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# Proton energy loss in multilayer graphene and carbon nanotubes

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#### ABSTRACT

Results of a study of electronic energy loss of low keV protons interacting with multilayer graphene targets are presented. Proton energy loss shows an unexpectedly high value as compared with measurements in amorphous carbon and carbon nanotubes. Furthermore, we observe a classical linear behavior of the energy loss with the ion velocity but with an apparent velocity threshold around 0.1 a.u., which is not observed in other carbon allotropes. This suggests low dimensionality effects which can be due to the extraordinary graphene properties.

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#### 1. Introduction

The use of the interaction of energetic particles with solid matter is a useful tool in fun-28 damental areas of physics and technological applications in material science, biomedicine, 29 nuclear and space industry, material characterization and new spectroscopies applied to 30 nano-science (1). Particle interaction with nanostructures is one of the most interesting phe-31 nomena, where the crucial parameter to study is the amount of deposited energy and how 32 it is transferred to the media. Nowadays, the emergence of true possibilities to obtain ultra-33 thin films allows us to study the phenomena of electronic excitations by energetic ions at 34 very low energies. New research on flat nanostructures, such as graphene (2,3), presents 35 extensive possibilities to study their extraordinary physical and chemical properties under 36 different configurations and environments, including radiation exposure. 37

Graphene is emerging as one of the most attractive materials for particle sieving includ ing gases, liquids and other kind of molecules. A recent review of graphene-based mem branes covers these topics (4). Graphene in a pristine state is impermeable to almost all ther mal atoms and molecules, especially hydrogen and helium under ambient conditions (4–6).
 On the other hand, particles with high kinetic energy can pass through graphene layers,
 but interacting with highly dense electron cloud, transfer much more energy to graphene

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than to bulk systems (6). This conclusion is supported by the recent time-dependent density functional theory (TD-DFT) calculations. For energetic ions, like proton and helium, the
transferred energy per unit path length through electronic excitations turns out to be at
least twice as high as that corresponding to graphite and amorphous carbon (7–9).

51 The kinetic energy, charge state and trajectories of the particles are affected in a different 52 manner by materials which have a well-differentiated electronic and crystalline structure. 53 For particles at energies less than 10 keV, these interactions play a significant role in sput-54 tering, ion implantation and in various characterization techniques (e.g. Low Energy Ion 55 Scattering-LEIS), in which particles probe the first few atomic layers of nanostructured mate-56 rials (10,11). In the energy range considered in this work, experimental data on the stopping cross section in any material is very scarce and deserves to be studied to extend tests of the 57 reliability of theoretical models and semi-empirical data compilations (12,13). These results 58 59 which may present considerable differences with bulk results deserve to be investigated.

In this progress report, we discuss experimental measurements of the electronic energy
 loss of proton beams transmitted through multilayer graphene (MLG) films. The proton inci dent energy range goes from 1 to 10 keV. To our knowledge, there exists no other similar
 data on such a material, to which our experiments could be compared. Hence, we com pare our experiments with data obtained for protons in amorphous carbon and nanotubes
 obtained in our laboratory and abroad. In the Figure 1, we show a sketch of our experiment
 where ions interact with graphene and nanotubes.

#### 2. Experiment

#### 2.1. Sample description

71 We obtained a set of commercial samples from the Graphenea company (14). Briefly, as 72 the company claims, graphene layers were synthesized by CVD method on a Cu substrate. Q9 73 The MLG films were prepared by transferring and stacking independently 10 (10) graphene 74 layers on a user requested substrate, in a multiple transfer procedure (non-AB Bernal config-75 uration). In our case, all samples were transferred onto Quantifoil gold TEM grid substrates 76 (15) and suspended over 2  $\mu$ m holes. The transfer procedure of these samples follows the 77 method described in the work of Ochoa et al. (16) and patented by Graphenea. The nomi-78 nal thickness of these films is 3.45 nm, considering that the theoretical graphene thickness 79 is 0.345 nm. Raman analysis of the stacked graphene layers shows spectra characteristic of 80 graphene, which means the layers are not interacting among themselves. The same result was 81 obtained recently by Chen et al. in fabricating two stacked monolayers (17). 82

