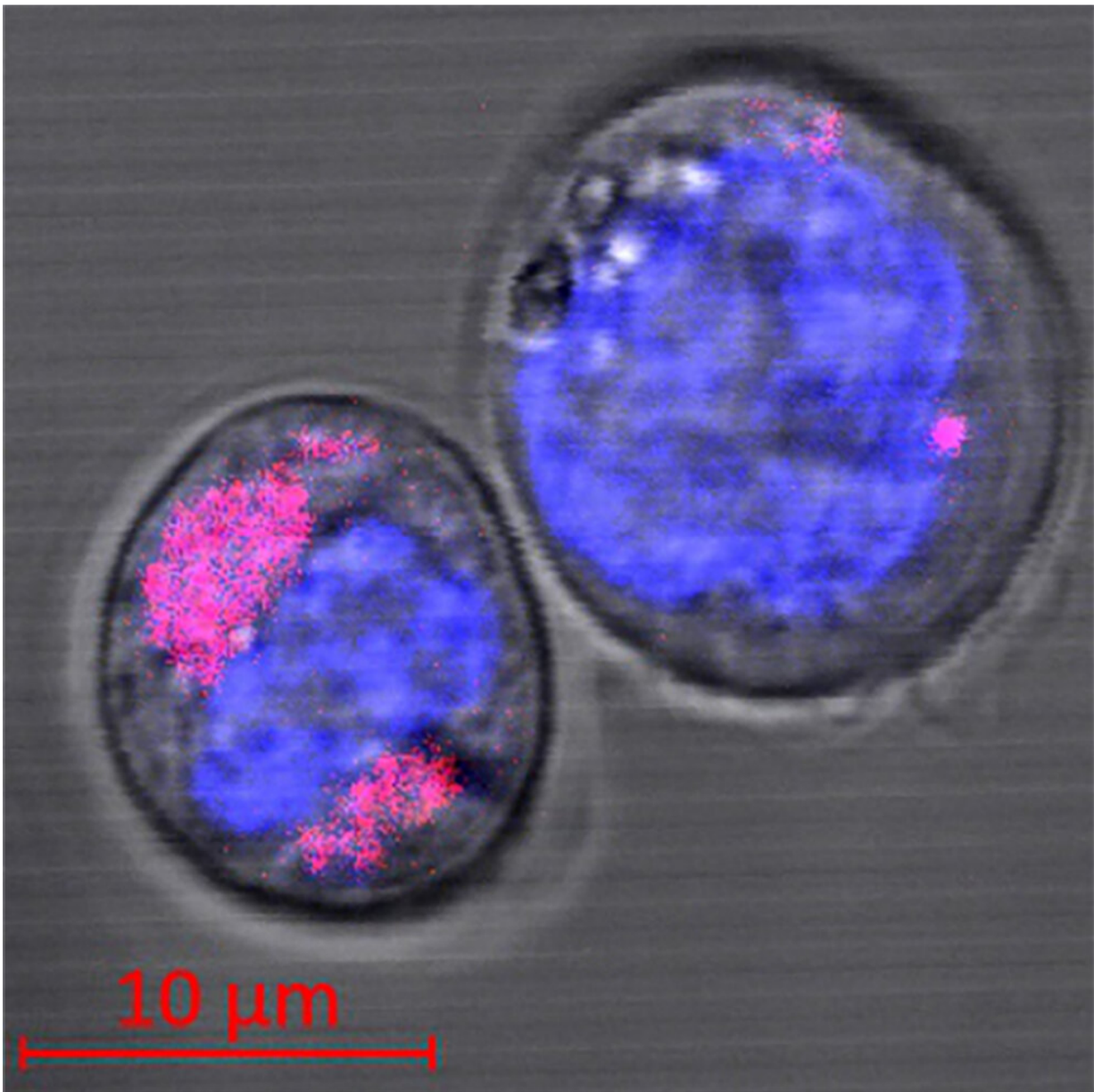


Biological risk potential of nanoparticles studied

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Two CD34+ stem cells containing carbon nanoparticles (coloured magenta); the cell nuclei can be seen in blue. The researchers found that the nanoparticles are encapsulated in the cell lysosomes. Credit: HHU / Stefan Fasbender

