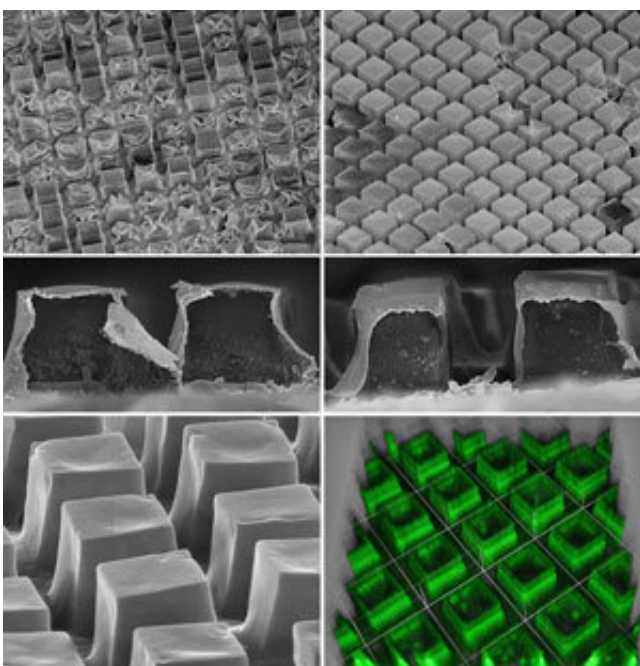


Layered chambers open a window for drug release

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The top and middle rows show microchambers without (left) and with (right) incorporated graphene oxide after dissolution of the templates. The bottom row shows microchambers with graphene oxide after peeling away the template, both scanning electron microscopy (left) and confocal laser scanning microscopy (right) images. Credit: Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim

Implantable arrays of microchambers show potential capacity for holding and releasing precisely controlled quantities of drugs on command, report A*STAR researchers with colleagues in Singapore, Russia and the United Kingdom.

A near-infrared laser beam acts to break open selected microchambers at the required time. "This [near-infrared light](#) is the perfect way to trigger drug release as it has the maximum penetration into [biological tissues](#)," says Maxim Kiryukhin of the A*STAR Institute of Materials Research and Engineering. The required wavelengths fall within the 'therapeutic window' that allows light for medical uses to reach safely into the body.

The team make the microchambers from composites of polymers and [graphene oxide](#). "My research group pioneered the manufacture of microchamber arrays using techniques called [nanoimprint lithography](#) and layer-by-layer assembly," says Kiryukhin. The lithography step makes templates with a desired pattern of microwells imprinted into their surface. Layers of polymers and graphene oxide are then built up on the templates to make a composite material. The templates can be dissolved or peeled away, creating the polymer/graphene-oxide chambered arrays that can be sealed with a layer of plastic.

If they are to contain drugs for delivery into the body, the chambers need to be mechanically robust. "Failure, followed by sudden release of the entire [drug](#) payload, could be catastrophic," Kiryukhin points out. Incorporating graphene oxide layers into the polymer layers is the critical innovation that makes the chambers sufficiently stable and responsive to near-infrared light.

The researchers have already developed techniques that can be used to load the chambers with a range of chemical solutions; selected chambers can then be disrupted using targeted laser light (see image). This would give clinicians fine control over the rate of [drug release](#) to suit different patients and conditions.

This proof-of-concept work lays the foundations for moving to tests with real drugs in animals and then humans. Kiryukhin explains that the team are relying on their collaborating research groups in the UK, China and

Russia to address this challenge.

Meanwhile the A*STAR researchers are pursuing wider possibilities. "We are interested in using the chambered arrays in sensing technologies, such as detecting the level of freshness of food or diagnosing the condition of wounds and diseased tissues," Kiryukhin explains. The microchambers could release a signal, such as fluorescence, in response to the changes being sensed, for example.

More information: Alexey Ermakov et al. Polyelectrolyte-Graphene Oxide Multilayer Composites for Array of Microchambers which are Mechanically Robust and Responsive to NIR Light, *Macromolecular Rapid Communications* (2018). [DOI: 10.1002/marc.201700868](https://doi.org/10.1002/marc.201700868)

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