

Living drugs new frontier for cancer patients out of options

June 12 2017, by Lauran Neergaard



In this photo taken March 29, 2017, Dr. David Maloney of the Fred Hutchinson Cancer Research Center is greeted by patient Ken Shefveland, whose lymphoma was successfully treated with CAR-T cell therapy. Immune therapy is the hottest trend in cancer care and its next frontier is creating "living drugs" that grow inside the body into an army that seeks and destroys tumors. (AP Photo/Elaine Thompson)

Ken Shefveland's body was swollen with cancer, treatment after treatment failing until doctors gambled on a radical approach: They removed some of his immune cells, engineered them into cancer



assassins and unleashed them into his bloodstream.

Immune therapy is the hottest trend in cancer care and this is its next frontier—creating "living drugs" that grow inside the body into an army that seeks and destroys tumors.

Looking in the mirror, Shefveland saw "the cancer was just melting away." A month later doctors at the Fred Hutchinson Cancer Research Center couldn't find any signs of lymphoma in the Vancouver, Washington, man's body.

"Today I find out I'm in full remission—how wonderful is that?" said Shefveland with a wide grin, giving his physician a quick embrace.

This experimental therapy marks an entirely new way to treat cancer—if scientists can make it work, safely. Early-stage studies are stirring hope as one-time infusions of supercharged immune cells help a remarkable number of patients with intractable leukemia or lymphoma.

"It shows the unbelievable power of your immune system," said Dr. David Maloney, Fred Hutch's medical director for cellular immunotherapy who treated Shefveland with a type called CAR-T cells.

"We're talking, really, patients who have no other options, and we're seeing tumors and leukemias disappear over weeks," added immunotherapy scientific director Dr. Stanley Riddell. But, "there's still lots to learn."

T cells are key immune system soldiers. But cancer can be hard for them to spot, and can put the brakes on an immune attack. Today's popular immunotherapy drugs called "checkpoint inhibitors" release one brake so nearby T cells can strike. The new cellular immunotherapy approach aims to be more potent: Give patients stronger T cells to begin with.



Currently available only in studies at major cancer centers, the first CAR-T cell therapies for a few blood cancers could hit the market later this year. The Food and Drug Administration is evaluating one version developed by the University of Pennsylvania and licensed to Novartis, and another created by the National Cancer Institute and licensed to Kite Pharma.

CAR-T therapy "feels very much like it's ready for prime time" for advanced blood cancers, said Dr. Nick Haining of the Dana-Farber Cancer Institute and Broad Institute of MIT and Harvard, who isn't involved in the development.



In this photo taken March 28, 2017, Dr. Stanley Riddell describes how a patient's large tumor rapidly shrank in an immunotherapy study at the Fred Hutchinson Cancer Research Center in Seattle. Riddell's team studies "living drugs" that are genetically engineered from patients' own immune cells to make them better cancer killers. (AP Photo/Elaine Thompson)



Now scientists are tackling a tougher next step, what Haining calls "the acid test": Making T cells target far more common cancers—solid tumors like lung, breast or brain cancer. Cancer kills about 600,000 Americans a year, including nearly 45,000 from leukemia and lymphoma.

"There's a desperate need," said NCI immunotherapy pioneer Dr. Steven Rosenberg, pointing to queries from hundreds of patients for studies that accept only a few.

For all the excitement, there are formidable challenges.

Scientists still are unraveling why these living cancer drugs work for some people and not others.

Doctors must learn to manage potentially life-threatening side effects from an overstimulated immune system. Also concerning is a small number of deaths from brain swelling, an unexplained complication that forced another company, Juno Therapeutics, to halt development of one CAR-T in its pipeline; Kite recently reported a death, too.

And, made from scratch for every patient using their own blood, this is one of the most customized therapies ever and could cost hundreds of thousands of dollars.

"It's a Model A Ford and we need a Lamborghini," said CAR-T researcher Dr. Renier Brentjens of New York's Memorial Sloan Kettering Cancer Center, which, like Hutch, has a partnership with Juno.

In Seattle, Fred Hutch offered a behind-the-scenes peek at research underway to tackle those challenges. At a recently opened immunotherapy clinic, scientists are taking newly designed T cells from the lab to the patient and back again to tease out what works best.



"We can essentially make a cell do things it wasn't programmed to do naturally," explained immunology chief Dr. Philip Greenberg. "Your imagination can run wild with how you can engineer cells to function better."

TWO LONG WEEKS TO BREW A DOSE



In this photo taken March 29, 2017, Ken Shefveland smiles as he talks about the success of his treatment in a research study run by the Fred Hutchinson Cancer Research Center in Seattle. Shefveland had some of his immune cells genetically reprogrammed into an army of "living drugs" that could better seek and destroy his cancer, the next frontier in immunotherapy. (AP Photo/Elaine Thompson)

The first step is much like donating blood. When leukemia patient Claude Bannick entered a Hutch CAR-T study in 2014, nurses hooked him to a machine that filtered out his white blood cells, including the T



cells.

Technicians raced his bag of cells to a factory-like facility that's kept so sterile they must pull on germ-deflecting suits, booties and masks just to enter. Then came 14 days of wait and worry, as his cells were reprogrammed.

