles were deposited on to Si (polished and textured) and interdigitated Au electrodes (Au-Cr/SiO$_2$/Si). Scanning electron microscopy (SEM) images of Cu/MLG nanoparticles ($D_m = 52$ nm) grown on polished Si, textured Si and interdigitated Au electrodes fabricated on SiO$_2$/Si substrates are shown in Figs. 7a–c. Optical and electronic properties of the Cu/MLG nanoparticles were also studied. Reflectance spectrum of Cu/MLG nanoparticles ($D_m = 52$ nm) deposited on polished Si is shown in Fig. 7a (inset). Reflectance spectrum of Cu/MLG nanoparticles deposited on polished Si shows a large drop in total reflectance in comparison to the bare Si. Electronic properties of Cu/MLG nanoparticles were carried out in a field effect transistor (FET) structure where n-doped Si substrate is acting as back gate. The source-drain resistance versus back-gate voltage $V_{\text{BG}}$ curve for the Cu/MLG
nanoparticles at room temperature is shown in Fig. 7c (inset). Dirac point at a negative gate voltage indicates unintentional n-type doping of the Cu/MLG nanoparticles.[41]

**Discussion**

On the basis of the results of the structural, morphological and Raman studies, formation of M/MLG core shell nanoparticles in both the Pd-C and Cu-C material systems can be understood in terms of temperature dependence of carbon solubility in metals and the relative surface energy values. The solubility of carbon in different metals forming solid solution is given by the following equation based on Henrian behavior for the carbon activity[42]

$$x_c = \frac{\exp \left( -\frac{\Delta H}{RT} \right) \exp \left( \frac{\Delta S}{R} \right)}{1 - 2x_c}$$  

(1)

Where $x_c$ is the carbon content in mole fraction, $\Delta H$ is the partial molar enthalpy of carbon in solution, $\Delta S$ is the partial molar excess entropy of carbon in solutions, $R$ is the gas constant and $T$ is the absolute temperature. Using the reported values of $\Delta H$ ($-30.9$ kJ/mol) and $\Delta S$ ($-0.136$) an increase from $3$ ppm at RT to $15800$ ppm at $700$ °C of carbon solubility in Pd lattice is calculated.[43] Observed shift (Fig. 2a) in XRD peak positions of Pd indicate that Pd-C solid solution is formed in the intermediate growth stage. Carbon solubility in Cu at RT and $700$ °C is calculated to be $1.5 \times 10^{-4}$ ppm and $2.5$ ppm, respectively, using the reported values of $\Delta H$ ($-35.1$ kJ/mol) and $\Delta S$ ($-0.532$). Due to low carbon solubility in copper, no change in the XRD peak positions of the Cu (Fig. 2b) is observed in Cu-C system. Average surface energy of carbon in different phases (fullerene, graphite, and carbon black) is quite low (%<250 mJ/m²) in comparison to Pd ($2.05$ J/m²) and Cu ($1.83$ J/m²)[44] In Pd-C system, a solid solution of Pd-C is formed on sintering, higher the sintering temperature more is the carbon solubility. As particles come out of the sintering zone, temperature decreases and hence carbon atoms segregate out of the Pd lattice and migrate to surface due to their low surface energy. Lattice fringes corresponding to Pd-C and Pd (Figs. 3e and 3h) are observed in unsintered agglomerates and nanoparticles sintered at $700$ °C, respectively. In case of Cu-C system, initially agglomerates consist of copper and carbon primary nanoparticles. At high temperatures Cu-C composite nanoparticle having Cu, C and mixed phases are formed, as confirmed by lattice fringes in the HRTEM images (Figs. 3a and 3b) of unsintered agglomerates and sintered agglomerates at $300$ °C. As Cu-C composite nanoparticle is sintered at higher temperature carbon segregates to the outer surface due to lower surface energy value while Cu segregates to the inner core. As both sintering and cooling takes place in-flight in the homogeneous gaseous medium, quasi spherical nanoparticles having Pd/Cu core and MLG shell are formed. It is necessary to mention that in both the material systems optimized sintering temperature is important for the growth of MLG shell around the spherical metal nanoparticle core. Under-sintering leads to incomplete compaction while over-sintering is observed to result in re-evaporation of metallic phases, both of which affect the size and size distribution of metal core and MLG shells. Large solubility values of C in Fe, Co and Ni at higher temperatures and large surface energy values of most of the metals (e.g. Pt, Ni, Co, W, Fe, Cr, and Ti) in comparison to carbon allows this method to be used as a general technique for M/MLG core shell nanoparticle growth[45]. Controlled growth of M/MLG core shell nanoparticles having a well defined size and narrow size distribution can be achieved by just manipulating the solubility and surface energy of two components with in-flight sintering temperature.

**Gas phase deposition methods** are ideally suitable for large scale and continuous deposition[46]. Recent report on the continuous growth of semiconductor nanorods onto size selected seed nanoparticles is a novel demonstration of the versatility of the gas phase deposition methods. A careful control of homogenous (Au nanoparticle) and heterogeneous nucleation (GaAs nanorods onto Au) in the gas has been utilized to grow Au-GaAs nanostructures[47]. The present method is a major step forward towards the advantage of depositing graphene based nanostructures in the gas phase mode in comparison to deposition of graphene layer on metal substrate and formation of graphene-metal nanoparticles by multi step chemical and physical processes. In the present study metal-carbon nanoagglomerates were formed by spark generation process using metal (Cu or Pd) and carbon as the two electrodes. Metal and carbon concentration in the intermediate alloy/composite nanoparticle was thus fixed by the relative ablation rates of the two materials during spark generation step[48]. It might be possible to vary the metal-carbon composition by preparing metal agglomerates and then sintering them in the presence of carbon precursor gases (e.g. CH₂). It is expected that this will result in a better control of number of graphene layers in the shell. Further, the M/MLG core shell nanoparticle prepared by the present method can also be incorporated into or onto growing thin films prepared by vacuum or plasma deposition techniques[49]. After etching the metal core, size-dependent electronic and magnetic properties of graphene shells can also be studied[50]. Moreover, the etching step for removing Cu or Pd core might be carried out in gas phase itself so that graphene shells can be deposited onto a desired substrate in the final step.