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## 2.2. Energy loss measurements

To determine the electronic energy loss, we use the transmission geometry, where ions pass through very thin self-supported films, with thicknesses preferably less than 20 nm, which are less than the proton penetration depth in our low energy range. A brief description of the experimental arrangement is given hereafter. Multilayer graphene samples are placed in front of the beam with a five-axis precision manipulator. The operating pressure in the ion gun system is  $5 \times 10^{-4}$  Pa and the energy loss measurements were performed in the collision chamber with a pressure around of  $4 \times 10^{-6}$  Pa.



121 lons are generated by a Colutron hot discharge source (18), then accelerated, focused, 122 123 and mass and charge selected, to obtain protons with energies in the range of 1 to 10 keV. 124 To avoid damage to the samples, a common practice in this kind of experiment is to dimin-125 ish to a minimum value the ion beam current, which is reduced to obtain fluencies less than  $6 \times 10^9$  protons/cm<sup>2</sup>. The proton energy is measured using a spherical sector electrostatic 126 127 energy analyzer with a resolution of less than 1% at FWHM. Protons are detected using mul-128 tichannel plates MCP in a Chevron configuration. Ions entering the analyzer were detected 129 with an angular acceptance of 0.5°, at 0° and at 3° with respect to the ion beam direction. 130 Detection at 3° is used to avoid an overlap of the incident energy distribution with that 131 coming from the target and used to prevent excessive bombardment of the MCP detectors 132 located at 0°. This overlapping is due to the presence of pinholes in the sample, allowing 133 passage of the incident beam.

In Figure 2, we show the energy distributions for protons after passing through the MLG target, measured at 0° (blue full circles) and 3° (red empty circles). The incident energy was 5 keV. A slight difference in energy is observed which is due to possible path length enlargements caused by the effect of multiple scattering phenomena. The nuclear loss is neglected due to the small scattering angles with respect to the incident direction. Roughly speaking,



Figure 2. This plot shows energy distributions of transmitted protons through the MLG target. These distributions were measured at 0° (blue full circles) and 3° (red empty circles), with respect to the incident direction, with an angular acceptance of 0.5°. The incident beam, indicated by an arrow at 5000 eV, passes through pinholes in the target and its energy distribution overlaps with the right wing of the proton distribution coming from the target. Elastic loss, or nuclear stopping power, is negligible in this case.

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passage through 10 layers of carbon means an estimated nuclear loss of 20 eV at 3° with pro-159 tons at 5 keV (12). We consider that these differences fall within the experimental error. We 160 can observe that the present energy distributions show large tails at the lower energy side 161 in contrast to the energy distributions observed in metallic films which have a Gaussian-like 162 shape (19). To evaluate the proton electronic energy loss, we use the most probable energy 163 in the energy distribution (peak position) and the energy loss, in eV/A, is calculated using 164 the nominal thickness of MLG sample, which corresponds to 3.45 nm. The uncertainty in 165 our energy measurements has been minimized and it is less than 1%, which is equivalent 166 to 10-20 eV. 167

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## 1691703. Experimental results and discussion

171 Figure 3, shows the most probable electronic energy loss, in eV/A, as a function of the pro-172 ton velocity (in atomic units) for protons interacting with carbon allotropes. Red symbols 173 correspond to our measurements on MLG targets. Our results show a linear behavior as 174 a function of velocity, with a surprising and not expected feature, an apparent velocity threshold merge at 0.1 a.u. (250 eV) velocity. Another interesting characteristic is that the 175 176 proton energy loss in MLG is larger as compared with the values obtained for amorphous 177 carbon and nanotubes (12,13,20,21). The isolated point at 0.63 a.u. (10 keV) corresponds 178 to a test measurement to verify linearity. In between 5, and 10 keV, we did not perform 179 measurements, in order to avoid target damage. For comparative purposes, we include 180 data for protons energy loss in amorphous carbon (blue symbols), obtained from data 181 compilations (13). Also shown are the experimental energy losses for protons in carbon 182 nanotubes of different dimensions. Green symbols correspond to proton energy loss data 183 in a multi-wall carbon nanotube with an internal diameter of 5 nm and an external diame-184 ter of 27 nm, which correspond to a wall thickness 11 nm (20). Black symbols correspond to



