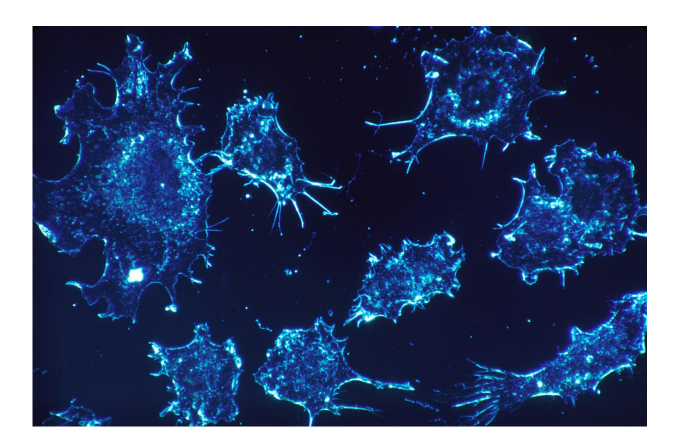


New study reveals how cancer manipulates our immune system to become harder to treat

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Many factors affect cancer treatment outcome, such as the size and location of the tumor, availability of effective treatments, and timing of intervention. But some cancers are so aggressive that outcome is poor, even after early diagnosis and chemotherapy. Researchers have focused



their attention on trying to understand what makes some cancers less treatable than others. Now, researchers at Children's Hospital Los Angeles reveal a mechanism by which some cancers trick healthy cells into protecting tumors.

Yves DeClerck, MD, of the Children's Center for Cancer and Blood Diseases and the Saban Research Institute at Children's Hospital Los Angeles, has dedicated his career to understanding how <u>cancer cells</u> interact with the surrounding normal tissue to escape the effects of therapy. Research has shown that tumors with high levels of a protein called Plasminogen Activator Inhibitor 1 (PAI-1) are more aggressive and are associated with poorer outcomes. In the new study, published November 20th in the journal *Cell Reports*, DeClerck's team demonstrated that cancer <u>cells</u> use PAI-1 to trick the body's immune system into supporting the cancer.

DeClerck and his team, led by postdoctoral research fellow Marta Kubala, Ph.D., characterized a relationship between tumors and the immune system. "In this study, we focused on the role of immune cells called macrophages and how PAI-1 affects their activity," explains Kubala. As important players in the immune system, macrophages find and destroy cancer cells or foreign invaders like bacteria. While macrophages are normally considered anti-cancer, DeClerck's team showed that PAI-1 pushes macrophages into an alternate, pro-cancer state (called M2) by recruiting common players in the immune system—IL-6 and STAT3—effectively signaling to the macrophages to support rather than attack tumor cells.

"A macrophage can either be a friend or an enemy to cancer cells," explains DeClerck, who is also a professor of pediatrics at the Keck School of Medicine of the University of Southern California. "The cancer communicates with the macrophages, telling them to become friendly. So, the macrophages change their behavior and support the



tumor." In altering the function of surrounding, healthy tissue, the cancer is better able to survive and spread.

The team around DeClerck also shows that cancer cells can use PAI-1 to promote movement of these pro-cancer M2 macrophages into the tumors, where they protect the cancer and repair any damage that chemotherapy may have inflicted. This symbolic one-two punch culminates in a stronger, more difficult-to-treat cancer.

DeClerck and his team have uncovered what appears to be a very common mechanism used by many cancers to commandeer part of the body's immune system. In order to investigate how broadly this cellular communication could impact cancer treatment, DeClerck's team studied the National Institute of Health's Cancer Genome Atlas, a library of genetic information from more than 11,000 patient samples, and found that many different cancers have this relationship. "We looked at patient data from neuroblastoma and breast, prostate, colon, and lung cancers. Every time we see higher levels of PAI-1, we see more evidence of procancer M2 macrophages," says DeClerck.

This new understanding of just how PAI-1 communicates with macrophages to change their activity has the potential to alter our approach to cancer treatment since these findings are applicable to most types of cancer. "It is clear that the <u>tumor</u> microenvironment, including cells of the <u>immune system</u>, is crucial in cancer development," explains Kubala.

While cancers vary widely in terms of location, treatment, and survival rates, their manipulation of macrophages represents a common thread, which is important for devising improved treatments for aggressive cancers. "Targeting PAI-1 could be beneficial in <u>cancer</u>," DeClerck says, "but much more work needs to be done." He cautions that the answer is not as simple as eliminating PAI-1, which is also made in healthy tissue



and is an important part of the blood clotting process. But these results, which uncover a complete pathway of communication between tumors and <u>macrophages</u>, lay the foundation for a promising avenue of research.

Provided by Children's Hospital Los Angeles

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