Synthesis of M/MLG nanoparticles having well defined dimensions and the possibility of their direct deposition on to different substrates will result in new and novel applications. MLG cap around the noble metal nanoparticles can result in significantly reducing the consumption of noble metals in large scale fuel cell applications[11,13]. Most recently, carbon-coated metal nanoparticles have shown exceptionally long term thermal and chemical stability against acidic erosion, reliable linking chemistry and biocompatibility, important for biomedical applications[51]. A well defined MLG shell can enhance the functionalization of drug molecules to metal nanoparticle core, which is useful in a number of drug delivery and bio imaging applications[52]. The enhanced surface contact between metal and graphene layers is known to result in enhancing the H₂ sensing response in graphene-Pd/Pt nanoparticles[53,54]. Graphene-metal (Au/Ag) nanoparticles with efficient molecule absorption property combined with strong surface enhanced Raman sensitivity are excellent substrates for nano-molar level molecular detection[55,56]. M/MLG nanoparticles may have superior electromagnetic interference shielding properties in comparison to individual metal or graphene layers[57]. M/MLG nanoparticles on solar cell surfaces may have superior antireflection properties due to anti-reflection (graphene layers) and surface plasmon (metal nanoparticles) effects[58,59]. SEM images of Cu/CMLG core shell nanoparticles deposited onto textured silicon substrates show possible uniform growth on substrates having non-planar surface using the present method.

**Methods**

The schematic diagram of the integrated gas phase setup used for growing Cu/MLG and Pd/MLG core shell nanoparticles is shown in Fig. 1. It comprises of a spark generator for forming aerosol consisting of metal-carbon nanoagglomerates, particle charger for charging the agglomerates, DMA for selecting the agglomerates of constant mobility equivalent diameter (Dm), sintering furnace for compacting the agglomerates into M/MLG core shell nanoparticles and an ESP for depositing the M/MLG core shell nanoparticles onto any desired substrate. In the spark generator, electrical sparks are generated by applying a pulsating high voltage ($\sim 3$ kV) to the carbon and Cu/Pd electrodes separated by a distance of $1.8$ mm which is kept fixed during material ablation by an electrical stepper motor. Electrical sparks result in the ablation of the electrode material. Spark current controls the electrode ablation rate during material ablation by an electrical stepper motor. Electrical sparks result in the ablation of the electrode material. Spark current controls the electrode ablation rate and can be changed by just varying the spark frequency. Due to very short duration of the pulsed discharge, evaporated material rapidly cools down, resulting in super saturation, leading to the nucleation, condensation and finally to the formation of nanoagglomerates having a large size distribution. Average value of size and size distribution depends on the frequency and carrier gas flow. High purity nitrogen (N₂) with 2% hydrogen gas (as a carrier gas) is used in the complete process starting from the initial step of agglomerate formation to final step of nanoparticle deposition. In the particle charger, agglomerates were charged using a Kr-85 based radioactive isotope. Particle charger is basically a radioactive Kr-85 source which charges the nanoparticles both positively and negatively and is also called a neutralizer. DMA is used to
select the nanoparticle sizes as per their Dm values which are controlled by the flow conditions and voltage applied to the DMA electrodes. By measuring the size distribution of agglomerates just after the spark generator, frequency was adjusted to obtain the peak at the desired Dm value. In the in-flight sintering step, compaction, alloy formation and phase separation processes resulted in the formation of M/MLG core shell nanoparticles. Finally, M/MLG core shell nanoparticles were deposited on the Si substrates to form TEM grids for further characterization.

Three different Pd-C/Cu-C agglomerates having Dm = 35, 52 and 68 nm were selected using the DMA. Diameter of C and Pd/Cu electrodes was kept fixed at 3.2 mm. Spark frequency was kept constant at 200 Hz throughout the experiment. Carrier flow (2% H2 + 98% N2) in the spark generator and aerosol flow in the sintering furnace were kept fixed at 2.5 l/min and 1 l/min, respectively. In the sintering furnace, agglomerates were sintered in-flight at 700 °C, 820 °C and 950 °C, respectively. In the ESP, a constant voltage of 1 kV was applied to the substrate with a fixed distance of 1.5 mm between the substrate and the nozzle.

TEM (FEI-Technai-G20 with a LaB6 filament, operated at 200 kV) was used to study the size and structural properties of nanoparticles. GAXRD (Philips X’Pert, PRO-PAD) measurements at a glancing angle of 1° were carried out to investigate the structural properties. Micro-Raman measurements were performed to study the vibrational properties, using a Renishaw spectrometer at 514 nm of wavelength.

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Author contributions
S.K.S. carried out the majority of the research work towards synthesis and characterization of graphene-metal nanoparticles. R.K. carried out the electrical and optical measurements. V.S. assisted in the synthesis of nanoparticles on different substrates. The present study was conceived and supervised by B.R.M., Schlumberger Chair Professor, Thin Film Laboratory, Physics Department, I. I. T. Delhi, New Delhi, India.

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