

Team generates long-lasting blood vessels from reprogrammed human cells

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Massachusetts General Hospital (MGH) researchers have used vascular precursor cells derived from human induced pluripotent stem cells (iPSCs) to generate, in an animal model, functional blood vessels that lasted as long as nine months. In their report being published in *PNAS Early Edition*, the investigators describe using iPSCs – reprogrammed adult cells that have many of the characteristics of embryonic stem cells – from both healthy adults and from individuals with type 1 diabetes to generate blood vessels on the outer surface of the brain or under the skin of mice.

"The discovery of ways to bring <u>mature cells</u> back to a 'stem-like' state that can differentiate into many different types of tissue has brought enormous potential to the field of cell-based regenerative medicine, but the challenge of deriving functional cells from these iPSCs still remains," says Rakesh Jain, PhD, director of the Steele Laboratory for Tumor Biology at MGH and co-senior author of the study. "Our team has developed an efficient method to generate vascular precursor cells from human iPSCs and used them to create networks of engineered <u>blood</u> <u>vessels</u> in living mice."

The ability to regenerate or repair blood vessels could make a crucial difference in the treatment of cardiovascular disease—which continues to be the number one cause of death in the U.S.—and other conditions caused by <u>blood vessel damage</u>, such as the vascular complications of diabetes. In addition, providing a vascular supply to newlygenerated tissue remains one of the greatest barriers facing efforts to build solid organs through tissue engineering.

Several previous studies have generated from iPSCs the types of cells required to build blood vessels—<u>endothelial cells</u> that line vessels and <u>connective tissue cells</u> that provide structural support—but those cells could not form long-lasting vessels once introduced into animal models. "The

biggest challenge we faced during the early phase of this project was establishing a reliable protocol to generate endothelial cell lines that produced great quantities of precursor cells that could generate strong, durable blood vessels," says co-senior author Dai Fukumura, MD, PhD, also of the Steele Lab.

The MGH team adapted a method originally used to derive endothelial cells from human <u>embryonic</u> <u>stem cells</u> (hESCs). But while that method used a single protein marker to identify vascular progenitors, the researchers sorted out iPSCderived cells that expressed not only that protein but also two other protein markers of vascular potential. They then expanded that population using a culture system that team members had previously developed to differentiate endothelial cells from hESCs and confirmed that only iPSCderived cells expressing all three markers generated endothelial cells with the full potential of forming blood vessels.

To test the capacity of those cells to generate functional blood vessels, they implanted onto the surface of the brains of mice a combination of the iPSC-derived endothelial precursor cells expressing the three markers with the mesenchymal precursors that generate essential structural cells. Within two weeks, the implanted cells had formed networks of blood-perfused vessels that appeared to function as well as adjacent natural vessels and continued to function for as long as 280 days in the living animals. While implantation of the combined precursor populations under the skin of the animals also generated functional blood vessels, it required implantation of five times more cells, and the vessels were short-lived, an observation consistent with the team's previous studies of vessel generation in these two locations.

Because patients with <u>type 1 diabetes</u> (T1D), which can damage blood vessels, could benefit from the ability to make new blood vessels, the researchers



wanted to determine whether iPSCs derived from the cells of such patients could help generate functional blood vessels. As with cells from healthy individuals, precursors derived from T1D-iPSCs were able to generate functional, long-lasting blood vessels. However, the researchers note, different lines of the T1D-iPSCs – including different lines derived from the same patient – showed differences in cell-generating potential, indicating the need to better understand the underlying mechanisms.

"The potential applications of iPSC-generated blood vessels are broad – from repairing damaged vessels supplying the heart or brain to preventing the need to amputate limbs because of the vascular complication of diabetes," says co-lead author Rekha Samuel, MD, of the Steele Laboratory, now at the Christian Medical College, Vellore, India. "But first we need to deal with such challenges as the variability of iPSC lines and the long-term safety issues involved in the use of these <u>cells</u>, which are being addressed by researchers around the world. We also need better ways of engineering the specific type of endothelial cell needed for specific organs and functions."

More information: Generation of functionally competent and durable engineered blood vessels from human induced pluripotent stem cells , *PNAS*, <u>www.pnas.org/cgi/doi/10.1073/pnas.1310675110</u>

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