

Possible new principle for cancer therapy

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A study published in *Science Translational Medicine* shows that small molecules that specifically inhibit an important selenium-containing enzyme may be useful in combating cancer. When researchers at Karolinska Institutet in Sweden treated cancer in mice using these molecules, they observed rapid tumoricidal effects. Researchers now



hope that this new principle for cancer treatment will eventually be developed for use in humans.

Humans need the chemical element selenium for good health. The selenium-containing enzyme thioredoxin reductase 1 (TrxR1) can be used to support the growth of cells and to protect them from harmful forms of oxygen radicals that cause oxidative stress. Selenium intake has long been connected with <u>cancer</u>, although the correlation between selenium and cancer growth is extremely complex and not fully understood.

Raised levels of TrxR1 can be seen in several forms of cancer and are linked to worse prognoses in head and neck, lung and breast cancers. Researchers therefore analysed almost 400,000 <u>molecules</u> in the search for new and more specific TrxR1 inhibitors than those previously available. They discovered three molecules that met their search criteria. Sure enough, when tested, these same molecules also proved to be active as cancer medicines. The researchers were able to treat over 60 types of cancer cells under laboratory conditions. Normal cells were, however, much less sensitive to these molecules.

"This effectiveness against cancer may be a result of cancer cells' seemingly greater sensitivity to <u>oxidative stress</u> when compared to <u>normal cells</u>, which in turn can be utilised in cancer therapy," says Professor Elias Arnér, Department of Medical Biochemistry and Biophysics at Karolinska Institutet, who led the study.

Researchers also found rapid tumoricidal effects when treating head and neck or breast cancers in mice, without obvious side-effects. The new molecules have yet to be tested on humans, but it is already known that several different cancer medicines currently in use, including cisplatin and melfalan, inhibit TrxR1. Thus far, whether or not this inhibition of the enzyme is important to the effectiveness of these medicines has



remained unknown, but this study suggests that this might well be the case. Researchers now intend to further develop the new TrxR1 inhibitors to offer a new form of cancer therapy.

"My hope is that we will be able to develop new treatments, effective against multiple forms of cancer but with few side-effects. This seems to work in mouse models and we are therefore hopeful that this principle for treatment can be developed for humans, even if this will require many years of further research," says Elias Arnér.

More information: "Irreversible inhibition of cytosolic thioredoxin reductase 1 as a mechanistic basis for anticancer therapy". William C Stafford, Xiaoxiao Peng, Maria Hägg Olofsson, Xiaonan Zhang, Diane K Luci, Li Lu, Qing Cheng, Lionel Trésaugues, Thomas S Dexheimer, Nathan P Coussens, Martin Augsten, Hanna-Stina Martinsson Ahlzén, Owe Orwar, Arne Östman, Sharon Stone-Elander, David J Maloney, Ajit Jadhav, Anton Simeonov, Stig Linder och Elias S.J. Arnér. *Science Translational Medicine*, online 14 February 2018. <u>stm.sciencemag.org/lookup/doi/... scitranslmed.aaf7444</u>

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