

Researchers rescue photoreceptors, prevent blindness in animal models of AMD

 Blood Cell
 Induced
Pluripotent
Stem Cell
(iPS cell)
 RPE Cell

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The researchers will take a patient's own blood cells, and in a lab, convert them into iPS cells capable of becoming any type of cell in the body. The iPS cells are then programmed to become retinal pigment epithelial cells, the type of cell that dies early in the geographic atrophy form of AMD. Credit: National Eye Institute

Using a novel patient-specific stem cell-based therapy, researchers at the National Eye Institute (NEI) prevented blindness in animal models of geographic atrophy, the advanced "dry" form of age-related macular degeneration (AMD), which is a leading cause of vision loss among

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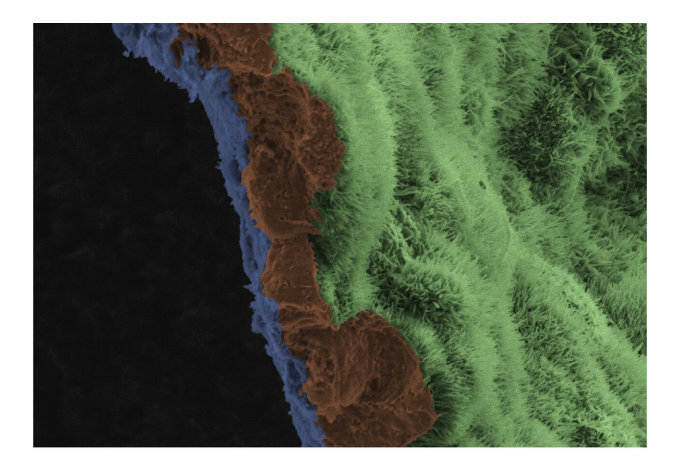
people age 65 and older. The protocols established by the animal study, published January 16 in *Science Translational Medicine*, set the stage for a first-in-human clinical trial testing the therapy in people with geographic atrophy, for which there is currently no treatment.

"If the clinical trial moves forward, it would be the first ever to test a stem cell-based therapy derived from induced pluripotent <u>stem cells</u> (iPSC) for treating a disease," said Kapil Bharti, Ph.D., a Stadtman Investigator and head of the NEI Unit on Ocular and Stem Cell Translational Research. Bharti was the lead investigator for the animalmodel study published in STM. The NEI is part of the National Institutes of Health.

The therapy involves taking a patient's blood <u>cells</u> and, in a lab, converting them into iPS cells, which can become any type of cell in the body. The iPS cells are programmed to become retinal pigment epithelial cells, the type of cell that dies early in the geographic atrophy stage of macular degeneration. RPE cells nurture photoreceptors, the light-sensing cells in the retina. In geographic atrophy, once RPE cells die, photoreceptors eventually also die, resulting in blindness. The therapy is an attempt to shore up the health of remaining photoreceptors by replacing dying RPE with iPSC-derived RPE.

Before they are transplanted, the iPSC-derived RPE are grown in tiny sheets one cell thick, replicating their natural structure within the eye. This monolayer of iPSC-derived RPE is grown on a biodegradable scaffold designed to promote the integration of the cells within the retina. A specially designed surgical tool was built for the task of inserting the patch of cells between the RPE and the photoreceptors.





A scanning electron micrograph image shows a polarized RPE monolayer on a biodegradable scaffold. The image is colored to highlight the scaffold in blue, three RPE cells (brown), and the apical process of cells in RPE monolayer are light green. Credit: Kapil Bharti, Ph.D., NEI

In the STM paper, Bharti describes testing the approach in rat and pig models. Ten weeks after the human iPSC-derived RPE patches were implanted in the animals' retinas, imaging studies confirmed that the labmade cells had integrated within the animal retina.

The investigators report the transplanted cells functioned properly. Immunostaining confirmed that the iPSC-derived RPE expressed the gene RPE65, suggesting the lab-made cells had reached a crucial stage of



maturity necessary to maintain <u>photoreceptor</u> health. RPE65 is necessary for the regeneration of visual pigment within the photoreceptors and is an essential component for vision.

Further tests showed that the transplanted RPE cells were pruning photoreceptors via phagocytosis, another RPE function that helps keep photoreceptors healthy. In addition, electrical responses recorded from photoreceptors rescued by RPE patches were normal; whereas photoreceptors treated with a control empty scaffold had died.

The animal studies inform the processes for conducting a first-in-human clinical trial testing the treatment. Key to that workflow are good manufacturing practice (GMP) protocols, which were established to ensure that the iPSC-derived RPE were a safe and efficacious, clinical-grade product. GMP protocols are also crucial for making the therapy reproducible, a requirement for scaling up production and Food and Drug Administration approval.

"Adhering to the protocols helps ensure that the transplanted cells function reliably, and that unintended consequences are minimized," Bharti said.

A key concern with any stem cell therapy is its oncogenic potential: the ability for cells to multiply uncontrollably and form tumors. The researchers genetically analyzed the iPSC-derived RPE cells and found no genetic mutations linked to tumor growth.

"The protocol also minimizes the chance of rejection by developing the iPSC-RPE with an individual's autologous (own) blood cells," Bharti said.

The planning of a Phase I clinical trial testing the safety of the iPSCbased <u>therapy</u> for geographic atrophy is underway and will be initiated



after U.S. Food and Drug Administration approval.

More information: R. Sharma el al., "Clinical-grade stem cell–derived retinal pigment epithelium patch rescues retinal degeneration in rodents and pigs," *Science Translational Medicine* (2018). <u>stm.sciencemag.org/lookup/doi/... scitranslmed.aat5580</u>

Provided by National Eye Institute

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