

Scientists aim to understand why men with prostate cancer are at higher risk of Alzheimer's

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Dr. Qin Wang and Augusta University graduate student Mae Aida. Credit: Michael Holahan, Augusta University

Researchers at the Medical College of Georgia at Augusta University are



searching for a better way to understand why many men with prostate cancer end up with Alzheimer's disease, and whether it's the standard hormone therapy treatment or an overactive immune response that actually contributes to the problem.

The <u>hormone therapy</u>, <u>androgen deprivation therapy</u>, known as ADT, treats the cancer by reducing testosterone, which the cancer needs to grow. But androgen is a key regulator of amyloid metabolism and when it's removed from the equation, more amyloid is left to form the plaques that are a hallmark of Alzheimer's.

"We know that <u>prostate cancer</u> itself also largely affects men over age 65, which is a population that's already at a higher risk of Alzheimer's, simply due to their age," said Qin Wang, MD/Ph.D., Georgia Research Alliance Eminent Scholar in Neurodegeneration and inaugural director of the Program for Alzheimer's Therapeutic Discovery at MCG. "What is not largely understood is the role ADT may play in the context of cancer and Alzheimer's disease."

Wang and her research team suspected that the body's over-active immune response and subsequent inflammation may be at play.

To better understand, they created an <u>animal model</u> with Alzheimer's disease and cancer, then delivered the ADT for eight weeks, while monitoring androgen levels and tumor size. The study is <u>published</u> in the journal *Science Advances*.

"We wanted to first validate our model and also check and see if there were cognitive deficits in that group," Wang said.

They also developed other animal models. A so-called wild type (without Alzheimer's or cancer), a group with just Alzheimer's, and a group with just cancer that received ADT therapy—helped the team further



delineate this complex interplay. The researchers also regularly monitored changes in the blood looking for immune markers.

At the end of eight weeks, they also looked for increased amyloid in the brain, but "to our surprise, we didn't actually see a significant difference in the plaque load in any of the groups."

They did, however, note a high amount of reactivity in <u>glial cells</u> of the groups with just cancer and the groups treated with ADT. Glial cells are a part of the central nervous system and typically support nerve cells, helping them function properly. Their hyperactivity is a key indicator of inflammation in the brain, Wang said.

They also noted an increase in pro-inflammatory cytokines, small proteins that trigger an increase in inflammation throughout the body, and a decrease in anti-inflammatory cytokines, particularly in the animals with Alzheimer's and cancer that received ADT.

The team also looked at the animals' blood-brain barrier and noted that the same group had significant damage. Under normal circumstances, that barrier protects the brain from harmful substances while allowing essential molecules to pass through. "That would explain why there is so much more inflammation in that group," Wang and the team reported.

"The ADT treatment is actually making the blood-brain barrier more permeable. Then these immune cells, which are circulating in higher amounts because of the cancer and the subsequent treatment, can go into the brain—we call it immune cell infiltration—and cause inflammation, which is associated with cognitive decline."

To intervene and stop the immune cell infiltration, Wang and her team turned to a drug that is already on the market to treat multiple sclerosis and Crohn's disease. Natalizumab is a disease modifying therapy that



works by attaching to <u>immune cells</u> and preventing them from moving across the blood-brain barrier and entering the central nervous system.

Treating the mice who had cancer and Alzheimer's and cancer with a combination of ADT and natalizumab not only reduced the infiltration but subsequently improved the integrity of the <u>blood-brain barrier</u>.

"Then we looked at inflammation markers and those levels were reduced as well," Wang said. "We essentially reduced the pro-inflammatory cycle and improved cognitive function. We now know that it's not just about the amyloid plaques. The immune system's response is the contributing factor here."

Since the drug is already on the market, next steps could include a clinical trial in patients who are undergoing ADT for prostate cancer.

More information: Chao Zhang et al, Androgen deprivation therapy exacerbates Alzheimer's-associated cognitive decline via increased brain immune cell infiltration, *Science Advances* (2024). DOI: 10.1126/sciadv.adn8709

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