

Wyss Institute awarded DARPA contract to further advance sepsis therapeutic device

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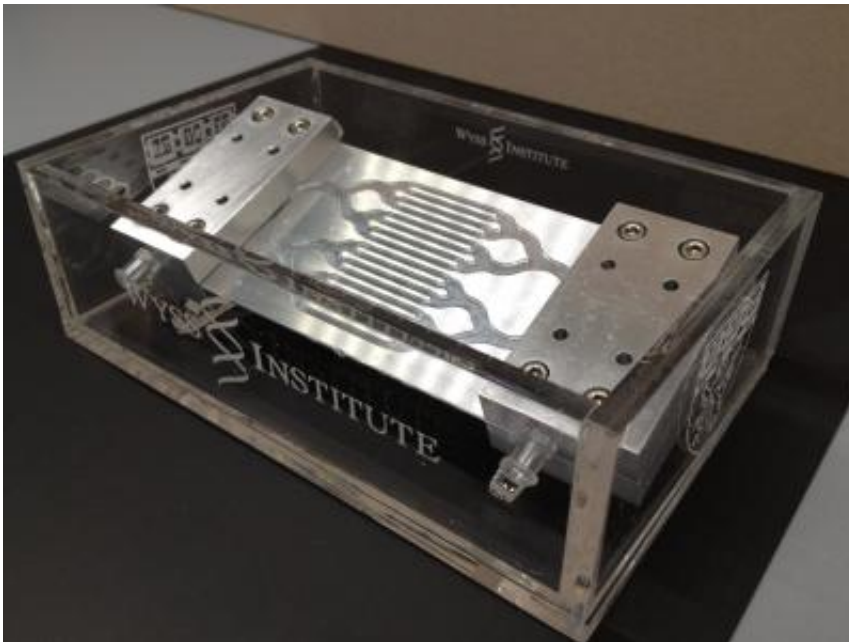


The microfluidic "Spleen-on-a-chip" developed at the Wyss Institute at Harvard University will be used to rapidly treat bloodstream infections. Credit: Wyss Institute, Harvard University

The Wyss Institute for Biologically Inspired Engineering at Harvard University announced today that it was awarded a \$9.25 million contract

from the Defense Advanced Research Projects Agency (DARPA) to further advance a blood-cleansing technology developed at the Institute with prior DARPA support, and help accelerate its translation to humans as a new type of sepsis therapy.

The device will be used to treat [bloodstream infections](#) that are the leading cause of death in critically ill patients and soldiers injured in combat.



This image shows the microfluidic "Spleen-on-a-chip" developed at the Wyss Institute at Harvard University. Credit: Wyss Institute, Harvard University

To rapidly cleanse the blood of pathogens, the patient's blood is mixed with magnetic nanobeads coated with a genetically engineered version of a human blood 'opsonin' protein that binds to a wide variety of bacteria, fungi, viruses, parasites, and toxins. It is then flowed through microchannels in the device where [magnetic forces](#) pull out the bead-

bound pathogens without removing human blood cells, proteins, fluids, or electrolytes – much like a human spleen does. The cleansed blood then flows back to the patient.

"In just a few years we have been able to develop a suite of new technologies, and to integrate them to create a powerful new device that could potentially transform the way we treat sepsis," said Wyss founding director and project leader, Don Ingber, M.D., Ph.D. "The continued support from DARPA enables us to advance our device manufacturing capabilities and to obtain validation in large animal models, which is precisely what is required to enable this technology to be moved towards testing in humans."

The team will work to develop manufacturing and integration strategies for its core pathogen-binding opsonin and Spleen-on-a-Chip fluidic separation technologies, as well as a novel [coating technology](#) called "SLIPS," which is a super-[hydrophobic coating](#) inspired from the slippery surface of a [pitcher plant](#) that repels nearly any material it contacts. By coating the inner surface of the channels of the device with SLIPS, blood cleansing can be carried out without the need for anticoagulants to prevent blood clotting.

In addition to Ingber, the multidisciplinary team behind this effort includes Wyss core faculty and Harvard School of Engineering and Applied Science faculty member Joanna Aizenberg, Ph.D., who developed the SLIPS technology; Wyss senior staff member Michael Super, PhD., who engineered the human opsonin protein; and Mark Puder, M.D., Ph.D., Associate Professor of Pediatric Surgery at Boston Children's Hospital and Harvard Medical School who will be assisting with animal studies.

Provided by Harvard University

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