Graphene Enhanced Secondary Ion Mass Spectrometry (GESIMS)

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The following invention - Graphene Enhanced Secondary Ion Mass Spectrometry - (pending European patent application no. EP 16461554.4) is related to a method of analysing a solid substrate by means of Secondary Ion Mass Spectrometry (SIMS). It comprises the steps of providing a graphene layer over the substrate surface and analysing ejected secondary anions through mass spectrometry analysis. The graphene layer acts as a kind of filament that emits a lot of secondary electrons during the experiment which significantly increases the negative ionization probability and thus the intensity of the SIMS signal can be more than two orders of magnitude higher than that of a similar sample without graphene. The method is particularly useful for the analysis of surfaces, 2D materials and ultra-thin films. The intensity of dopants and contamination signals can be enhanced up to 35 times, which approaches the detection limit of $10^{15}$ atoms/cm$^3$, otherwise unreachable in a standard static SIMS analysis.

SIMS is a very precise analytical technique for determining the elemental composition of a sample$^{1-6}$. The sample is bombarded with a primary ion beam that leads to the sputtering of the surface. Small parts of the sputtered particles are ionized (secondary ions) and the sample composition can be determined by means of mass spectral analysis. During the sputtering subsequent layers of the sample are removed and thus it is possible to obtain information about changes in the composition as a function of depth, thus creating a depth profile.

SIMS is well known for its excellent detection limits of trace elements$^{7-12}$. For most materials it is reported to be in the range of $10^{15}$–$10^{16}$ atoms/cm$^3$11, sometimes even as good as $10^{12}$ atoms/cm$^3$32. These optimum detection limits, however, are achieved during the dynamic SIMS (dSIMS) mode. In this mode a very dense ion beam is used and a substantial amount of material is sputtered simultaneously, allowing more ions to be detected. Drawbacks include very poor depth resolution and limitation to thick materials.

Surfaces, 2D materials and ultra-thin films are analysed ideally in a special mode called static SIMS (sSIMS)$^{13-15}$, in which a low density ion beam ensures that ions are emitted only from monolayers 1 to 3. This mode, however, has physical limitations. Because a relatively small amount of matter is sputtered, and even less ionized, there are not enough secondary ions extracted from the sample to achieve as optimum detection limit as the dynamic mode does. In most cases it is difficult to exceed a limit of 1 ppm ($5 \times 10^{16}$ atoms/cm$^3$ or $10^9$ atoms/cm$^2$).

There are many ways to enhance the ionization probability and thus the detection limit. It has been noted that using oxygen as primary ions significantly enhances the formation of positive ions$^{16-19}$, whereas cesium enhances the formation of negative ions$^{17-20}$. For some elements the difference can be up to four orders of magnitude. Although most elements can be detected as both positive and negative ions, each element usually gives better results in terms of detection limit in either positive or negative mode. In some experiments, oxygen flooding can further enhance the secondary ion yields of some negative ions$^{21-23}$. Using these techniques yields acceptable results, but in the cases of surfaces, 2D materials and ultra-thin films, it is often not enough to reach the desired detection limit, particularly in the case of trace elements.

Several SIMS measurements on graphene (Gr) have already been reported$^{24-31}$. More specifically, SIMS has been used to map the surface of the graphene layer$^{24,25}$, to analyse the mass spectra of graphene$^{26,27}$, and to create a depth profile of single-layer graphene grown on substrates that do not contain carbon$^{26,28}$. Furthermore, SIMS has been used for the depth profiling of multilayer graphene of a thickness of more than 3 nm on SiC$^{30}$ and, recently, of hydrogen-intercalated graphene on SiC having a theoretical thickness of 0.69 nm$^{31}$. All these methods employed SIMS for characterising the graphene layer, while the composition of support material underneath was not analysed.

In this work we present Graphene Enhanced Secondary Ion Mass Spectrometry (GESIMS) - a new technique for enhancing the detection limit in the analysis of thin materials. It has been found that providing a graphene
layer on a substrate significantly reduces the emission of matter from the substrate but at the same time increases the negative ionization probability. Destruction of a small part of the graphene layer in a dynamic SIMS mode significantly reduces the initial strong blocking effect of graphene, while the enhancement of the ionization probability of the sputtered material is still observed. Under these conditions the ionization probability greatly prevails, resulting in an SIMS signal with an intensity that is unexpectedly high compared with a sample without graphene and thus able to to reach better detection limits.

