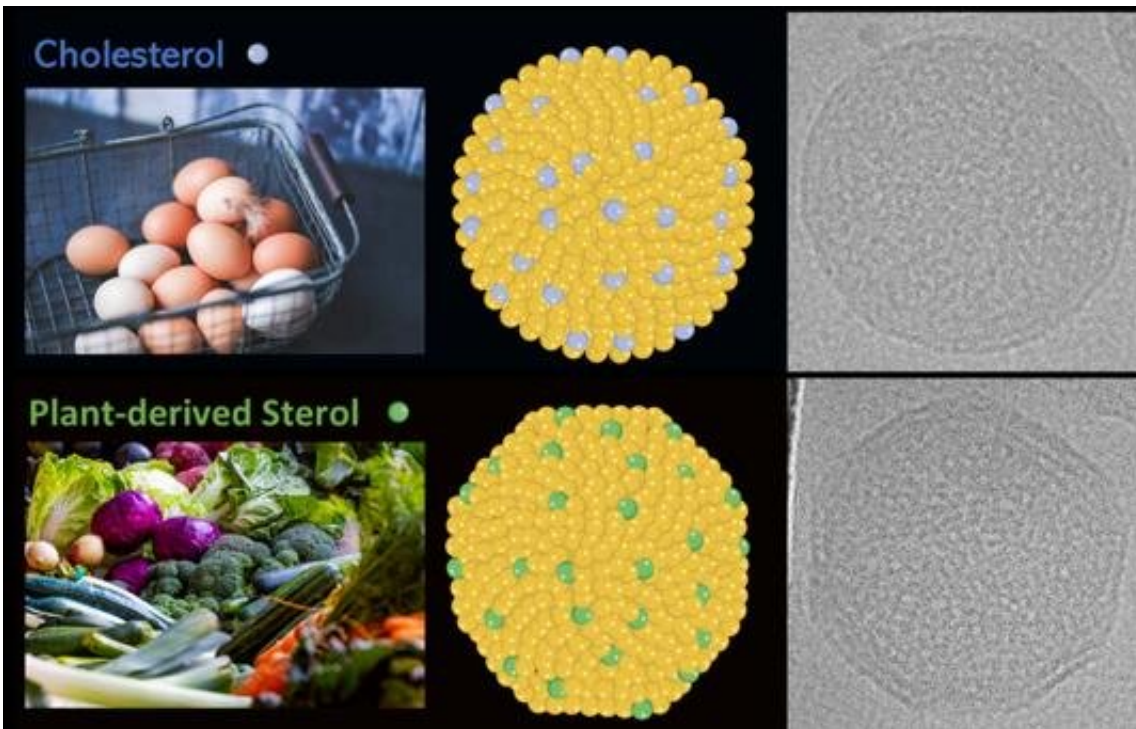


Plant-based relatives of cholesterol could give boost to gene therapy

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Gene-infused nanoparticles used for combating disease work better when they include plant-based relatives of cholesterol because their shape and structure help the genes get where they need to be inside cells. Credit: Gaurav Sahay, OSU College of Pharmacy.

Gene-infused nanoparticles used for combating disease work better when they include plant-based relatives of cholesterol because their shape and structure help the genes get where they need to be inside cells.

The findings by Oregon State University researchers, published today in *Nature Communications*, are important because many illnesses that can't be treated effectively with conventional drugs can be treated genetically—delivering nucleic acids to diseased [cells](#) so they can make the correct proteins needed for health.

Those genetic treatments rely on the transport devices reaching their destination with a high success rate and releasing their cargo effectively.

Gaurav Sahay, assistant professor of pharmaceutical sciences in the OSU College of Pharmacy, studies lipid-based nanoparticles as a gene delivery vehicle, with a focus on cystic fibrosis.

Cystic fibrosis is a progressive genetic disorder that results in persistent lung infection and afflicts 30,000 people in the U.S., with about 1,000 new cases diagnosed every year.

More than three-quarters of patients are diagnosed by age 2, and despite steady advances in alleviating complications, the median life expectancy of cystic fibrosis patients is still just 40 years.

One faulty gene—the cystic fibrosis transmembrane conductance regulator, or CFTR—causes the disease, which is characterized by lung dehydration and mucous buildup that blocks the airway.

Two years ago, Sahay and other scientists and clinicians at OSU and Oregon Health & Science University demonstrated proof-of-concept for a new, improved cystic fibrosis therapy: loading chemically modified CFTR messenger RNA into lipid-based nanoparticles, creating molecular medicine that could simply be inhaled at home.

The mRNA-loaded nanoparticle approach causes cells to make the correct protein, allowing cells to properly regulate chloride and water

transport, which is critical to healthy respiratory function.

Cholesterol, a waxy substance the body uses to make healthy cells, is thought to provide stability in these gene nanocarriers. In the latest study, Sahay and collaborators boosted gene delivery by using plant-based analogs of cholesterol instead. Another plus of these plant-derived sterols is a cardiovascular health benefit, he adds.

The type of nanoparticle used to deliver genes in this study has already been clinically approved; it's being used in a drug, trade-named Onpattro, given to patients with a progressive genetic condition called amyloidosis, which disrupts organ function through harmful deposits of the amyloid protein.

Sahay and graduate student Siddharth Patel, first author on the study, found that phytosterols—plant-based molecules chemically similar to cholesterol—change the shape of the nanoparticles from spherical to polyhedral and cause them move faster.

That's important because once inside a cell, the nanoparticles need maneuverability for the escape they need to make: from a cell compartment known as an endosome into the cytosol, where the delivered genes can perform their intended function.

"One of the biggest challenges in the delivery of genes is that less than 2% of the nanoparticles reach the cytosol," said Sahay, who also holds an adjunct faculty position with OHSU. "If you up the dose to get more genes there, now you have problems with toxicity, plus the cost goes higher. But the nanoparticles' [shape changes](#) because of these naturally occurring cholesterol analogs, and the new shape helps them deliver [genes](#) better. The analogs boost gene delivery 10-fold and sometimes 200-fold."

The finding can be used to make inhalable particles that can cross several barriers in the lung in a [cystic fibrosis](#) patient, enabling patients to be treated with much higher efficacy, Sahay added.

"In this latest research, we hypothesized that with the analog inclusions, there would be shape changes and changes with how the nanoparticles interact with the cell and how the cell perceives them," Patel said. "For instance, the sterols might help them get to the ribosomes for translation faster. This opens up a whole new area of research—the shape and structure and composition of the liquid [nanoparticles](#) now become quite relevant. We're just scratching the surface on the way to building LNPs with a rational design to get different properties for treating different diseases with cell-type specificity."

More information: *Nature Communications* (2020). [DOI: 10.1038/s41467-020-14527-2](#)

Provided by Oregon State University

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