

DNA-based nanotechnology stimulates potent antitumor immune responses

September 10 2020



L-R: Wistar scientists Drs. David Weiner & Dan Kulp Credit: The Wistar Institute

Combining their expertise in protein engineering and synthetic DNA technology, scientists at The Wistar Institute successfully delivered



nanoparticle antitumor vaccines that stimulated robust CD8 T cell immunity and controlled melanoma growth in preclinical models. Study results were published online in Cancer Immunology Research, a journal of the American Association for Cancer Research, and support exploration of this immunotherapy approach for additional cancer types.

Nanovaccines consist of extremely small (nano) particles—similar in size to bacteria and viruses—used to display multiple copies of an antigen and able to elicit strong immune responses. The team previously reported on using DNA instructions to launch in vivo production of nanoparticle vaccines (DLnano-vaccines).

DLnano-vaccines assembled in the body produced stronger immune responses than protein based nanoparticle vaccines in an infectious disease setting, especially inducing CD8 T cell responses.

"We wanted to test DLnano-vaccines for <u>cancer</u> immunotherapy and obtain proof of concept that this platform could be successfully applied in the cancer field, thanks to its effectiveness at prompting CD8 T <u>cells</u> responses," said Daniel Kulp, Ph.D., associate professor in Wistar's Vaccine & Immunotherapy Center and co-corresponding author of the study, who specializes in nanotechnology and protein engineering for vaccine development.

Due to their ability to specifically kill malignant cells, CD8 T cells play a pivotal role in anticancer immunity, therefore engagement of these cells represents a necessary step for the success of anticancer vaccine approaches, although this type of immune response is typically difficult to achieve by vaccination with proteins or inactivated virus.

Researchers designed DLnano-vaccines displaying 60 copies of protein parts derived from the melanoma-specific antigens Trp2 and Gp100 and tested these in mouse models of melanoma, observing prolonged survival



that depended on CD8 T cell activation both in therapeutic and prophylactic settings.

"One of the advantages of synthetic DNA technologies over other methods is the versatility of the platforms," said Ziyang Xu, Ph.D., a recent doctoral graduate working at Wistar and the first author of the study. "DLnano-vaccines may be designed for various cancer targets and our study shows this is a promising strategy for cancer immunotherapy that may warrant further testing."

To elucidate the mechanism through which DLnano-vaccines activate CD8 T cells, the team studied the effects of the DNA-launched version of a previously described HIV nanoparticle vaccine (eOD-GT8-60mer). They observed that DLnano-vaccines administered via electroporation resulted in transient muscle cell apoptosis that attracted macrophage infiltration at the injection site, which in turn was instrumental to activate CD8 T cells.

DLnano-vaccines were developed using synthetic DNA technology in collaboration with the lab of David B. Weiner, Ph.D., Wistar executive vice president, director of the Vaccine & Immunotherapy Center, and the W.W. Smith Charitable Trust Professor in Cancer Research and also a co-senior author on the study.

More information: Ziyang Xu et al. A DNA-launched nanoparticle vaccine elicits CD8+ T-cell immunity to promote in vivo tumor control. *Cancer Immunology Research.* (2020) DOI: 10.1158/2326-6066.CIR-20-0061

Provided by The Wistar Institute



Citation: DNA-based nanotechnology stimulates potent antitumor immune responses (2020, September 10) retrieved 25 April 2024 from https://medicalxpress.com/news/2020-09-dna-based-nanotechnology-potent-antitumor-immune.html

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