

New signaling role for key protein may contribute to wound healing, tumor growth

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A key protein may represent a new way to use the immune system to speed healing and counter inflammatory, infectious and autoimmune diseases, according to study led by researchers at the Icahn School of Medicine at Mount Sinai and published in the December issue of *Cell Reports*.

The current study results revolve around proteinases, enzymes that break down proteins as part of cellular life. Matrix metalloproteinases or MMPs specifically target the extracellular matrix, the non-cell, structural framework within tissues. Beyond that role, the new study found that one member of this family, MMP-2, has another signaling role related to the human immune system. It may shift a set of cells to become part of immune response that accelerates healing in some cases, but may worsen inflammatory disease in others.

Drug designers may be able to leverage the newfound MM-2 mechanisms to prevent the contribution of inflammatory signals to tumor growth and autoimmune diseases, or to promote wound healing.

"Our results show that MMP-2 uses a multitude of mechanisms to modulate the immune system," says the study's lead investigator Nina Bhardwaj, MD, PhD, Director of Immunotherapy, Tisch Cancer Center at Mount Sinai, Professor of Medicine, Hematology and Oncology, Icahn School of Medicine at Mount Sinai. "These data provides context to how this mechanism happens and could lead to novel treatments."



The backdrop for the study finding is the body's initial, vague reaction to any invading organism expands into a precise and massive counterattack. That expansion starts when a dendritic cell "swallows" a piece of any invader encountered, ferries it to the nearest lymph node and presents it there for notice by lymphocytes (B cells and T cells), the workhorses of the immune system. DCs exposed to MMP-2 send signals that change gene expression such that more of a protein called OX40L is made, according to the study authors. OX40L is a signaling protein that links into a receptor protein on the surface of <u>dendritic cells</u>, like a key fitting into a lock, with their interaction changing the shape of the receptor such that other signals get sent. Among these is one that causes more <u>immune cells</u> within a certain part of the <u>immune system</u>, TH2 cells, to be made.

Though vital to effective immune responses to worms, Th2 cells have a dark side, in some cases triggering B cells to produce antibodies that recognize the body's own cells as foreign and that destroy the kidneys of lupus patients and the joints of patients with rheumatoid arthritis. Shifting T cells in the Th2 path generally encourages inflammation.

If MMP-2 signaling leads to the presence of more Th2 T cells, it means Th2 cells will secrete more inflammatory TH2 cytokines like TNFalpha, and interleukins 4 and 13, which fight parasite infections, but that can also contribute to <u>autoimmune diseases</u> if present in too large amounts. MMP-2 is also over-expressed in tumors, where it promotes cancer progression. This may in part be due to its ability to skew T cells toward Th2 differentiation, a process called type-2 polarization.

Th2 <u>cells</u> also contribute to wound healing because they produce cytokines, which fight infection and ultimately repair tissue. The tissue repair functions of MMP-2's may be due to the ability to drive Th2 immune response.

"MMP-2-mediated signaling either through direct injection or through



mimics could promote wound healing but these proposed functions will require further investigation in disease-specific models," says Dr. Bhardwaj. "Further discovery of new MMP-2 functions could lead to new reagents of inflammatory responses."

Provided by The Mount Sinai Hospital

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