Figure 3. Proton electronic energy loss in eV/Å as a function of ion velocity in a.u. Exhibited data correspond to targets of graphene multilayers (MLG), amorphous carbon and nanotubes, for two different dimensions, see text. The dotted lines are given to guide the eye to the intercept point of the linear extrapolation. Red symbols correspond to our measurements. Blue symbols are proton energy losses in amorphous carbon (*13*). Green and black symbols correspond to proton energy loss in MWCNT nanotubes, see text (*20,21*).

proton energy loss data in nanotubes with an internal diameter of 74 nm and an external
diameter of 85 nm, corresponding to a wall thickness of 5.5 nm (21). In the case of amorphous carbon and carbon nanotubes, we observe a linear behavior for the energy loss, as a
function of proton velocity, with different slopes.

Our experimental results are compared with recent *ab initio* calculations. These approaches combine classical molecular dynamics and TD-DFT to describe proton inter-action with graphene (7,8). Their study is focused on the energy transfer to graphene when protons travel perpendicular to the hexagonal structure of graphene in two specific points, in the middle of the C-C bond and in the middle of the hexagon. A common feature of their results is the large energy per unit path length transferred to graphene in the energy range of our experiment and the non-linearity of energy loss as a function of velocity. The energy loss as a function of the incident energy (see Figure 4) turns out to be highest for passage between the middle of the C-C bond.

In Figure 5, we repeat part of the information appearing in Figure 3. In this plot, we show a comparison between our experiment and theory. To compare our experiment with the mentioned theories, we take the energy transfer value, corresponding to a single graphene layer as given by those calculations. We multiply that energy transfer by the number of layers of the MLG and then divided by the total thickness of the target, which is 34.5 Å. Symbols in Figure 5 are blue symbols (up and down triangles) correspond to calculations made by Bubin et al. (8), where up triangles correspond to energy transferred by the proton to electrons belonging to the C-C bond and would correspond to the maximum particle energy loss in Figure 4. Down triangles would correspond to the minimum energy loss due to the low electron density present in the hexagon. Red symbols are the same as in Figure 3. Black symbols correspond to calculations performed by Krasheninnikov (7), and have the same meaning as the blue ones. Also shown in Figure 5, for reference, a black line indicates







Figure 5. Proton electronic energy loss in eV/Å as a function of ion velocity in a.u. For comparison with our measurements, we include data from recent theoretical calculations, which combine both classical molecular dynamic and time-dependent density functional theory. Blue data correspond to data from ref. (8) and black symbols correspond to data calculated in the same framework, ref. (7), see text for detailed explanation of symbols. The red line is the linear approximation to our experimental data. The amorphous carbon data are approximated by a black solid line, which also represent, approximately, data obtained from the MWCNT (5,22), see Figure 3.

proton energy loss in amorphous carbon. Both calculations indicate that the energy losses
in graphene should be higher than in amorphous carbon. Our experimental data appear to
lie close to the high limit (middle C–C bond) of the energy losses, predicted by reference (7)
and, the ring center contribution to energy loss, which corresponds to the lowest electron
density region, agree with the amorphous carbon data.

Despite similar methods of calculations, there are significant differences between results
of the two theoretical but discussion of these approaches is beyond the scope of this work.
However, both theories agree in their prediction of high energy transfer to graphene by
energetic protons.

