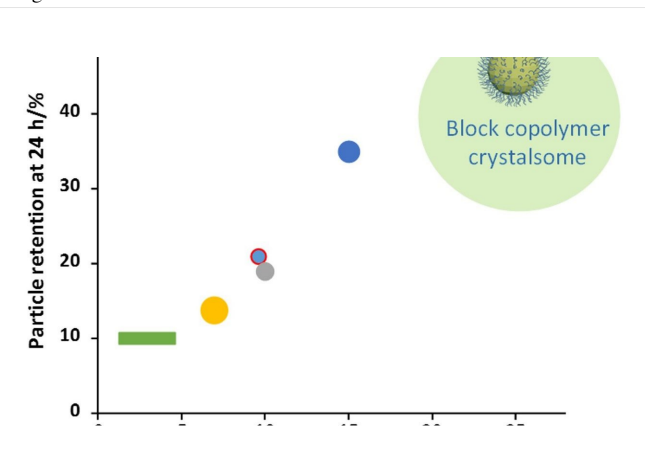


Tests show 'crystalsome' nanoparticle lasts longer in bloodstream



Polymer crystalsome nanoparticles have a 24-hour half-life and can last in the bloodstream for more than 96 hours, figures that far exceed current injectable medication. Credit: Drexel University

Selecting the right packaging to get precious cargo from point A to point B can be a daunting task at the post office. For some time, scientists have wrestled with a similar set of questions when packaging medicine

August 1 2018



for delivery in the bloodstream: How much packing will keep it safe? Is it the right packing material? Is it too big? Is it too heavy? Researchers from Drexel University have developed a new type of container that seems to be the perfect fit for making the delivery.

Intravenous medication has taken important leaps in recent years as a way of directly targeting ailments where they are occurring inside the body. But getting the medicine through the bloodstream to the right place and releasing it at the right time is no easy task. The body is designed to detect and eliminate foreign objects, so successfully designing a vessel for targeted <u>drug delivery</u> requires equal parts engineering and cunning.

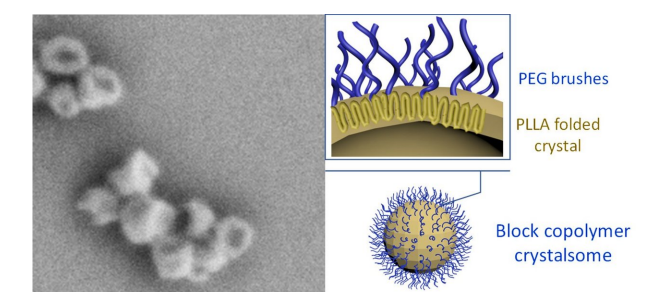
"Delivery vessels have traditionally been designed to avoid recognition by the immune system by mimicking naturally occurring materials in the body, such as cells or liposomes," said Christopher Li, Ph.D., a materials science professor in Drexel's College of Engineering. "But the problem with the previously reported artificial carriers is that they're not always durable enough to get to the far reaches of the body."

Li and Hao Cheng, Ph.D., an assistant professor in the College of Engineering lead a group of researchers who have been developing a polymer crystal casing for intravenous medicine delivery. Their work, which was recently published in the journal *Nature Communications* shows how these "crystalsomes," designed to be durable enough for long, intravenous journeys, can outlast current artificial nanoparticle packaging—which means doctors can use it to directly treat maladies in the body, with precisely the right amount of medication.

"Crystalsomes structurally mimic the classical liposome and polymersomes used for drug delivery, yet mechanically they are more robust thanks to their single crystal-like shell," Li said.



In blood circulation and biodistribution experiments, Li's polymer crystalsomes have a 24-hour half-life and can last in the bloodstream for more than 96 hours—figures that far exceed current injectable medication.



Researchers at Drexel University designed a polymer nanoparticle, called a "crystalsome," whose unique combination of polymer structures makes it exceptionally durable in the bloodstream, and an excellent candidate for intravenous drug delivery. Credit: Drexel University

"Crystalsomes are closely sealed so that medication will not be released until it reaches the target sites. Thus medication can be delivered in higher doses, as desired, to afflictions in the body, without causing severe side-effects associated with the early release of the medicine," Li said. "And a more direct intravenous delivery means that treatments are likely to be more effective."

Li's group combined its unique work on growing crystal spheres and self-



assembled nanobrushes to produce this special capsule that is just thick enough to safely encase the medicine, and also features an array of <u>polymer strands</u> that can ward off the proteins that flag foreign bodies for removal.

The method for creating the crystalsomes, which Li's Soft Materials Lab initially developed in 2016, looks something like combining oil and water to create suspended liquid beads. In this application, the beads encapsulate two types of polymer strands that, when cooled, condense into the solid, egg shell-like spherical crystalsome, protecting the yolklike cargo inside.

While one set of polymers, called poly L-lactide acid or PLLA, are drawing together to form the corrugated casing of the sphere, the other variety, poly ethylene glycol or PEG, coming to attention like whiskers on its surface. PEG polymers are known to prevent proteins from attaching to solid surfaces, so the uniform distribution of these polymers on the outside of the crystalsome prevent it from being flagged by the proteins of the immune system as a bodily invader.

"Taken together, these characteristics give the crystalsome its superior staying power in the bloodstream," said Cheng, whose research group specializes in <u>engineering molecules for intravenous drug delivery</u>.

The discovery provides a strategy for producing long-circulating nanomaterials, which could lead to a new class of polymer nanoparticle carriers for drug delivery and gene therapy, according to the researchers.

"The ingenious, curved, polymer crystal nanocapsules reported here remain robust while circulating in the blood, a potentially important feature for delivering medications and gene therapies," said Andrew Lovinger, the materials research program officer who oversaw National Science Foundation funding of the work. "NSF is proud to have



supported this important research, which integrates the agency's missions to promote the progress of science as well as contribute to advancing the nation's health."

More information: Hao Qi et al, Block copolymer crystalsomes with an ultrathin shell to extend blood circulation time, *Nature Communications* (2018). DOI: 10.1038/s41467-018-05396-x

Provided by Drexel University

Citation: Tests show 'crystalsome' nanoparticle lasts longer in bloodstream (2018, August 1) retrieved 3 May 2024 from <u>https://phys.org/news/2018-08-crystalsome-nanoparticle-longer-bloodstream.html</u>

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