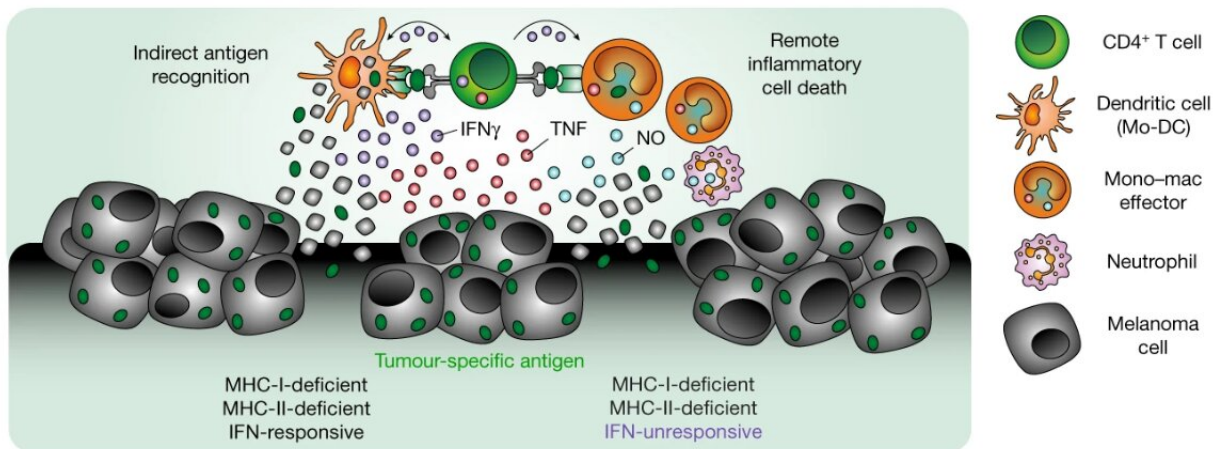


Researchers discover a new mechanism of cancer immune defense

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Graphical summary of inflammatory cell death induction of MHC-deficient and IFN-unresponsive tumors by CD4+ T cells in cooperation with iNOS-expressing myeloid cells. Credit: *Nature* (2023). DOI: 10.1038/s41586-023-06199-x

An interdisciplinary team of researchers at the Otto-von-Guericke University in Magdeburg has gained new insights into how inflammatory mediators of pathogen defense can remotely drive cancer cells into death—an important contribution to improving cancer immunotherapies.

Modern immunotherapies boost the body's own defenses against cancer. They activate killer T cells of the immune system that can specifically recognize and destroy [cancer cells](#). In many patients, however, cancer

cells adapt and become invisible to killer T cells so that the treatment is no longer effective. An interdisciplinary team of researchers from Magdeburg has now discovered a new mechanism that enables the immune system to also eliminate such invisible cancer cells. These findings open up new possibilities for the development of improved cancer immunotherapies. The results have now been published in the journal *Nature*.

"In our work, we have been looking for strategies to target such cancer cells that are 'invisible' to killer T cells. In doing so, we came across the special abilities of the so-called helper T cells," says Prof. Dr. Thomas Tüting, Professor of Dermatology at the University Hospital Magdeburg and leader of the study team.

Very few helper T cells are more effective than many killer T cells

Using an experimental cancer model, the researchers observed that a small number of helper T cells can eradicate advanced cancers as effectively as a much larger number of killer T cells. The helper T cells were also able to eliminate cancer cells that had become invisible to killer T cells.

Using cutting edge microscopy techniques to study [immune cells](#) in living [cancer tissue](#) revealed that helper T cells behave fundamentally differently from killer T cells. "Killer T cells penetrate into cancer tissues and interact directly with cancer cells, while helper T cells are mainly found at the edge of cancer tissues, where they exchange signals with other immune cells," says Prof. Dr. Andreas Müller from the Institute of Molecular and Clinical Immunology at the University of Magdeburg.

Helper T cells engage scavenger cells

Further investigations revealed that helper T cells secrete chemical mediators that attract scavenger cells of the immune system and induce them to support the destruction of cancer cells on their behalf. Together, these two [cell types](#) can effectively fight bacterial and viral infections. Their cooperation can also be exploited to mobilize the full arsenal of immune defense against cancer cells.

In search for the underlying mechanisms of action, the researchers found that the interaction between helper T cells and scavenger cells enhances their ability to release [inflammatory mediators](#) which act remotely to drive cancer cell death, as if they were infected by a pathogen. Exactly how this happens is still not completely understood, and the significance of this mechanism for [cancer immunotherapies](#) will have to be elucidated.

The research results reveal a mechanism for how the ability of the [immune system](#) to defend against the spread of pathogens in the body can be utilized to destroy cancer cells. Based on these findings, the researchers in Magdeburg are developing new strategies for cancer immunotherapy that are also effective in patients with cancers that have become invisible to killer T cells.

More information: Bastian Kruse et al, CD4+ T cell-induced inflammatory cell death controls immune-evasive tumours, *Nature* (2023). [DOI: 10.1038/s41586-023-06199-x](https://doi.org/10.1038/s41586-023-06199-x)

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