

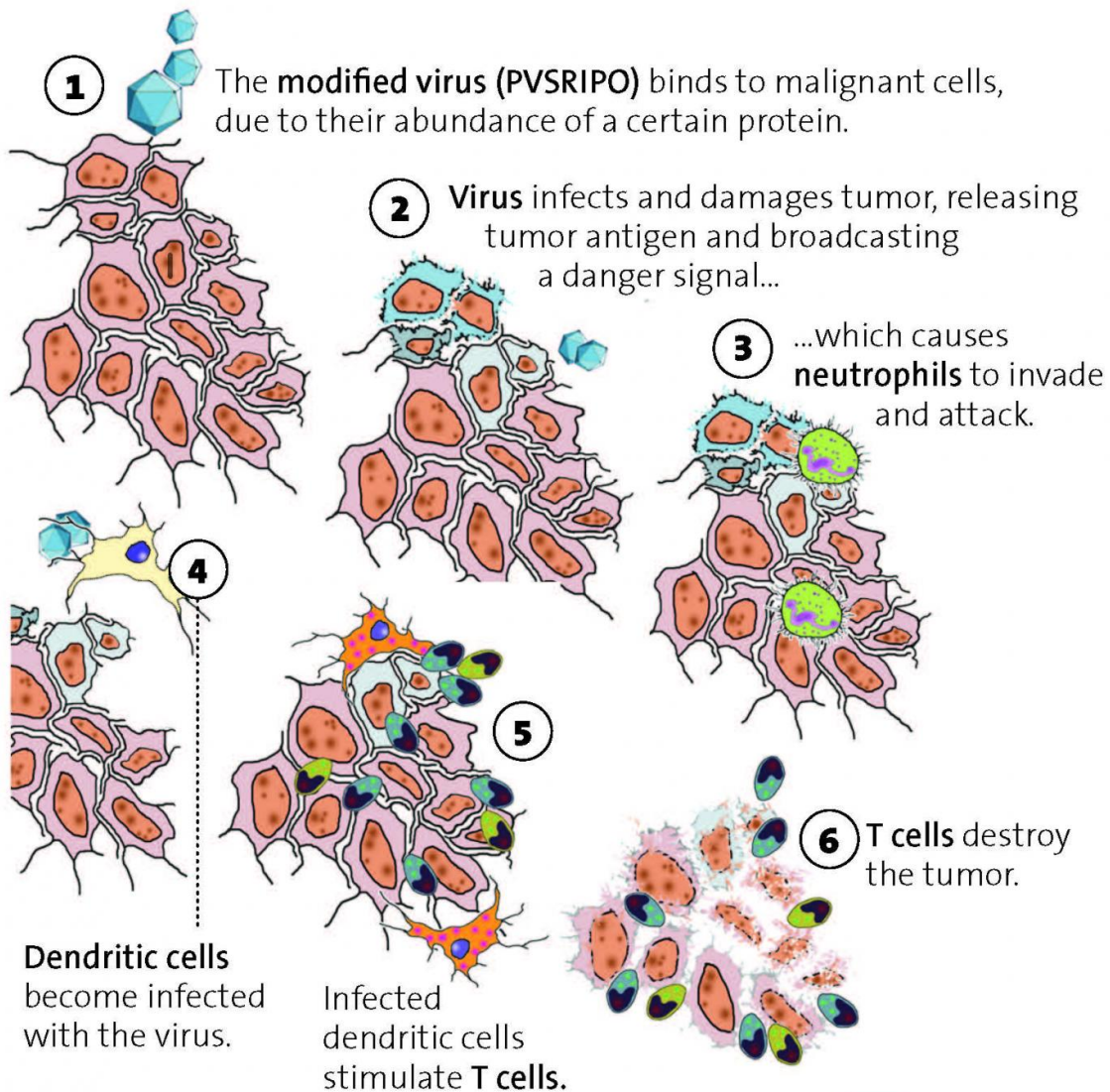
# **Poliovirus therapy induces immune responses against cancer**

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## An old virus fighting a new battle

A modified polio vaccine has shown success in clinical trials at killing cancer cells and destroying tumors. Scientists have learned more details in how this virus engages the bodies' own immune system to attack.



Duke Health



A modified polio vaccine has shown success in clinical trials at killing cancer cells and destroying tumors. Scientist have learned more details in how this virus engages the bodies' own immune system to attack. Credit: Duke Health

An investigational therapy using modified poliovirus to attack cancer tumors appears to unleash the body's own capacity to fight malignancies by activating an inflammation process that counter's the ability of cancer cells to evade the immune system.

Describing this process in a paper published Sept. 20 in the journal *Science Translational Medicine*, Duke Cancer Institute researchers provide the first published insight into the workings of a therapy that has shown promise in early clinical trials in patients with recurrent glioblastoma, a lethal form of brain [cancer](#). The modified [poliovirus](#) received a breakthrough therapy designation from the Food and Drug Administration last year, expediting research.

"We have had a general understanding of how the modified poliovirus works, but not the mechanistic details at this level," said co-senior author Matthias Gromeier, M.D., a professor in the Duke Department of Neurosurgery who developed the therapy. "This is hugely important to us. Knowing the steps that occur to generate an [immune response](#) will enable us to rationally decide whether and what other therapies make sense in combination with poliovirus to improve patient survival."

Gromeier, with expertise in cancer biology, collaborated with fellow Duke researcher and co-senior author Smita Nair, Ph.D., an immunologist and professor in the Department of Surgery. The research team elucidated how the poliovirus works not only to attack cancer [cells](#) directly, but also to trigger a longer-lasting immune response that appears to inhibit regrowth of the [tumor](#).

Using human melanoma and breast cancer cell lines, and then validating the findings in mouse models, the researchers found that the modified poliovirus therapy starts by attaching to malignant cells, which have an

abundance of CD155 protein. The CD155 protein is otherwise known as the poliovirus receptor. The modified virus then begins to attack the [tumor cells](#), directly killing many, but not all. This releases tumor antigens.

The second phase of assault is more complicated. By killing the cancer cells, the modified poliovirus triggers an alarm within the immune system, alerting the body's defenses to go on the attack.

This appears to occur when the modified poliovirus infects [dendritic cells](#) and macrophages. Dendritic cells then present tumor to T cells to launch an immune response. Once the immune system is activated against the poliovirus-infected tumor, the [cancer cells](#) can no longer hide and they remain vulnerable to ongoing immune attack.

"Not only is poliovirus killing tumor cells, it is also infecting the antigen-presenting cells, which allows them to function in such a way that they can now raise a T-cell response that can recognize and infiltrate a tumor," Nair said. "This is an encouraging finding, because it means the poliovirus stimulates an innate inflammatory response."

Nair and Gromeier said further studies will focus on the additional immune activity following exposure to the modified virus.

**More information:** M.C. Brown et al., "Cancer immunotherapy with recombinant poliovirus induces IFN-dominant activation of dendritic cells and tumor antigen-specific CTLs," *Science Translational Medicine* (2017). [stm.sciencemag.org/lookup/doi/10.1126/scitranslmed.aan4220](http://stm.sciencemag.org/lookup/doi/10.1126/scitranslmed.aan4220)

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