

A novel cancer immunotherapy; vaccination against tumor vascular extracellular vimentin

Huijbers EJM, Van Beijnum JR, Van Loon K, Griffioen AW

Angiogenesis Laboratory, Cancer Center Amsterdam, Department of Medical Oncology, VU University Medical Center, Amsterdam UMC, Amsterdam, The Netherlands

Angiogenesis, the development of neovasculature, is required to sustain tumor growth and metastases of solid tumors. The tumor neovasculature expresses specific markers, which are selectively overexpressed in tumor endothelial cells compared to normal healthy adult endothelium and are therefore ideal targets for vaccination.

One of these tumor vascular markers, the cytoskeletal protein vimentin, was found to externalize from tumor endothelial cells, while expression in all other cells in the body is exclusively intracellular. Extracellular vimentin (eVim) is pro-angiogenic and functionally mimics vascular endothelial growth factor (VEGF) action, while concomitantly acting as inhibitor of leukocyte-endothelial interactions, thereby hampering leukocyte infiltration into the tumor. eVim is overexpressed in the vasculature of different solid tumors, but not present in normal healthy tissue. This makes extracellular vimentin (eVim) an exclusive target for therapy.

Using our iBoost conjugate vaccine technology for induction of efficient antibody responses against self-antigens, we were able to generate strong eVim specific humoral immune responses, resulting in inhibition of tumor growth in preclinical models without affecting the normal healthy vasculature. Furthermore, in an ongoing clinical study in client owned dogs with spontaneous bladder cancer our vaccine targeting eVim shows effective and safe inhibition of angiogenesis and tumor growth.

Targeting of extracellular vimentin by vaccination therefore presents a promising antiangiogenic immunotherapy strategy against cancer.