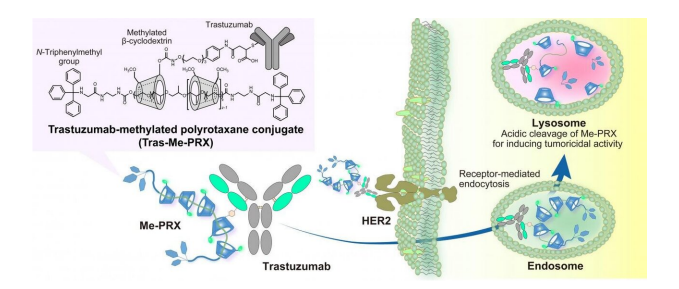


## **Opening an autophagy window as the apoptosis door starts to close**

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The conjugate of Trastuzumab, a monoclonal antibody against HER2 expressed in various malignant tumors, and methylated β-cyclodextrin-threaded polyrotaxane (Me-PRX) is designed for specific delivery of Me-PRX in HER2-expressing tumor cells. Because Me-PRX is known to induce endoplasmic reticulum stress-related autophagic cell death, Trastuzumab-Me-PRX conjugates are regarded as a new class of antibody-drug conjugates that would contribute to the chemotherapy of cancers through the induction of autophagic cell death. Credit: Department of Organic Biomaterials,TMDU

Many people across the globe are working hard to get the better of cancer; however cancer is always working too. Cancer cells can become resistant to the methods that have been adopted to kill them, so



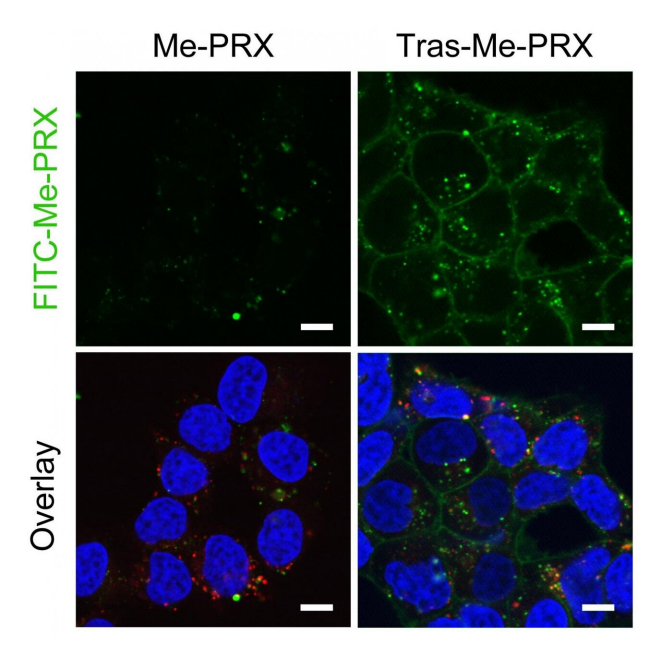
identifying drugs that act in different ways is part of the push to outsmart this ubiquitous disease. TMDU researchers have engineered a material that can identify cancer cells and mount an attack that they are not yet ready for. Their findings are published in the *Journal of Materials Chemistry B*.

Most of the anti-tumor drugs available work by inducing a cell <u>death</u> process known as apoptosis, and unfortunately cancer <u>cells</u> are developing resistance to drugs that work by this mechanism. However, other modes of cell death are known and focusing on alternative mechanisms is one way for researchers to stay a step ahead.

The TMDU researchers previously reported a supermolecule (Me-PRX) that acts as a drug by inducing autophagic cell death. The structure of Me-PRX can be thought of as lots of rings threaded onto a piece of string. These rings are kept on the string by adding stoppers to each end. However, the stoppers are specifically designed so that the rings are released at the acidic pH inside the cell. The release of the rings inside the cell causes endoplasmic reticulum stress, which leads to autophagic cell death.

Me-PRX has significant potential, but until now lacked a way of targeting the specific <u>cancer</u> cells requiring treatment. It was also too small to stay in the bloodstream for extended periods. In their recent work, the researchers have attached their supramolecular drug to the Trastuzumab antibody, which is able to identify the HER2 gene that is overexpressed by many tumor cells.





Fluorescence microscopic images of HER2-positive BT-474 cells treated with FITC-Me-PRX (green) and FITC-Tras-Me-PRX (green) for 24 h (Scale bars: 10 µm). The nuclei (blue) and endosomes/lysosomes (red) are stained with Hoechst 33342 and LysoTracker Red, respectively. The cellular uptake of Me-PRX in HER2-positive tumor cells is remarkably enhanced by the conjugation with Trastuzumab, leading to potentiating the autophagic cell death of Me-PRXs. Credit: Department of Organic Biomaterials,TMDU



The size increase brought about by forming this antibody-drug conjugate (ADC) also stops it being filtered out by the kidneys, giving it more time to act. And being larger helps the ADC to be passively taken up by tumors.

"We compared our Trastuzumab-modified Me-PRX (Tras-Me-PRX) with Me-PRX alone and Me-PRX modified with an antibody unable to target HER2-overexpressing <u>cancer cells</u>," study lead author Atsushi Tamura explains. "And we found that the binding of Trastuzumab to HER2 played an important role in getting Me-PRX into cells."

The researchers also found that lower concentrations of Tras-Me-PRX were needed to have an impact on the number of live healthy cells than those required for Me-PRX alone. This shows that the delivery of Tras-Me-PRX led to enhanced autophagic cell death.

"Our demonstration of targeted delivery and cell death using an ADC is a significant step towards an autophagic cell death <u>drug</u>," study corresponding author Nobuhiko Yui explains. "Many ADCs have been approved for clinical use; therefore we are hopeful that future in vivo investigations with our material will ultimately lead to a tangible option in the clinic."

**More information:** Kei Nishida et al, An antibody–supermolecule conjugate for tumor-specific targeting of tumoricidal methylated  $\beta$ -cyclodextrin-threaded polyrotaxanes, *Journal of Materials Chemistry B* (2020). DOI: 10.1039/D0TB00575D

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