For a recent CSHL Cancer Center study, Adjunct Professor Mikala Egeblad (now a Bloomberg Distinguished Professor with Johns Hopkins University) and postdoc Xue-Yan He (now Assistant Professor of Cell Biology & Physiology at Washington University School of Medicine in St. Louis) teamed with CSHL Professor Linda Van Aelst. Above: lung cancer metastasis in a mouse that underwent experiments designed to simulate the stress that cancer patients experience. Credit: Egeblad lab/Cold Spring Harbor Laboratory

Stress is inevitable. But too much of it can be terrible for our health. Chronic stress can increase our risk for heart disease and strokes. It may
also help cancer spread. How this works has remained a mystery—a challenge for cancer care.

Xue-Yan He, a former postdoc in Cold Spring Harbor Laboratory (CSHL) Adjunct Professor Mikala Egeblad's lab, says, "Stress is something we cannot really avoid in cancer patients. You can imagine if you are diagnosed, you cannot stop thinking about the disease or insurance or family. So it is very important to understand how stress works on us."

Now, He and Egeblad may have reached a breakthrough in understanding exactly that. The work has been published in Cancer Cell.

Working with CSHL Professor Linda Van Aelst, they discovered that stress causes certain white blood cells called neutrophils to form sticky web-like structures that make body tissues more susceptible to metastasis. The finding could point to new treatment strategies that stop cancer's spread before it starts.

The team arrived at their discovery by mimicking chronic stress in mice with cancer. They first removed tumors that had been growing in mice's breasts and spreading cancer cells to their lungs. Next, they exposed the mice to stress. What He observed was shocking.

"She saw this scary increase in metastatic lesions in these animals. It was up to a fourfold increase in metastasis," Egeblad recalls.
Cancer spread faster and more furiously in stressed mice (middle column) than in a control group (left column). By comparison, cancer cells in stressed mice treated with an enzyme called DNase I (right column) were largely non-proliferating, and the treatment caused a significant reduction in stress-induced metastasis. Credit: Egeblad lab/Cold Spring Harbor Laboratory

The team found that stress hormones called glucocorticoids acted on the neutrophils. These "stressed" neutrophils formed spider-web-like structures called NETs (neutrophil extracellular traps). NETs form when neutrophils expel DNA. Normally, they can defend us against invading microorganisms. However, in cancer, NETs create a metastasis-friendly environment.

To confirm that stress triggers NET formation, leading to increased metastasis, He performed three tests. First, she removed neutrophils from the mice using antibodies. Next, she injected a NET-destroying
drug into the animals. Lastly, she used mice whose neutrophils couldn't respond to glucocorticoids. Each test achieved similar results. "The stressed mice no longer developed more metastasis," He says.

Notably, the team found that chronic stress caused NET formation to modify lung tissue even in mice without cancer. "It's almost preparing your tissue for getting cancer," Egeblad explains.

To Van Aelst, the implication, though startling, is clear. "Reducing stress should be a component of cancer treatment and prevention," she says.

The team also speculates that future drugs preventing NET formation could benefit patients whose cancer hasn't yet metastasized. Such new treatments could slow or stop cancer's spread, offering much-needed relief.

**More information:** Chronic stress increases metastasis via neutrophil-mediated changes to the microenvironment, *Cancer Cell* (2024). [DOI: 10.1016/j.ccell.2024.01.013](https://www.cell.com/cancer-cell/fulltext/S1535-6108(24)00037-0)

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