Bannick, 67, says he "was almost dead." Chemotherapy, experimental drugs, even a bone marrow transplant had failed, and "I was willing to try anything."

GENETICALLY ENGINEERING CELLS

The goal: Arm T cells with an artificial receptor, a tracking system that can zero in on identifying markers of cancer cells, known as antigens. For many leukemias and lymphomas, that's an antigen named CD19.

Every research group has its own recipe but generally, scientists infect T cells with an inactive virus carrying genetic instructions to grow the desired "chimeric antigen receptor." That CAR will bind to its target cancer cells and rev up for attack.

Millions of copies of <u>engineered cells</u> are grown in incubators, Hutch technicians pulling out precious batches to monitor if they're ready for waiting patients.

If they work, those cells will keep multiplying in the body. If they don't, the doctors send blood and other samples back to researchers like Riddell to figure out why.

WHAT'S THE DATA?

Small, early studies in the U.S. made headlines as 60 percent to 90



percent of patients trying CAR-Ts as a last resort for leukemia or lymphoma saw their cancer rapidly decrease or even become undetectable. Last week, Chinese researchers reported similar early findings as 33 of 35 patients with another blood cancer, multiple myeloma, reached some degree of remission within two months.



In this photo taken March 21, 2017, lymphoma patient Peter Bjazevich receives cellular immunotherapy as part of a study at the Fred Hutchinson Cancer Research Center in Seattle. Immune therapy is the hottest trend in cancer care and this is its next frontier, "living drugs" that grow inside the body into an army that seeks and destroys tumors. (AP Photo/Elaine Thompson)

Too few people have been studied so far to know how long such responses will last. A recent review reported up to half of leukemia and lymphoma patients may relapse.



There are long-term survivors. Doug Olson in 2010 received the University of Pennsylvania's CAR-T version for leukemia. The researchers were frank—it had worked in mice but they didn't know what would happen to him.

"Sitting here almost seven years later, I can tell you it works," Olson, now 70, told a recent meeting of the Leukemia and Lymphoma Society.

Bannick, the Hutch patient treated in 2014, recalls Maloney calling him "the miracle man." He had some lingering side effects that required blood-boosting infusions but says CAR-T is "giving me a second life."

SCARY SIDE EFFECTS

"The more side effects you have, that sort of tells everybody it's working," said Shefveland, who was hospitalized soon after his treatment at Hutch when his blood pressure collapsed. His last clear memory for days: "I was having a conversation with a nurse and all of a sudden it was gibberish."

As CAR-T cells swarm the cancer, an immune overreaction called "cytokine release syndrome" can trigger high fevers and plummeting blood pressure and in severe cases organ damage. Some patients also experience confusion, hallucinations or other neurologic symptoms.

Treatment is a balancing act to control those symptoms without shutting down the cancer attack.

Experienced cancer centers have learned to expect and watch for these problems. "And, most importantly, we've learned how to treat them," said Dr. Len Lichtenfeld of the American Cancer Society, who is watching CAR-T's development.



FIGHTING SOLID TUMORS WILL BE HARDER



In this photo taken March 29, 2017, cell production associate Herley Beyene places containers of immune cells in a centrifuge at the Fred Hutchinson Cancer Research Center in Seattle. Researchers are genetically reprogramming patients' immune cells to create "living drugs" that better seek and destroy cancer. (AP Photo/Elaine Thompson)

CAR-Ts cause collateral damage, killing some healthy <u>white blood cells</u>, called B cells, along with cancerous ones because both harbor the same marker. Finding the right target to kill solid tumors but not healthy organ tissue will be even more complicated.

"You can live without some normal B cells. You can't live without your lungs," Riddell explained.



Early studies against solid tumors are beginning, targeting different antigens. Time-lapse photos taken through a microscope in Riddell's lab show those new CAR-T cells crawling over aggressive breast cancer, releasing toxic chemicals until tumor cells shrivel and die.

CARs aren't the only approach. Researchers also are trying to target markers inside <u>tumor cells</u> rather than on the surface, or even gene mutations that don't form in healthy tissue.

"It's ironic that the very mutations that cause the cancer are very likely to be the Achilles heel," NCI's Rosenberg said.

And studies are beginning to test CAR-Ts in combination with older immunotherapy drugs, in hopes of overcoming tumor defenses.

HOW WILL PATIENTS GET THE FIRST CAR-T THERAPIES?

If the FDA approves Novartis' or Kite's versions, eligible leukemia and lymphoma patients would be treated at cancer centers experienced with this tricky therapy. Their T <u>cells</u> would be shipped to company factories, engineered, and shipped back. Gradually, more hospitals could offer it.

Because only certain patients would qualify for the first drugs, others would have to search for CAR-T studies to try the treatment. A drug industry report lists 21 CAR-T therapies in development by a dozen companies.

"This is the hope of any <u>cancer</u> patient, that if you stay in the game long enough, the next treatment's going to be just around the corner," said Shefveland, the Hutch patient.



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Citation: Living drugs new frontier for cancer patients out of options (2017, June 12) retrieved 5 May 2024 from <u>https://medicalxpress.com/news/2017-06-drugs-frontier-cancer-patients-options.html</u>

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