

Breaking the stroma barrier: Study shows a new way to hit cancer with radiation

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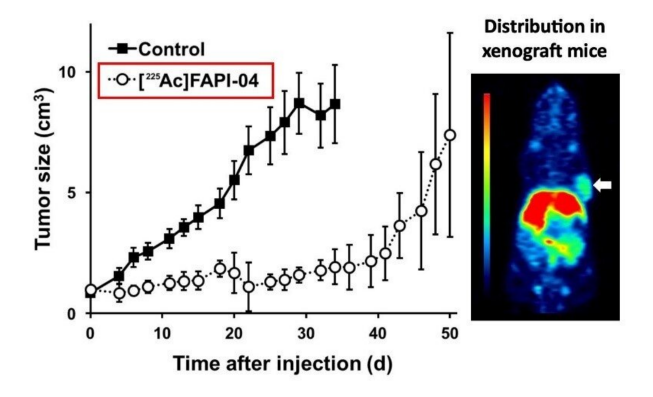


Fig.1. (a) Human pancreatic cancer cells were grafted onto mice, which were then given targeted alpha radiation therapy (²²⁵Ac-FAPI-04). Compared with mice given a control treatment (black squares), mice given ²²⁵Ac-FAPI-04 (open circles) showed a marked decrease in tumor growth. (b) Mice were injected with targeted positron radiation (⁶⁴Cu-FAPI-04) to enable positron emission tomography (PET) imaging. The imaging shown here indicates uptake of ⁶⁴Cu-FAPI-04 into the pancreatic tumor (arrow). Credit: Osaka University



Pancreatic cancer is one of the leading causes of cancer-related deaths worldwide. Patients with pancreatic cancer often receive chemotherapy or radiation therapy, which are not always effective and can have toxic side effects. In a collaborative research between Osaka University and the University of Heidelberg, researchers are exploring a new method of treatment that brings powerful yet targeted radiation therapy directly to cancer tissue. The method may offer an effective treatment for pancreatic cancer with fewer side effects in surrounding organs.

Chemotherapy and <u>radiation therapy</u> typically work by stopping cancer cells from dividing. Side effects often come about because the treatment also stops healthy cells from dividing and thriving.

"With traditional anti-cancer therapies, there's a tradeoff between efficacy against cancer cells and off-target effects in non-cancerous cells," explains Dr. Tadashi Watabe, lead author of the study and a researcher in <u>nuclear medicine</u> and tracer kinetics at Osaka University Graduate School of Medicine. "We're focused on finding ways to rebalance this tradeoff in radiotherapy, by increasing the dose of radiation delivered to <u>cancer cells</u> while keeping it localized to those cells as much as possible."

One way this can be done is by making a radioactive "homing" molecule that, when injected into a patient, travels straight to the site of a tumor to deliver its cancer-killing payload. The research team did precisely that—in this case, using a molecule that goes directly to pancreatic tumors.

"We used a molecule with an isotope that emits <u>alpha particles</u>," says Dr. Watabe. "These particles travel a shorter distance than the beta particles emitted by traditional isotopes, which means their off-target effect is limited. Our molecule binds to a protein called FAP, which is found almost exclusively on stroma cells that surround pancreatic tumors and



various types of cancers. Together, this allows us to target a high-energy dose of radiation directly to the site of the tumor."

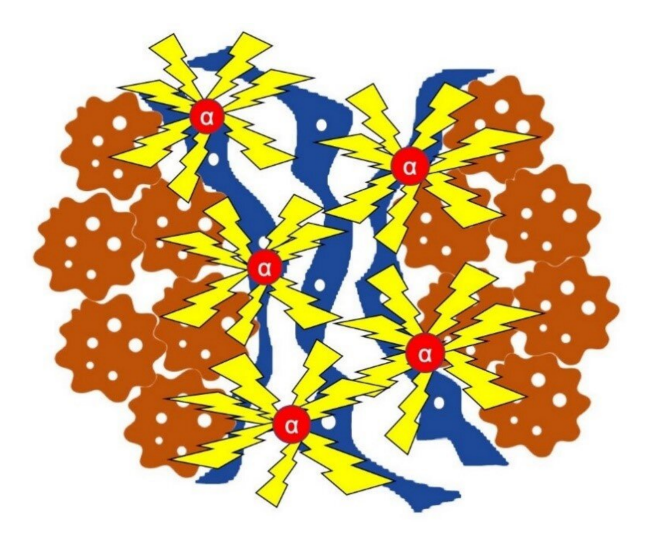


Fig.2. Schematic depiction of the mode of action for targeted alpha radiation in skin cell-derived cancers. Cancer cells (brown) are associated with several other cell types in the tumor microenvironment, such as fibroblasts (blue). ²²⁵Ac-FAPI-04 (red) is localized to the cancer through its binding to fibroblasts, resulting in targeted irradiation of the tumor. Credit: Department of Nuclear Medicine and Tracer Kinetics, Osaka University Graduate School of Medicine



The stroma is part of a unique environment that encases solid tumors, acting as a kind of barrier. Treatments for pancreatic cancer are often stymied by this barrier but, remarkably, this doesn't seem to be the case with the new experimental treatment.

In a study published in the *Journal of Nuclear Medicine*, the team offers a proof of concept that their technique can act as a powerful cancer treatment. They injected their FAP-binding molecule into mice that had human pancreatic cancer grafted under their skin. The result? A dramatic decrease in tumor growth with no apparent effect on the body weight of the mice, the latter indicating that the treatment likely had few toxic side effects.

"We are very encouraged by these initial results," Dr. Watabe concludes. "We think the approach has enormous therapeutic potential, particularly for patients with <u>pancreatic</u> cancer who've exhausted their other treatment options. What's especially exciting is that our method of targeting the stroma can in principle work against many other types of <u>cancer</u>. We think this could represent a new path forward in radiation therapy."

The article, "Theranostics targeting fibroblast activation protein in the tumor stroma: ⁶⁴Cu- and ²²⁵Ac-labelled FAPI-04 in <u>pancreatic cancer</u> xenograft mouse models" was published in the *Journal of Nuclear Medicine*.

More information: TTadashi Watabe et al. Theranostics targeting fibroblast activation protein in the tumor stroma: 64Cu and 225Ac labelled FAPI-04 in pancreatic cancer xenograft mouse models, *Journal of Nuclear Medicine* (2019). DOI: 10.2967/jnumed.119.233122



Provided by Osaka University

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