Results
The basic concept of GESIMS method is presented on Fig. 1. To understand the process correctly it is necessary to remember that the intensity of SIMS signals depends on the secondary ion yield, which is defined as the number of emitted ions $A^{-}/A^{+}$ (multiple ionization is also possible) per incident ion. In turn, secondary ion yield is defined as the product of the partial sputter yield (number of emitted species A per incident ion) and ionization probability. The presence of graphene on a substrate surface clearly reduces the partial sputter yield of a material from the substrate, since for given conditions it is expected that sputtered species are emitted from the ~3 top monolayers of material and graphene acts as a barrier to emission. If despite this fact a significant increase in the SIMS signal intensity is still observed, the only possible explanation is that the presence of graphene drastically increases the negative ionization probability. Fig. 2 presents a comparison of mass spectra for the Ge substrate with and without graphene layer for optimal GESIMS conditions. The intensity of the SIMS signal can be enhanced more than two orders of magnitude.

Figure 1. The basic concept of GESIMS analysis. Graphene blocks the emission of the matter from the substrate but significantly increases the ionization probability. By creating some defects in graphene layer the emission increases while the ionization ability is preserved and thus more ions can reach the detector, resulting in enhanced SIMS signal.

Figure 2. A comparison of mass spectra of the germanium substrate with and without graphene layer for optimal GESIMS conditions. The intensity of the SIMS signal can be enhanced more than two orders of magnitude.
\[
\alpha = \frac{GF}{Y}
\]

wherein GF is the gain factor, \(Y_{Gr}\) and \(Y\) denote partial sputter yield for a sample with and without graphene, respectively, while \(\alpha_{Gr}\) and \(\alpha\) denote ionization probabilities (again, a sample with and without graphene, respectively). Under optimum conditions \(Y_{Gr} < Y\) and \(\alpha_{Gr} > \alpha\), whereby the latter effect is much stronger, so that \(GF > 1\) and thus the intensity of the SIMS signal can be enhanced.

Although increasing the intensity of the matrix element signal is of interest from a scientific viewpoint, it does not seem to have much practical application in SIMS-based surface analysis. As already mentioned above, the actual benefit is the enhancement of the detection limit of trace elements and thus new sets of samples have been considered and tested. Each of the sets consisted of a pair of samples, one having graphene transferred onto its surface, and the other - without graphene - serving as a reference. Signal enhancement was observed for all of them and the maximum gain that was reached during the experiments has been presented in Table 1 along with standard and enhanced detection limits.

As already mentioned, the effect was not observed at the beginning of the experiment, when a fresh graphene layer was transferred on a sample. This is not surprising since graphene is a strong material and the blocking effect was expected to be high. Thus the reduction in the partial sputter yield was higher than the increase of the ionization probability. To overcome it, the primary beam intensity had to be increased to several hundred pA for a short time. Under such conditions dynamic SIMS mode was achieved and the entire surface was slowly eroded. When only a small part of the graphene sheet was destroyed, the blocking effect was significantly reduced while the enhancement of the ionization probability was still observed. Figure 3 shows the gain change during sputtering in dSIMS mode with the primary beam intensity being set to 200 pA for all samples presented in Table 1. Three regions can be identified: at the beginning the blocking effect is very strong but after the partial destruction of graphene it is reduced and the enhanced ionization probability is dominant. Thus the gain is much larger than a factor of 1. After the complete destruction of the graphene layer, no difference between the two samples is observed (GF = 1). The shape of all curves and the position of the maximum depends on the partial destruction of the graphene layer and thus are similar for all samples.

| Substrate | Dopant | Concentration (atoms/cm\(^2\)) | Maximal gain factor | Standard detection limit (atoms/cm\(^2\)) | Enhanced detection limit (atoms/cm\(^2\)) |
|-----------|--------|-------------------------------|--------------------|------------------------------------------|------------------------------------------|
| GaAs      | Te     | \(1 \times 10^{18}\)         | 35.7               | \(7.04 \times 10^{14}\)                   | \(1.97 \times 10^{15}\)                   |
| Si        | Sb     | \(1 \times 10^{18}\)         | 15.1               | \(4.86 \times 10^{14}\)                   | \(3.02 \times 10^{15}\)                   |
| AlGaAs    | Si     | \(2 \times 10^{18}\)         | 13.8               | \(6.06 \times 10^{14}\)                   | \(4.39 \times 10^{15}\)                   |
| Si        | As     | \(5 \times 10^{18}\)         | 7.1                | \(1.35 \times 10^{14}\)                   | \(1.90 \times 10^{15}\)                   |

