

Cancer Immunotherapy Drug Simultaneously Targets Two Proteins

April 27, 2018 By [National Cancer Institute](#)

A growing number of patients with cancer have benefited from drugs that help the immune system fight cancer, such as [immune checkpoint inhibitors](#). But most patients with cancers do not respond to these treatments.

To try to develop more effective [immunotherapies](#), two groups of researchers, working independently, have developed a type of drug that simultaneously targets two proteins involved in suppressing the body's immune response against tumors.

The researchers created the drugs by fusing a [receptor](#) for a protein called TGF-beta to a [monoclonal antibody](#) that targets a checkpoint protein, such as PD-L1 or CTLA-4. The result is a single hybrid molecule that blocks two [pathways](#) used by tumors to evade the immune system.

TGF-beta is often found in the [tumor microenvironment](#) and may, like immune checkpoint proteins, regulate the activity of immune cells. The TGF-beta receptor portion of the new immunotherapy drug binds to TGF-beta, "trapping" the protein and rendering it useless.

"TGF-beta is well known to be a bad actor in the tumor microenvironment, because it inhibits immune cells called [T cells](#), which are the good guys," said Jeffrey Schlom, PhD, of NCI's [Center for Cancer Research](#) (CCR), who has led several [preclinical studies of the new immunotherapy drug](#).

The drug, which is a type of therapy called a bifunctional [fusion protein](#) or a Y-trap, may be more effective than an immune checkpoint inhibitor alone while not substantially increasing the side effects for patients, according to the results of several recent studies.

Based on the NCI preclinical studies, Julius Strauss, MD, of CCR and his colleagues began to test the drug, which targets PD-L1, in clinical trials. Meanwhile, a new study from researchers at the Johns Hopkins School of Medicine has [also demonstrated the potential of two Y-traps](#) tested in mouse models of cancer, including one that, like NCI's drug, targets PD-L1.

"Targeting both a checkpoint inhibitor and other molecules, such as TGF-beta, gives you a much more effective way to block the immune suppression we so often see in tumors," said Jeffrey Sosman, MD, of the Robert H. Lurie Comprehensive Cancer Center of Northwestern University, who investigates immunotherapies but was not involved in the NCI clinical trials or the Johns Hopkins study.

Using a single drug to treat a patient's cancer has some advantages over using multiple drugs, such as reducing potential side effects, Sosman noted.

Encouraging Results from Clinical Trials

In January 2018, NCI researchers reported [the results of a phase 1 clinical trial of this approach with a molecule called M7824](#) in 19 patients with advanced cancer that had been treated previously. Based on the findings, including one patient who had a [complete response](#), the researchers have expanded the trial to include additional patients.

Inhibiting both PD-L1 and TGF-beta is a promising therapeutic strategy "because these key pathways have independent and complementary roles in suppressing the immune response," said CCR's James Gulley, MD, PhD, an immunotherapy researcher who co-led the NCI clinical trial.

"This is a very exciting approach," he added, noting that in clinical trials the [side effects](#) of the new technology were similar to those of inhibitors that block the interaction of PD-1 and PD-L1 such as [pembrolizumab \(Keytruda\)](#) and [nivolumab \(Opdivo\)](#) and were largely manageable.

The researchers are also conducting early-phase clinical trials of [M7824 in patients with pancreatic cancer](#) and [patients whose cancers are associated with the human papillomavirus](#). Additional phase 2 studies are in development, according to Schlom.

Testing Multiple Y-Traps in Mouse Models

In experiments with mice, the two Y-traps developed at Johns Hopkins were more effective than the corresponding immune checkpoint inhibitors, including the anti-CTLA-4 antibody [ipilimumab \(Yervoy\)](#) and the anti-PD-L1 antibodies [atezolizumab \(Tecentriq\)](#) and [avelumab \(Bavencio\)](#), the researchers reported in Nature Communications on February 21.

For example, a Y-trap targeting CTLA-4 and TGF-beta slowed the growth of tumors that did not respond to ipilimumab. Similarly, a Y-trap targeting PD-L1 and TGF-beta slowed the growth of tumors that did not respond to atezolizumab and avelumab.

"The antibody goes after CTLA-4 or PD-L1, and at the same time it can trap a molecule of interest, such as TGF-beta, that is helping to suppress the immune system," said Atul Bedi, MD, who led the Johns Hopkins team.

The researchers initially became interested in targeting TGF-beta as a way to limit the production of [regulatory T cells](#), a type of immune cell that suppresses the activity of the immune system. Regulatory T cells infiltrate many advanced cancers, and this infiltration is associated with a poor prognosis.

"You need a powerful set of tools to address this situation," he added. "With the Y-trap, we now

have the ability to counteract a key mechanism of resistance to the immune response.”

Developing a Family of Y-Traps

Bedi noted, however, that regulatory T cells are not the only way tumors evade the immune system. The Johns Hopkins team is developing a family of Y-traps that can be used to target multiple factors that play a role in the body’s immune response to tumors.

Johns Hopkins has licensed the Y-trap technology to a new company that could begin to test the approach in patients, according to Bedi.

“The concept of using combined therapies for patients with cancer is a central theme in the development of new immunotherapies,” said Sosman. “The beauty of this approach is that it combines multiple therapies in a single molecule.”

The idea of using a single molecule for this purpose is not new, Sosman added. “But we now have examples of this approach that are getting into the clinic and showing activity in patients. We have molecules that clearly work.”

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