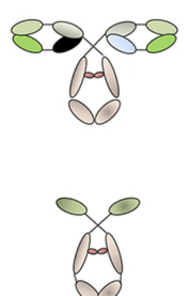


## Camel, alpaca antibodies target anticancer viruses directly to tumors

February 19 2015, by Julia Evangelou Strait



Typical antibodies (top) unfold in the harsh environment of the cell. Camelid antibodies (bottom) are smaller and more stable. Credit: S. KALIBEROV

Using antibodies from camels and alpacas, scientists at Washington University School of Medicine in St. Louis have found a way to deliver anticancer viruses directly to tumor cells, leaving other types of cells uninfected.

The research appears Feb. 18 in Molecular Therapy—Oncolytics.



The new study was conducted in <u>human cells</u> grown in the lab. According to the investigators, it demonstrates the possibility of directly delivering genetically engineered <u>viruses</u> to specific cells. The goal is to infect only cancer cells and then trigger the virus to replicate until the cells burst, killing them and releasing more of the targeted viruses.

The scientists showed that unlike <u>human antibodies</u> or those of most other animals, the antibodies of camels and alpacas survive the <u>harsh</u> <u>environment</u> inside cells and retain the ability to seek out targets, potentially solving a longstanding problem in the field of gene therapy.

"For decades, investigators have been putting human or mouse antibodies on viruses, and they haven't worked—the antibodies would lose their targeting ability," said senior author David T. Curiel, MD, PhD, distinguished professor of radiation oncology. "It was a technical problem. During replication, the virus is made in one part of the cell, and the antibody is made in another. To incorporate the two, the antibody is dragged through the internal fluid of the cell. This is a harsh environment for the antibodies, so they unfold and lose their targeting ability."

Antibodies are proteins of the immune system that travel through the bloodstream and recognize potential threats to the body, whether bacteria, viruses or <u>abnormal cells</u>. Most antibodies have a characteristic Y shape. The tips of the Y form a "lock" that binds to a specific "key" carried by foreign bodies that the immune system should destroy.

According to Curiel, recent work by other groups has identified an unusually small and stable class of antibodies made by camels, alpacas and related species collectively classified as camelids. The "lock" of camelid antibodies consists of the stem of the Y only, so it can't unfold in the harsh internal environment of the cell.



"We found that when we incorporated the camelid antibodies into the virus, they retained their binding specificity," Curiel said. "This opens the door to targeting these antibodies to specific tumor markers."

Currently available viral-based cancer therapeutics and those in human trials are not targeted directly to <u>tumor cells</u>. Indeed, these therapeutic viruses can infect almost any cell type. But they are genetically engineered to replicate only inside tumor cells. The viral DNA code is rewritten so that only tumor cells carry the molecular switch that activates the virus's replication machinery. Once that switch is flipped, the virus replicates until the cell bursts, killing the cell and releasing more viruses that can potentially infect other cells.

Limiting virus replication and cell death only to <u>cancer cells</u> is helpful in reducing side effects of the treatment. But the technology is still inefficient because so much of the injected virus never finds the tumor and is essentially wasted.

"We want this new level of targeting specificity because it would allow us to inject the virus into the bloodstream, where it would exclusively infect and replicate in tumor cells, even if they are disseminated throughout the body," Curiel said. "These viruses are already engineered to replicate only in tumors. These camelid antibodies would enable them to become even more tumor-specific and open the door for use in metastatic cancer."

After demonstrating this proof of concept, the next step is to design viruses with camelid <u>antibodies</u> against a specific tumor and test it in an animal model of the disease, Curiel said.

**More information:** van Erp EA, Kaliberova LN, Kaliberov SA, Curiel DT. "Retargeted oncolytic adenovirus displaying a single variable domain of camelid heavy-chain-only antibody in a fiber protein."



*Molecular Therapy—Oncolytics*. Feb. 18, 2015. DOI: <u>10.1038/mto.2015.1</u>

## Provided by Washington University School of Medicine in St. Louis

Citation: Camel, alpaca antibodies target anticancer viruses directly to tumors (2015, February 19) retrieved 6 May 2024 from <u>https://medicalxpress.com/news/2015-02-camel-alpaca-antibodies-anticancer-viruses.html</u>

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