Table 1. Summary of Graphene Enhanced Secondary Ion Mass Spectrometry measurements performed on various materials and dopants.
It was essential in these experiments for the uniformity and the intensity of the beam to be well defined and reproducible and thus it was not surprising that the shape of all curves and the position of the maximum were similar for all measured samples as they depended on the partial destruction of the graphene layer and not to the type of the dopant and the substrate. Only the actual enhancement was material related (both the substrate and the type of dopant) and thus each sample had a different maximal gain factor.

The result is fully reproducible - Fig. 3 shows the average of 15 measurements performed separately on both types of sample (with and without graphene) for each sample set. We wish to emphasize that to achieve a good reproducibility of the method, samples must be cleaned prior to the measurement by annealing them at elevated temperatures and reduced pressure. In the experiment described above, the samples were subjected to annealing at 200 °C and pressure of $10^{-8}$ mbar for two hours. Figure 4 shows a suitable comparison of the gain factors measured for As dopant in Si substrate samples with and without cleaning procedures. The intensity of the primary beam was set to 150 pA and the duration of the dSIMS mode was varied between 2 and 4 minutes. Five different measurements were taken at various spots of the sample, leading to a straightforward conclusion that for the cleaned sample the method is reproducible, with only small variations.

The best results, as presented in Table 1, were obtained for higher values of primary ion energy (usually above 10 keV). This is not surprising because as more energy is transferred to the substrate, a larger collision cascade is generated and thus the matter from the substrate has higher energy to overcome the graphene blocking effect. Figure 5 shows a comparison of maximal gain factor versus primary ion energy for Te dopant in GaAs substrate. When the optimal conditions are reached, the beam intensity should be decreased and sSIMS measurement can be performed. Figure 6 presents a suitable comparison of SIMS measurement for Sb dopant in Si substrate with and without graphene. Two sample sets were analysed: with high ($1 * 10^{18}$ atoms/cm$^2$) and low

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**Figure 4.** The gain change during sputtering in dSIMS mode for As dopant in Si substrate measured at five different spots for a clean and a dirty sample. For the cleaned sample, the method is reproducible, with small variations only.

**Figure 5.** Maximal gain factor as a function of primary ion energy for Te dopant in GaAs substrate. For high-energy ions, a bigger collision cascade is generated and thus it is easier to overcome the graphene blocking effect.
(1 * 10^16 atoms/cm^3) dopant concentration (Sample A and B, respectively). As can be clearly seen for Sample A, the intensity of the SIMS signal was significantly increased and thus a better detection limit was achieved (3.02 * 10^15 instead of 4.86 * 10^16 atoms/cm^3). In the case of Sample B, the concentration was too low to be measured using a standard sSIMS condition, whereas it was possible for the GESIMS method. As can be seen, the enhancing effect does not depend on the dopant concentration and thus GESIMS method is quantifiable.

The same procedure was used for a O_2^+ primary beam with positive ion detection. As shown on Fig. 7 only the blocking effect of graphene was observed with no enhancement. Changing the impact energy influenced the sputter rate whereas the general shape of the profile was the same. It was therefore concluded that the positive ionization probability remained the same for the whole experiment and thus the Y_{Gr}/Y ratio is being monitored. For smaller impact energies lower sputter rate is observed.

A new experiment was devised for AlGaAs substrate with Si dopant: the sample was sputtered in dSIMS mode with Cs^+ ions and the gain factor change was obtained just like it was shown on Fig. 3. Every ten seconds the Cs^+ beam was switched off and a very brief sSIMS measurement with O_2^+ beam was performed so that the Y_{Gr}/Y ratio could be determined as shown on Fig. 8. We noted that the Y_{Gr}/Y ratio increased rapidly from ~0.27 to ~0.43 close to the optimal condition which apparently led to a significant increase of the gain factor.