Carbon nanoparticles are a promising tool for biomedical applications, for example, for targeted transportation of biologically active compounds into cells. A team of researchers from the Physics, Medicine and Chemistry departments at Heinrich Heine University Düsseldorf (HHU) has now examined whether these particles are potentially dangerous for the organism and how cells cope with them once they have been incorporated. The findings of the interdisciplinary study have just been published in the journal *Scientific Reports*.

Nanoparticles are smaller than five nanometres—a nanometre being one millionth of a millimetre—which corresponds approximately to the size of macromolecules. Such tiny particles are very easily absorbed in body [cells](#). There are two aspects to this feature. Firstly, it makes nanoparticles good vehicles for transporting a broad range of compounds or substances attached to them into normal diseased cells in a targeted manner.

On the other hand, they can also pose [health risks](#), for example in connection with [particulate matter](#). One of the ways that particulate matter is created is in combustion processes, and part of that can be classified as nanoparticles. These extremely small particles can overcome the blood-air barrier and penetrate the body: The bronchial mucosa in the lungs doesn't filter out the particles. Instead, they make their way into the pulmonary alveoli and from there into the bloodstream.

Together with work groups from the Chemistry department, HHU

researchers from the Institute of Experimental Condensed Matter Physics working under Prof. Dr. Thomas Heinzl and from the Department of Haematology, Oncology and Clinical Immunology working under Prof. Dr. Rainer Haas have now studied what happens when body cells absorb such nanoparticles. The researchers used nanoparticles made from graphene; this is a special form of carbon that comprises two-dimensional layers of hexagonal carbon rings. They added these to special haematopoietic stem cells referred to as CD34+ stem cells. These cells are particularly sensitive to damaging environmental influences on account of their ability to divide throughout their lifespan. The assumption is that these cells would be damaged more by nanoparticles—if at all—than the more robust other cell types.

The interdisciplinary team of researchers based in Düsseldorf was able to demonstrate that the [carbon nanoparticles](#) get into the cells, where they are encapsulated in special organelles called lysosomes. The lysosomes serve as a type of waste removal unit for the body where foreign bodies accumulate and are normally broken down with the help of enzymes. However, the researchers didn't observe any such process over the duration of the experiments, which lasted for several days.

When comparing the active genes ("gene expression") of stem cells with and without the addition of nanoparticles, the researchers found that only one of a total of 20,800 recorded expressions had changed; minor effects were determined in a further 1,171 gene expressions.

Prof. Heinzl had this to say regarding the findings: "Encapsulation of the nanoparticles in the [lysosomes](#) ensures that these particles are stored securely at least for a few days—for the duration of our experiments—and cannot damage the cell. This means the cell remains viable without any major change in gene expression." This insight is important if [nanoparticles](#) are to be used to deliver drugs into the cell. The experimental framework used here does not allow for any long-term

statements to be made regarding any increased probability of cell mutation resulting in cancer.

The research was carried out as a close collaboration between HHU's Faculty of Mathematics and Natural Sciences and Faculty of Medicine and University Hospital Düsseldorf. Düsseldorf School of Oncology (headed up by Prof. Dr. Sebastian Wesselborg) funded the doctoral scholarship of first author Stefan Fasbender. Prof. Haas said: "The proximity of the Hospital and the University and their close links in terms of content provides HHU with a particularly fruitful environment for translational research, where insights and expertise from basic research are combined with aspects relevant to treatment."

More information: Stefan Fasbender et al, The Low Toxicity of Graphene Quantum Dots is Reflected by Marginal Gene Expression Changes of Primary Human Hematopoietic Stem Cells, *Scientific Reports* (2019). [DOI: 10.1038/s41598-019-48567-6](https://doi.org/10.1038/s41598-019-48567-6)

Provided by Heinrich-Heine University Duesseldorf

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