A novel model for ex situ reperfusion of the human liver following subnormothermic machine perfusion

September 1, 2014 — A novel technology developed by researchers at the Massachusetts General Hospital enables the modeling of human liver transplantation in an experimental setting.

A team of scientists, physicians, and engineers from the Center for Engineering in Medicine (CEM) and the Transplant Center of the Massachusetts General Hospital (MGH) at the Harvard Medical School in Boston, MA report the development of a new technology that enables researchers to better study liver transplantation in a pre-clinical setting. The model specifically examines the reperfusion stage of transplantation, when the liver is surgically implanted into the recipient patient. This reperfusion stage of liver transplantation is particularly important because it causes significant injury to the organ, but it is also the first time the organ begins to function again inside the recipient’s body after a period of organ preservation. This model could be of significant value to those that aim to study the effects that organ preservation have on the function of the liver after transplantation, which was previously more difficult to study outside of the clinical setting. The report appears in the December 2017 issue of the journal TECHNOLOGY.

Conventionally, the liver is preserved in a cold box of ice, which allows surgeons to transplant organs after a limited period of organ preservation. With the increasing demand for high quality organs, new organ preservation techniques are rapidly being developed to both improve the quality of donor organs as well as prolong the duration of time that the liver can be kept outside of the body. Many of these new techniques have been evaluated in animal models, as well as in human liver that could not be used for transplantation. “While we were able to assess a human liver during the preservation period, we could not predict how it would function after transplantation without actually transplanting the liver,” says Dr. Bote Bruinsma, one of the lead authors on the paper, “to this end, we attempted to simulate the transplantation in the lab”.

The authors demonstrated the feasibility of the new technique in three human liver that were deemed unsuitable for transplantation. The livers were first tested in an experimental organ preservation technique called subnormothermic machine perfusion (SNMP), which was also developed by the lab at MGH. This technique connects the liver to a device that perfuses the organ with a nutrient-rich and oxygenated solution that aims to sustain the liver at a temperature of 21 degrees Celsius. After 3 hours of SNMP, the organ was connected to a new device that was used to simulate warm reperfusion of the liver as it would occur during transplantation. The liver was perfused with blood at body temperature for several hours, while the investigators took measurements relating to the function of the organ as well as the injury that it sustained.

During the reperfusion, the investigators noted production of bile as is normally observed during the clinical reperfusion period. “The biochemical changes that we observed in the blood during the reperfusion period were what we would expect to see during reperfusion clinically, which contributes to the validation of the model,” explains Dr. James Avruch, the lead author on the paper.

“We plan to use this new reperfusion model to evaluate the efficacy of experimental organ perfusion techniques in improving the outcome of liver transplantation. By applying this method, we can more safely and confidently transition new technologies into clinical practice,” explains Dr. Korkut Uygun, principal investigator on the study.
Additional co-authors of the TECHNOLOGY paper are Dr. Gautham V. Sridharan, Dr. Heidi Yeh, and Dr. James F. Markmann of the Massachusetts General Hospital, and Dr. Pepijn D. Weeder and Dr. Robert J. Porte of the University Medical Center of Groningen at the University of Groningen.

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Corresponding author for this study in TECHNOLOGY is Dr. Bote G. Bruinsma, Ph.D., botebruinsma@gmail.com.

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