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# Nd-Pulsed-Laser Irradiations Coupled with Sonication for Enhancing Growth Rate 2D-Graphene Formation

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# ABSTRACT

The graphene was synthesized by two routes the  $1^{st}$  one application of conventional sonication in case of Tri-chloro-acetic 75%H<sub>2</sub>SO<sub>3</sub> for 60hrs while  $2^{nd}$  is combined sonication (60 hrs) plus laser irradiations for 30hrs. The yields of two routes are collected and compared structurally to check and investigate the effect of laser on structure quality and amount of yield obtained. Results indicated that combined route gave yield higher than conventional route by ratio ~ 39\% .AFM-investigations were performed to characterize nano-structural features of produced graphene . .Furthermore raman spectra were measured to confirm graphene formation .

Keywords : Nd-Laser; Synthesis; Dispersion ; AFM ; Graphene; Raman spectra .

# **I.Introduction :**

Graphene has shown many amazing properties and has numerous numbers of synthesizing techniques [1-11].

In the chemical exfoliation process, the insertion of reactants in the inter-layer space weakens the van der Waals cohesive force. The loosened layer stacking is disrupted when the intercalant decomposition produces a high gas pressure of CO<sub>2</sub> by a rapid annealing to 1050 °C. As a result, the *sp2* lattice is partially degraded into *sp2-sp3* sheet that possesses less  $\pi$ - $\pi$  stacking stability.

Chemical exfoliation can be performed in a suspension known as graphite oxide. The most common method to produce graphite oxide was reported by Hummer [5], where graphite is dispersed into a mixture of concentrated sulfuric acid, sodium nitrate and potassium permanganate at 45  $\$  for a couple of hours. In this compound, the graphite layers remain largely intact and the quest molecules or atoms are located in between. To obtain a few or even single sheet materials, the intercalated reactant is decomposed to produce large amounts of gas in the van der Waals space by chemical or thermal means [6]. Then, rapid annealing to 1050  $\$  generates a CO<sub>2</sub> over pressure and splits the graphite oxide into individual sheets [7]. Graphene oxide solutions are yellow in colour or greenish-blue when non-oxidized graphene sheets are the major constituents [8,9]. GO could be regarded as graphene functionalized by carboxylic acid, hydroxyl and epoxide groups. Hence, these functionalized groups make GO easily dispersed into a few select polar solvents that form an intercalated composite with polar molecules [10, 11]. The intercalation of solvent causes the graphene sheet to swell and lose its mechanical integrity [12]. Nevertheless, large quantities of structural defects introduced by the oxidation process shifts the physical properties away from pristine graphene [13].

The major goal of the present work is to introduce new dispersed matrix for exfoliation of GO to yield graphene with reasonable degree of structural quality.

## **II.Experiments:**

## **II.I.Sample Preparation:**

Nano-graphene was prepared according to the following sequences. 2gm of sieved graphite powder with particles size  $\leq 100 \ \mu m$  (Merck product) was dispersed in 50ml of Tri-chloro-

acetic acid sulfonated by 75%  $H_2SO_3$ . Sonication of the dispersion process was performed via a (WiseClean WUC-DIOH 200W, 40 kHz, ) ultrasonic bath up to 60 hours with two stages of mechanical stirring 1<sup>st</sup> after 3 hrs and 2<sup>nd</sup> after 20hrs each of 1.5hrs duration . Dispersion by bath sonication provided the mechanical disruption that broke apart the graphite flakes into small pieces, which was then stabilized by the solvent system . After sonication process, the dispersion was dark grey in color as described in Fig.1 and left to stand for about 10 hours to allow small aggregates to form.

To remove these aggregates and stabilize the dispersion, the top 15ml of the dispersion was taken out and consequently centrifuged for 40 minutes at ~10,000 rpm. After this primary centrifugation, the top 20ml of the dispersion was decanted carefully by pipette, forming a homogeneous black dispersion which was retained for use.

These steps of synthesis procedures were identically repeated with only one difference each two hrs of sonication, Nd-pulsed laser- irradiations were parallel applied for 1hr with total time = 30 hrs (~  $30 \text{ W/cm}^2$  for 30hrs).



Fig.1 : Graphite dispersed in sulphonated Tri-Chloro-acetic acid.

No sedimentations were noticeable for the first 10 hrs that confirm graphene Oxide (GO) is relatively stable in the applied sulfonated tri-choloroacetic .Finally dimethylhydrazine solution (60%) was added to the warmed mixture ( $70 \,^{\circ}$ C) carefully drop wise at to get graphene in highly pure state, leave solution settled for 25 hrs. Then solution has to be centrifuged in order to dispose of the thicker flakes of nano-graphene as possible.

