Tip-enhanced Raman mapping (TERM) of single-walled carbon nanotubes and graphene

Günter G. Hoffmann, a,∗ Marcos Ghislandia,b and Gijsbertus de With a

aDepartment of Chemical Engineering and Chemistry, Laboratory of Materials and Interface Chemistry (SMG), Eindhoven University of Technology, PO Box 513, 5600 MB Eindhoven, The Netherlands. E-mail: g.g.hoffmann@tue.nl
bCETENE–Center for Strategic Technologies of the Northeast, Av. Prof. Luiz Freire 1 (CDU), Recife–PE, 50740-540, Brazil

Introduction
Single-walled carbon nanotubes (SWCNTs)1 and graphene2 have received tremendous research attention during the last few years. Both allotropes of carbon have matchless mechanical and electrical properties. The extraordinary tensile strength of SWCNTs ranges from 13 GPa to 126 GPa (much higher than of stainless steel with 0.38 GPa to 1.55 GPa and of Kevlar® with 3.6 GPa to 3.8 GPa); the strength of graphene is also about 100 times that of steel. This makes these materials very attractive to be used as fillers in polymer composites and thereby increase the polymer’s strength and electrical conductivity. The excellent electrical conductivity of graphene makes a number of enhanced applications in electronics possible, for example, touch screens, field effect transistors and a large range of novel sensors of many kinds. Metallic SWCNTs can, theoretically, carry a current of 4 × 109 A cm–2, which means that they conduct current about a thousand times better than copper. SWCNTs have a high thermal conductivity value of 3500 W m–1 K–1 along the axis of the tube, compared to 385 W m–1 K–1 of copper. The thermal conductivity of graphene is even more impressive, with a measured value of 5000 W m–1 K–1.

A SWCNT can be considered as a narrow sheet of graphene that has been rolled and fused. This is also apparent from the Raman spectrum: most features can be found in the spectra of both allotropes. The G-line, attributed to C–C stretching vibrations and named after its occurrence in graphite at 1582 cm–1, is split into two components in CNTs: the G’-line at 1590 cm–1, which is attributed to the stretching along the axis of the nanotube (longitudinal phonon mode) and the G’-line at 1570 cm–1, which arises from the transversal mode of the C–C stretching. The D-line at ~1350 cm–1 is only found if there are defects in the sample, so it is used to judge the quality of the material. In graphene it can also be used to detect the edges of the sheets, as these are considered “defects” in an infinite graphene sheet. The G’- or 2D-band at 2700 cm–1 is the first overtone of the D-band and can be used to analyse the number of graphene layers in a graphitic sample using its Raman shift and the 2D-/G-band intensity ratio. The radial breathing mode (RBM, at 100 cm–1 to 500 cm–1), which is a C–C bond-stretching mode that moves all carbon atoms radially out, can, of course, only be observed from the tubes. As its wave-number is inversely proportional to the tube’s diameter, it can be advantageously used to analyse a sample’s content of tubes with different diameters.

Tip-enhanced Raman spectroscopy (TERS) Raman spectroscopy and infrared spectroscopy are complementary techniques for the qualitative and quantitative analysis of materials which, in addition to chemical composition, yield a lot of useful information including crystallinity and other properties of the sample under test. In the form of confocal Raman spectroscopy and infrared microscopy, they can also be used to spatially map the sample; the former being the higher spatial resolution technique. Due to physical limitations, the resolution is limited to 0.2–1 µm, depending upon the wavelength of the Raman spectroscopy excitation laser used. Using a “trick”, this limit may be overcome: in the case of Raman spectroscopy, a noble metal tip is placed at the focus of a confocal Raman spectrometer. The resulting technique is therefore called tip-enhanced Raman spectroscopy (TERS) in the case of a stationary sample, or tip-enhanced Raman mapping (TERM), for the case of a scanned sample. Using TERM, carbon allotrope samples can be analysed with high spatial resolution.

The basic TERS experimental set-up for transparent samples is illustrated in Figure 1. Using a high numerical aperture (NA) microscope objective (of, for example, an inverted microscope), a laser beam is focused onto a sample. The sample can be moved in both the x- and y-directions by a high precision piezo sample stage. A very sharp needle (radius at tip ~30 nm), etched electrochemically from a thin gold wire, is attached to a miniature quartz tuning fork. By means of a second x–y-scanner in an atomic force microscope (AFM) shear force head, the gold tip is placed with high accuracy at the exact centre of the laser focus. Feedback from the tuning fork, vibrating at its resonance frequency, is used to keep the gold tip at a constant distance to the sample with a piezo fork.
The distance of the tip is kept constant by a feedback loop to a piezo and the mapping is done by a high precision xy-stage. The enhanced Raman spectrum is collected by the objective and fed into the spectrometer.

**TERM of graphene**

Figure 3 shows the experiments on a graphene sample which was grown by chemical vapour deposition (CVD) on nickel. After etching away the substrate, the sample was transferred to a thin microscope cover glass, where a confocal Raman map (a) was recorded using the shift position of the 2D line for mapping. The graphene area, surrounded by multiple layer material, could be identified as a flat area in the atomic force microscope’s phase image (b). A higher resolution Raman map could be obtained using TERS. Using the intensity of the 2D Raman band, an image (c) of the clearly defined graphene area could be obtained. If, in the TERS experiment, the shift position is again used for the mapping (d), the image shows more details: the graphene area is surrounded by a borderline. We can now pick special places from 1 to 4, where different spectra can be measured.

Figure 4 shows the Raman spectra that were taken at the positions shown in part (d) of Figure 3. At point 4 (blue curve) we measure a pure graphene spectrum,
easily recognised by the 2D-band position and the low G- to 2D-band intensity ratio. Point 3 (green curve) corresponds to the edge of the graphene area: a D-peak appears. Moving further to point 2 (yellow curve), more than one graphene layer can be detected from the breadth and the position of the 2D-band also the D-band increases. Finally, a graphite spectrum with high G- to 2D-band intensity ratio was acquired at point 1 (red curve).

Conclusion
Tip-enhanced Raman spectroscopy (TERS) and tip-enhanced Raman mapping (TERM) can be used advantageously to investigate the carbon allotropes graphene and single-walled carbon nanotubes, with a spatial resolution in the nanometre range. Even the localisation of defects in one single sheet or tube is possible, with a spatial resolution of ~35 nm.

Acknowledgements
This work is part of the Research Program of the Dutch Polymer Institute (DPI), Eindhoven, The Netherlands, projects #648, #692 and #756.

References