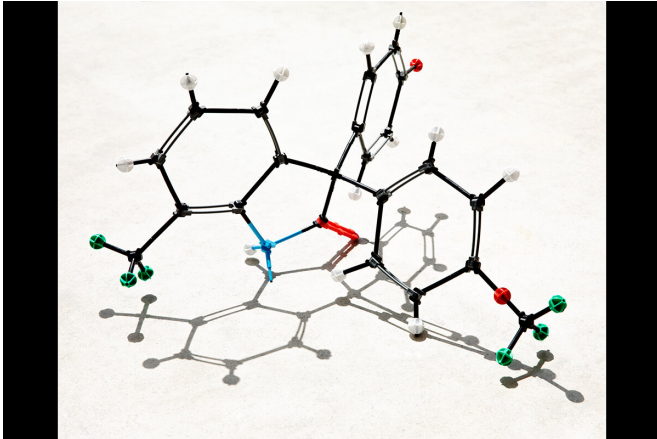


New approach eradicates breast cancer in mice

21 July 2021



Researchers discovered a small molecule, ErSO, that eradicates breast cancers in mice by targeting a pathway that protects cancer cells. Credit: L. Brian Stauffer

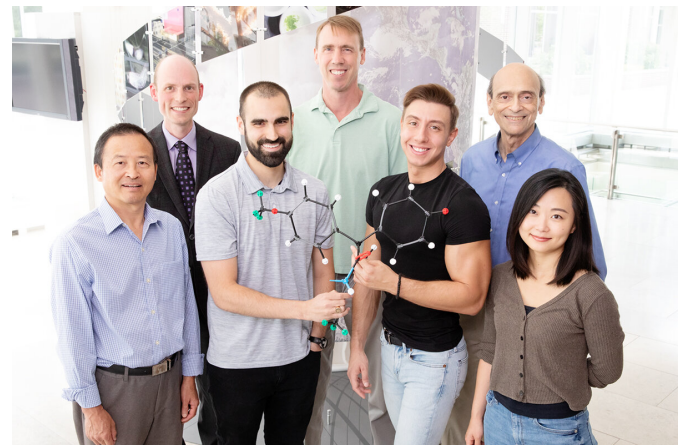
A new approach to treating breast cancer kills 95-100% of cancer cells in mouse models of human estrogen-receptor-positive breast cancers and their metastases in bone, brain, liver and lungs. The newly developed drug, called ErSO, quickly shrinks even large tumors to undetectable levels.

Led by scientists at the University of Illinois Urbana-Champaign, the research team reports the findings in the journal *Science Translational Medicine*.

"Even when a few [breast cancer cells](#) do survive, enabling tumors to regrow over several months, the tumors that regrow remain completely sensitive to retreatment with ErSO," said U. of I. biochemistry professor David Shapiro, who led the research with Illinois chemistry professor Paul Hergenrother. "It is striking that ErSO caused the rapid destruction of most lung, bone and liver metastases and dramatic shrinkage of brain metastases, since tumors that have spread to

other sites in the body are responsible for most breast [cancer](#) deaths," Shapiro said.

The activity of ErSO depends on a protein called the estrogen receptor, which is present in a high percentage of breast tumors. When ErSO binds to the estrogen receptor, it upregulates a [cellular pathway](#) that prepares cancer cells for rapid growth and protects them from stress. This pathway, called the anticipatory Unfolded Protein Response, or a-UPR, spurs the production of proteins that protect the cell from harm.



Illinois researchers on the study include, from front left, research scientist Chengjian Mao and graduate students Matthew Boudreau, Darjan Duraki and Ji Eun Kim. In the back row, from left, are molecular and integrative physiology professor Erik Nelson, chemistry professor Paul Hergenrother and biochemistry professor David Shapiro. Credit: L. Brian Stauffer

"The a-UPR is already on, but running at a low level, in many breast cancer cells," Shapiro said. "It turns out that this pathway shields cancer cells from being killed off by [anti-cancer drugs](#)."

Shapiro and former U. of I. medical scholar Neal

Andruska first identified the a-UPR pathway in 2014 and reported the development of a compound that pushed the a-UPR pathway into overdrive to selectively kill estrogen-receptor-containing breast cancer cells.

"Because this pathway is already on in cancer cells, it's easy for us to overactivate it, to switch the breast cancer cells into lethal mode," said graduate student Darjan Duraki, who shares first-author status on the new report with graduate student Matthew Boudreau.

While the original compound prevented breast cancer cells from growing, it did not rapidly kill them, and it had undesirable side effects. For the new research, Shapiro and Hergenrother worked together on a search for a much more potent small molecule that would target the a-UPR. Their analysis led to the discovery of ErSO, a small molecule that had powerful anticancer properties without detectable side effects in mice, further tests revealed.

"This anticipatory UPR is estrogen-receptor dependent," Hergenrother said. "The unique thing about this compound is that it doesn't touch cells that lack the estrogen receptor, and it doesn't affect healthy cells—whether or not they have an estrogen receptor. But it's super-potent against estrogen-receptor-positive cancer [cells](#)."

ErSO is nothing like the drugs that are commonly used to treat estrogen-receptor-positive cancers, Shapiro said.

"This is not another version of tamoxifen or fulvestrant, which are therapeutically used to block estrogen signaling in breast cancer," he said. Even though it binds to the same receptor that estrogen binds, it targets a different site on the estrogen receptor and attacks a protective cellular pathway that is already turned on in [cancer cells](#), he said.

"Since about 75% of breast cancers are estrogen-receptor positive, ErSO has potential against the most common form of breast cancer," Boudreau said. "The amount of estrogen receptor needed for ErSO to target a breast cancer is very low, so ErSO may also work against some breast cancers not

traditionally considered to be ER-positive." Further studies in mice showed that exposure to the drug had no effect on their reproductive development. And the compound was well tolerated in mice, rats and dogs given doses much higher than required for therapeutic efficacy, the researchers found.

ErSO also worked quickly, even against advanced, human-derived breast cancer tumors in mice, the researchers report. Often within a week of exposure to ErSO, advanced human-derived breast cancers in mice shrank to undetectable levels.

"Many of these breast cancers shrink by more than 99% in just three days," Shapiro said. "ErSO is fast-acting and its effects on breast cancers in mice are large and dramatic."

The pharmaceutical company Bayer AG has [licensed the new drug](#) and will explore its potential for further study in [human clinical trials](#) targeting [estrogen](#)-receptor-positive [breast](#) cancers, the researchers said. The researchers will next explore whether ErSO is effective against other types of cancers that contain [estrogen receptor](#).

More information: M.W. Boudreau et al., "A small-molecule activator of the unfolded protein response eradicates human breast tumors in mice," *Science Translational Medicine* (2021). stm.sciencemag.org/lookup/doi/10.1126/scitranslmed.abf1383

Provided by University of Illinois at Urbana-Champaign

APA citation: New approach eradicates breast cancer in mice (2021, July 21) retrieved 27 July 2021 from <https://medicalxpress.com/news/2021-07-approach-eradicates-breast-cancer-mice.html>

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