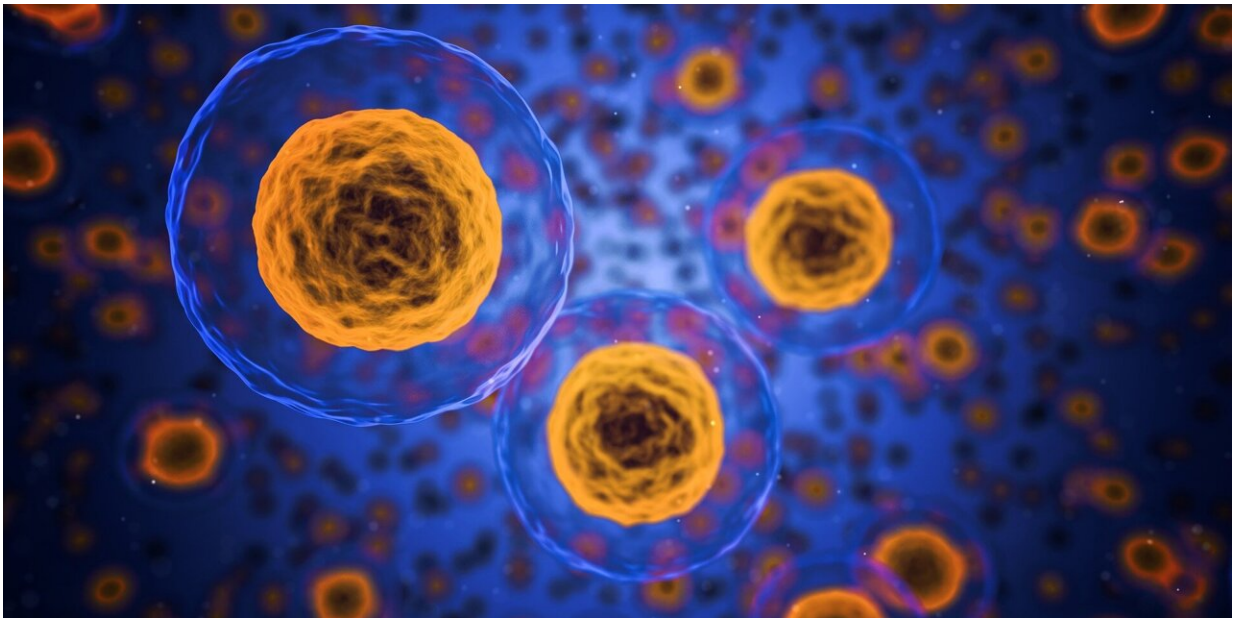


Duping antibodies with a decoy, researchers aim to prevent rejection of transplanted cells

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Researchers at UCSF have developed a novel, potentially life-saving approach that may prevent antibodies from triggering immune rejection of engineered therapeutic and transplant cells.

Rejection mediated by [antibodies](#)—as opposed to the chemical assault initiated by [immune cells](#)—has proven particularly tricky to resolve, a factor holding held back the development of some of these treatments.

The new strategy, described in the Monday, Jan. 2, 2023 issue of *Nature Biotechnology*, involved using a "decoy" receptor to capture the antibodies and take them out of circulation before they could kill the therapeutic [cells](#), that they treat as invading foreigners. The tactic may also be useful for [organ transplants](#).

"This antibody-mediated rejection is really hard to overcome," said Tobias Deuse, MD, the Julien I.E. Hoffman, M.D. Chair in Cardiac Surgery and senior author on the study. "So, rather than trying to suppress the patient's [immune system](#), we looked for ways we can alter the cells that the patient will receive and better enable them to survive."

Protecting friendly-but-foreign cells

The most celebrated cellular therapies in the U.S. are [chimeric antigen receptor](#) (CAR) T-cell therapies. These CAR-T therapies are often used to successfully treat specific forms of lymphomas, a type of often-deadly cancer. But deploying them against solid tumors has proved much more difficult.

Until recently, most CAR-T therapies have been made using the patient's own cells, but the long-term commercial viability of cellular therapies of all types will rely on "allogeneic" cells—mass-produced therapeutic cells grown from a source outside the patient.

As with transplanted organs, the recipient's immune system is likely to treat any outsider cells, or tissues developed from them, as foreign and to reject them, according to Deuse. "We have been through this with [organ transplantation](#), so we know what's coming for cellular transplantation," said Deuse, a cardiac transplant surgeon who is no stranger to the troubles caused by [immune rejection](#). "This issue is likely to be a severe obstacle in any type of allogeneic cell transplantation."

Clinical trials of allogeneic CAR-T therapies have had worse outcomes than treatments derived from the patients' cells, Deuse noted, adding that immunotherapy entails the added challenge of these free-floating cells being more exposed to immune attack than those in a transplanted organ.

"We have to find better ways to protect these cells," he said.

Duping antibodies with a decoy

Normally when an antibody binds to a cell, it acts as a sort of tag, calling out for an immune cell to bind to the antibody and set off an efficient process of destroying the tagged cell. To stop this [chain reaction](#), Deuse and his team devised a method to catch the antibodies before they bind to cells, preventing activation of the immune response.

The researchers genetically engineered three types of cells—insulin-producing pancreatic islet cells, thyroid cells, and CAR-T cells—so that each type made and displayed large numbers of a protein called CD64 on their surfaces.

On these engineered cells, CD64, which tightly binds the antibodies responsible for this type of immune rejection, acted as a kind of decoy, capturing the antibodies and binding them to the engineered cell, so they wouldn't activate immune cells.

"We saw that we can snatch up high levels of these antibodies, which resulted in very robust protection for the [therapeutic cells](#)," said Deuse. "This is clear proof-of-concept for this approach."

There's more work to do before the approach can be tested on cells that are designed to be therapeutics or transplanted cells, he said. While such cells are biologically sophisticated, they are also expensive and hard to manufacture.

"My hope is that our concept can help bring about the development of universally usable allogeneic cells." said Deuse. "That would make treatment with cellular therapies cheaper and more accessible, putting them within reach for many more patients."

Other UCSF authors are Alessia Gravina, Grigol Tediashvili, and Sonja Schrepfer of the Transplant and Stem Cell Immunobiology (TSI)-Lab, Raja Rajalingam of the Immunogenetics and Transplantation Laboratory, Zoe Quandt of the Division of Diabetes, Endocrinology and Metabolism. Chad Deisenroth of the U.S. Environmental Protection Agency, Center for Computational Toxicology & Exposure is also an author.

More information: Tobias Deuse, Protection of cell therapeutics from antibody-mediated killing by CD64 overexpression, *Nature Biotechnology* (2023). DOI: [10.1038/s41587-022-01540-7](https://doi.org/10.1038/s41587-022-01540-7). www.nature.com/articles/s41587-022-01540-7

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