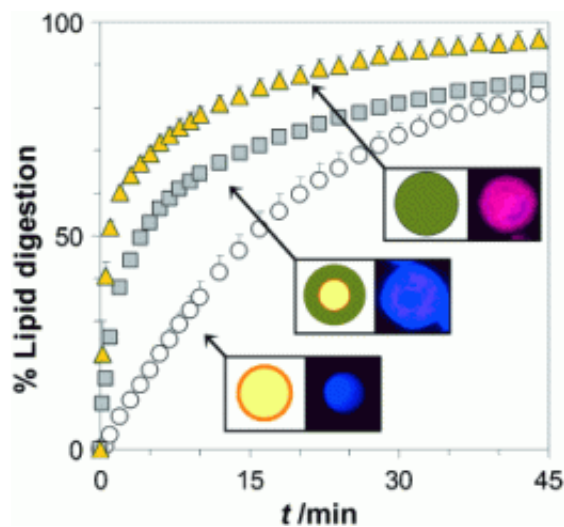


Simulated digestion: Nanomaterial made from lipids and silicon dioxide improves absorption of pharmaceuticals

April 27 2012



(Phys.org) -- Some medicines have to be taken either before, after, or during a meal because food ingredients can affect its absorption or bioavailability. Australian researchers have now encapsulated drugs in a matrix of silicon dioxide and lipids to simulate the administration of pharmaceuticals with food. As the researchers report in the journal *Angewandte Chemie*, drug absorption is increased through control of the enzymatic digestion of the lipid droplets.

For example, the body only absorbs fat-soluble vitamins A and D in the presence of some fat. Also, the bioavailability of many poorly water-soluble drugs increases when they are taken with high-fat meals. There are many techniques for the lipid-based delivery of pharmaceuticals, including emulsions, micelles and “packaging” in liposomes. These methods prevent the active ingredients from precipitating out and improve transportation to the absorption sites in the gastrointestinal tract. However, in order for the [drug](#) to become active in the body, it must be released from its [lipid](#) shell. The enzymatic decomposition of the lipid coating plays an important role in this process, but it has proven to be difficult to control. In addition, it is difficult to calculate the extent to which such lipid “packaging” really increases the bioavailability of a drug.

Clive Prestidge and a team at the University of South Australia and Monash University have now developed a controllable packaging type. Their material consists of a nanostructured network of [silicon dioxide](#) nanoparticles that contains nanoscopic [lipid droplets](#) containing the drug. This system is produced by generating a fine emulsion of the drug-containing oil droplets in an aqueous phase. The silicon dioxide particles collect around the droplets at the phase boundary. Spray-drying results in solid microparticles of entrapped lipid droplets.

The team has demonstrated that the lipid in these microparticles is enzymatically digested much more rapidly than pure lipid drops. This is because the nanostructured silicon dioxide network holds the enzymes close to their substrate. The size of the silicon dioxide particles used and the porosity of the resulting [matrix](#) determine how fast the enzymatic decomposition of the lipids occurs.

Animal trials with Celecoxib, a drug used to treat arthritis, showed a higher drug content in plasma when the pharmaceutical was orally administered in this new form rather than in its pure state or as drug-

containing lipid drops. In contrast to Celecoxib-containing lipid drops, the release rate did not vary from batch to batch or after a longer storage period.

The new nanomaterial imitates the food effect in a predictable fashion and allows for better control of drug release, it could minimise the food effect on drug absorption and enhance more predictable therapeutic responses.

More information: Clive A. Prestidge, Hybrid Nanomaterials that Mimic the Food Effect: Controlling Enzymatic Digestion for Enhanced Oral Drug Absorption, *Angewandte Chemie International Edition*, [dx.doi.org/10.1002/anie.201200409](https://doi.org/10.1002/anie.201200409)

Provided by Wiley

Citation: Simulated digestion: Nanomaterial made from lipids and silicon dioxide improves absorption of pharmaceuticals (2012, April 27) retrieved 18 April 2024 from <https://phys.org/news/2012-04-simulated-digestion-nanomaterial-lipids-silicon.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.