After graphene separation yield of dried graphene  $1^{st}$  route ~ 0.28g while  $2^{nd}$  route (sonication+laser irradiation) was ~ 0.39 g. This result indicates that  $2^{nd}$  route with laser irradiations is more efficient by 39.2%.

# **II.II.Nd-Laser Irradiation Source :**

The applied laser Nd-pulsed laser has the following parameters: wavelength  $\lambda$ = 1.06 µm, pulsed rate  $\eta$  = 10-3 s. The graphene solution target was irradiated by dose of Nd-laser beam irradiations ~ 30 W/cm<sup>2</sup> for 30hrs. The irradiation was carried out in air without any external heating .The energies of pulsed Nd-laser were sufficient to melt homogeneously the surface and near surface layers.

#### **Nd-Laser irradiations**



Ultrasonic Path with GO Solution



## **II.III. Raman Spectroscopy Measurements :**

The measurements of Raman spectra were carried out on the finally ground powders with Laser wavelength = 632.8 nm (He-Ne laser with power = 1mW) and laser power applied to the site of the sample = 0.4 mW with microscope objective = x20, accumulation time = 1000 - 4000s, up to more than an hour.

# II.IV.Atomic force microscopy (AFM):

High-resolution Atomic Force microscopy (AFM) is used for testing morphological features and topological map (Veeco-di Innova Model-2009-AFM-USA). The applied mode was tapping non-contacting mode. For accurate mapping of the surface topology AFM-raw data were forwarded to the Origin-Lab version 6-USA program to visualize more accurate three dimension surface of the sample under investigation. This process is new trend to get high resolution 3D- surface [14-16].

# **III.RESULTS & DISCUSSION**

The synthesized nano-graphene flakes were investigated by both of raman spectroscopy and x-ray diffraction to prove existence of graphene as shown in Figs.2<sub>a</sub> and 2<sub>b</sub> respectively. As it clear in Fig.2a graphene characteristic peaks (modes) appear in the zone ~ 1400-1650 cm<sup>-1</sup> which fully consistent with [17] while graphene oxide appears ~ 1000 and ~ 2600 cm<sup>-1</sup>.

From Fig.2a one can notify that original experimental raman profile is very crowded and there are many of raman modes overlapped with each other due to GO existence as clear in the refined smoothed raman profile .



Fig.2a : Raman spectra measured for ground graphene flakes .



Fig.2b: X-ray diffraction pattern recorded for graphene flakes .

Although XRD measurements is not accurate tool to identify graphene but in our case as it clear in high resolution XRD profile, it gives sharp peak at two theta ~ 24.8 which is fully consistent with literature [18]. The shift in fingerprint peak position of graphene is attributable to poly crystalline phases interference that existed together with graphene as graphene oxide and small traces from un-reacted graphite.

To characterize nano-structural features of obtained graphene AFM measurements were made applying tapping non-contact mode as shown in Fig.3a-c .

Fig.3a-c shows 2D and 3D-AFM image captured for tiny scanned area ( $0.1x0.1 \ \mu m^2$ ) of graphene. Fig.3c displays honey comb structure for graphene which described in literature by author himself applying STM atomic mode [17].



Fig.3a-c: (a) 3D- image for graphene applying tapping non-contact mode .

- (b) 2D-AFM image for graphene applying tapping non-contact mode .
- (c) Honey comb structure of graphene.



Fig.4: 3D-AFM-visualized image for synthesized nano-graphene .

For accurate mapping of the surface topology AFM-raw data were forwarded to the Origin-Lab version 6-USA program to visualize more accurate three dimension surface of the synthesized nano-graphene see Fig.4 .As it clear in Fig.4 which represent very narrow 3D-scanned area with dimensional 0.16x0.16x0.16  $\mu$ m .The accurate analysis of this figure one can conclude the following facts ; 1<sup>st</sup> the maximum heights gradient ranged in between (1.065 – 1.10  $\mu$ m) orange-red zones ,2<sup>nd</sup> the minimum depth gradient is ranged in between (0.96-0.995  $\mu$ m) pale – dark blue zones .3<sup>rd</sup> higher than 50 % of the scanned area moderate in heights and ranged in between 0.99-1.048  $\mu$ m those represented by blue-green colors .These accurate investigations interpret why graphene has huge unique exposure surface area (≈2600 m2/g) as reported in [19,20] with different gradients on the surface topology in contrast with others carbon-based materials .

# **IV. Conclusions:**

The route of application (sonication + laser irradiation) is more efficient by 39.2% than the ordinary sonication only. The energies of pulsed Nd-laser were sufficient to melt homogeneously the surface and near surface layers and as result enhance dispersed graphene to aggregate forming bigger particles .

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