It did not, however, explain why the graphene layer increase the negative ionization probability. To better understand the effect we repeated the whole procedure but instead of the graphene layer we deposited a thin layer of amorphous carbon. We tested it in both, negative and positive ion detection mode but no signal enhancement was observed, just the blocking effect and thus the enhancement could not be associated with the presence of carbon atoms only. To gain more insight we studied the electric properties of the graphene layer and amorphous carbon during the SIMS experiment: we transfered/deposited them on insulating materials like SiC, AlN and SiO_2 which normally required an electron flood gun for charge compensation. The intensity of the Cs^+ primary beam was once again set to 200 pA and it turned out that a thin layer of the amorphous carbon did not alter the...
experiment significantly; the charging effect was visible from the very beginning and the flood gun was needed to perform the analysis. However, in case of the graphene layer we were able to sputter the sample without the flood gun up to four and a half minutes before the charging effect occurred. It corresponded well with the time frame of the enhancement effect (see Fig. 3).

From Fig. 8 it could be further deduced that after four and half minutes of sputtering the graphene layer was significantly destroyed (the \( Y_{Gr}/Y \) ratio was above 0.8) and yet it was still able to provide enough electrons for charge compensation. We assumed that in a similar way the GESIMS effect could be explained: during the SIMS experiment a high voltage was applied to the sample holder (up to 5 kV) and thus the graphene layer acted as a kind of filament which emitted an excess of electrons during the ion bombardment leading to the enhanced ionization probability.

Discussion
GESIMS is a process for measuring and analysing thin substrates that involves the following steps:

- applying a graphene layer over the substrate surface;
- annealing the graphene-coated substrate at elevated temperature and reduced pressure;
- sputtering of the graphene-coated substrate in dSIMS mode, which leads to the partial destruction of the graphene layer;
- detecting and analysing ejected secondary anions by mass spectrometry analysis in sSIMS mode.

This method cannot be used for depth profiling of thick materials because the graphene layer will be destroyed within the first moments of the measurement and the enhancing property will be lost. Furthermore, the dSIMS analysis itself offers a high detection limit. It cannot be used for a high resolution imagining neither, since the presence of the partially destroyed graphene layer will reduce the spatial resolution. However, in the case of very thin materials, sSIMS reaches its physical limit and any enhancement is beneficial. GESIMS requires more preparation and optimization than the standard sSIMS analysis but improves by one order of magnitude the detection limits of trace elements, which may prove to be invaluable for the further development of 2D materials.

The effect does not take place for samples covered with a thin layer of amorphous carbon. It has been noted that the graphene layer has different electric properties during the SIMS experiment - even a significantly destroyed layer can provide enough electrons to compensate the charging effect which is typically present for insulating materials. This behaviour can suggest the explanation of the GESIMS effect: since a high voltage is applied to the sample holder the graphene layer acts as a kind of filament and emits a lot of secondary electrons during the ion bombardment and thus the negative ionization probability is increased.

Methods
Sample preparation. A pair of samples was created for every experiment described in this work, one having graphene grown or transferred onto its surface, and the other - without graphene - serving as a reference.

In the case of germanium substrate, graphene films were synthesized in a commercially available system by the CVD method as described by Paternak et al. As a substrate, (100)-oriented Ge layers deposited on Si(100) wafers were used. Methane gas in the mixture of Ar and \( H_2 \) in the ratio of 200:1 was used as a carbon precursor. Growth was preceded by the substrate's annealing in a pure hydrogen atmosphere in order to reduce native oxides \textit{in-situ}. During the growth process, 800 mbar of pressure were sustained.

In other samples, graphene films were synthesized on 35 \( \mu m \) thick copper foils by the CVD method using a Black Magic Pro system (Aixtron). To grow graphene on copper foils the following procedure was used: copper samples were first pretreated at 960 °C under an Ar gas flow and then a \( H_2 \) gas flow at 20 mbar of pressure. The purpose of this step was to improve the quality and enlarge the grain size of Cu substrates as well as to remove oxides from the copper surface. Next, methane was introduced into the reactor at an adequate flow rate for a few
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Author Contributions
P.P.M. conceived the experiment, P.P.M. and W.K. conducted the SIMS experiments, I.P. and W.S. prepared the samples. All authors reviewed the manuscript.

Additional Information
Competing Interests: The authors declare that they have no competing interests.

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