While the higher energy loss trend of the experimental data agrees on the whole with the
indications of theory that the energy losses in graphene are higher than in amorphous carbon, one also needs to consider other possible reasons such as contaminations. There are

different sources of contamination one can consider. These are water molecule and hydro-277 278 carbon adsorption, and also some residual PMMA in transferring graphene layers. The work 279 of Ochoa et al. (16), on which the Graphenea samples are based, suggests that the main contaminant should be water. Their model of a graphene multilayer derived from a detailed 280 281 spectroscopic ellipsometry study and XPS includes a circa 1.1 nm effective interlayer thick-282 ness of a contamination layer consisting predominantly of water and traces of the other 283 contaminants. Based on this, using the semi-empirical calculations from Ziegler (12), we 284 can estimate the contribution of 1.1 nm of mainly water to the energy loss at 5 to be 35 eV. 285 This would reduce the energy loss protons in Figure 3 and 5 by 0.35 eV/A, which would still 286 result in a significantly higher energy loss than that of amorphous carbon.

Note also that F. Mao et al. (9), using TD-DFT, also predict high energy losses for He<sup>+</sup><sub>2</sub> inter-287 288 acting with graphene layers, reaching three times the results obtained by Echenique et al. 289 (23). For instance, for helium ions at 1 keV (0.1 a.u. of velocity), the energy loss in graphene 290 predicted for by F. Mao is around 11 eV/Å, meanwhile, Echenique gives 3.3 eV/Å. Also, 291 recent experiments of slow highly charged ions interacting with graphene layers report 292 that the energy loss and charge exchange of ions in two-dimensional materials show sig-293 nificant differences with respect to bulk solids (24,25). The high energy losses differ by an 294 order of magnitude with respect to results obtained by TRIM (12).

The higher energy loss for protons, found in our experiments and which recent calculations would fairly explain, can be traced to the unusual properties of graphene. Assuming that the measured energy loss is due to electronic excitations, and graphene with a high electron mobility can react very fast to the presence of the intruder, near the surface, with its subsequent fast screening (25). This phenomenon produces a very high and fast flux of electrons against the energetic proton producing a high momentum transfer which translates into a high graphene stopping power on the particle.

Another interesting feature appearing in our experiments is the apparent velocity 302 303 threshold in the protons energy loss, which appears around 0.1 a.u. of velocity (see Figure 5). 304 From the theoretical point of view, calculations of stopping power at low energies in the frame of the free electron gas model, using linear response theory, non-linear DFT and the 305 306 transport cross section model (22,26), predict that the stopping power is linear with the ion 307 velocity. However, experiments show that this prediction is not necessarily true for protons 308 and helium in metals and insulators. For instance, the predicted proportionality with the ion 309 velocity of the stopping power of transition metals (Cu, Ag and Au) for protons breaks down 310 drastically at some very low velocities displaying two well-differentiated regimes (27). This 311 phenomenon is explained considering the existence of a threshold effect for electron-hole 312 pair excitation, where the valence electrons in these materials, mainly non-free d electrons, 313 need a minimum of energy to be excited. Recently, this phenomenon of non-linearity with 314 the ion velocity has been studied with TD-DFT calculations (28), obtaining a more realistic 315 description of stopping power for H and He in metals like Au. This work finds very good 316 quantitative agreement with experiments, describing the deviation from the ion velocity 317 proportionality.

In the case of large band-gap insulators, such as LiF and KCl, a threshold effect was found for protons, deuterium and helium ions. Data obtained indicated a velocity threshold of around 0.1 a.u., below which particles pass through the material without energy loss, *i.e.* no electron excitations. For insulators, the electron excitations are suppressed due to a minimum excitation energy, the energy band gap of these materials plays a fundamental role. The interaction of low energy protons, in these cases, was described by invoking the creation of negative ions and charge interchange via electron promotion (29–33).

In the case of protons interacting with graphene at low energies, there may be sufficient
time for successive charge transfer, leading to negative ion formation as this occurs on, *e.g.*graphite (*34*), in which case it is a negative ion and not a fast proton passing through a high
electron density cloud, which could then affect the scattering process and energy loss.

This work deserves to be extended to the case of different numbers of graphene layers in the target. We consider this work as a starting point to study several low dimensional structures under particle irradiation and its effect on particles dynamics. These interesting results might be used in technological applications in nanostructures material characterization,

radiation protection, 'solar protons cells' in space energy storage.

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#### 338 339 **Disclosure statement**

Q12340 No potential conflict of interest was reported by the authors.

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