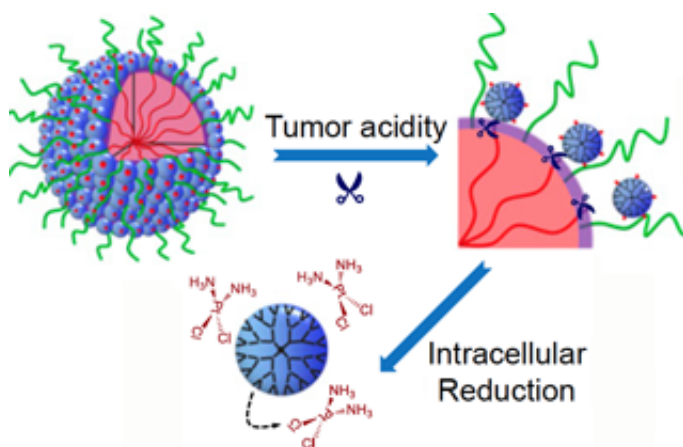


Nanoparticles deliver anticancer cluster bombs

March 29 2016



The particles start out large (100 nm) to enable smooth transport into the tumor through leaky blood vessels. Then, in acidic conditions found close to tumors, the particles discharge bomblets 5 nm in size. Inside tumor cells, a second chemical step activates cisplatin. Credit: Jinzhi Du/Wallace H. Coulter Department of Biomedical Engineering at Georgia Tech and Emory

Scientists have devised a triple-stage "cluster bomb" system for delivering the chemotherapy drug cisplatin, via tiny nanoparticles designed to break up when they reach a tumor.

Details of the particles' design and their potency against cancer in mice were published March 28 in *PNAS*. They have not been tested in humans, although similar ways of packaging cisplatin have been in clinical trials.

What makes these particles distinctive is that they start out relatively large—100 nanometers wide - to enable smooth transport into the tumor through leaky blood vessels. Then, in acidic conditions found close to tumors, the particles discharge "bomblets" just 5 nanometers in size.

Inside tumor cells, a second chemical step activates the platinum-based cisplatin, which kills by crosslinking and damaging DNA. Doctors have used cisplatin to fight several types of cancer for decades, but toxic side effects - to the kidneys, nerves and inner ear—can limit its effectiveness.

The *PNAS* paper is the result of a collaboration between and a team led by professor Jun Wang, PhD at the University of Science and Technology of China, and researchers led by professor Shuming Nie, PhD in the Wallace H. Coulter Department of Biomedical Engineering at Georgia Tech and Emory. Nie is a member of the Discovery and Developmental Therapeutics research program at Winship Cancer Institute of Emory University. The lead authors are graduate student Hong-Jun Li and postdoctoral fellows Jinzhi Du, PhD and Xiao-Jiao Du, PhD.

"The negative side effects of cisplatin are a long-standing limitation for conventional chemotherapy," says Jinzhi Du. "In our study, the delivery system was able to improve tumor penetration to reach more cancer cells, as well as release the drugs specifically inside cancer cells through their size-transition property."

The researchers showed that their nanoparticles could enhance cisplatin drug accumulation in [tumor tissues](#). When mice bearing human pancreatic tumors were given the same doses of free cisplatin or cisplatin clothed in pH-sensitive nanoparticles, the level of platinum in tumor tissues was seven times higher with the nanoparticles. This suggests the possibility that nanoparticle delivery of a limited dose of cisplatin could restrain the toxic [side effects](#) during cancer treatment.

The researchers also showed that the nanoparticles were effective against a cisplatin-resistant lung [cancer model](#) and an invasive metastatic breast cancer model in mice. In the lung cancer model, a dose of free cisplatin yielded just 10 percent growth inhibition, while the same dose clothed in nanoparticles yielded 95 percent growth inhibition, the researchers report.

In the [metastatic breast cancer](#) model, treating mice with cisplatin clothed in nanoparticles prolonged animal survival by weeks; 50 percent of the mice were surviving at 54 days with nanoparticles compared with 37 days for the same dose of free cisplatin. Enhanced efficacy in three different [tumor](#) models demonstrate that this strategy may be applicable to several types of cancer, Jinzhi Du says.

More information: Hong-Jun Li et al. Stimuli-responsive clustered nanoparticles for improved tumor penetration and therapeutic efficacy, *Proceedings of the National Academy of Sciences* (2016). [DOI: 10.1073/pnas.1522080113](#)

Provided by Emory University

Citation: Nanoparticles deliver anticancer cluster bombs (2016, March 29) retrieved 5 May 2024 from <https://phys.org/news/2016-03-nanoparticles-anticancer-cluster.html>

This document is subject to copyright. Apart from any fair dealing for the purpose of private study or research, no part may be reproduced without the written permission. The content is provided for information purposes only.
