RESEARCH HIGHLIGHTS



Virus-based anticancer strategies have attracted increasing interest in recent years, and oncolytic virus-based vaccine therapies have reached late-stage clinical trials. Now, reporting in Nature Medicine, the group of Richard Vile present a virus-based anticancer strategy that works predominantly by an immune-enhancing rather than an oncolytic mechanism. By cloning a cDNA library derived from normal prostate into the vesicular stomatitis virus (VSV), cure rates of up to 80% were achieved in mouse models of prostate cancer.

This cDNA library — termed an altered self antigen and epitope library (ASEL) — resulted in presentation of a broad repertoire of low-affinity antigens, rather than targeting one cancer-specific tumourassociated antigen. Interestingly, this did not induce autoimmunity.

To investigate its anticancer effects, ASEL was tested in a mouse model of prostate cancer induced by the injection of mouse prostate tumour TC2 cells. Prostate-specific ASEL treatment of mice with 7-day-old TC2 tumours significantly enhanced survival compared to treatment with VSV-green fluorescent protein (GFP), and intravenous (i.v.) injection of ASEL proved to be more efficient than intratumoural injection.

After nine i.v. injections of ASEL into mice with TC2 tumours, a cure rate of 80% was achieved. Three i.v. injections only typically induced tumour regression but with aggressive recurrence, which prompted an investigation into whether vaccination with a library derived from the recurrent tumour cells (TC2Rs), termed immune-escape epitope libraries (IEELs), was effective in this setting. To avoid neutralization

of virus particles in mice that were previously immunized with ASEL, the IEEL-containing virus was pre-loaded into CD8⁺ T cells (T(IEEL)). Indeed, a sequential treatment with ASEL and T(IEEL) delayed or prevented the development of recurrences, thus demonstrating that tumours that evade the initial immune response can be re-targeted.

The authors further examined the nature of the immune responses initiated, and found that the induction of a CD4+ T helper 17 ($T_{\rm H}$ 17) cell response was crucial for the response against TC2 cells following the initial ASEL treatment. However, the sequential response to TC2R tumours was dependent on a $T_{\rm H}$ 1-like interferon- γ response mediated by CD8+ cells. They also found that responses against the xenogeneic (human) 'altered' self antigens were more potent than treatment with VSV carrying a cDNA library derived from mouse prostate.

These experiments demonstrate that cDNA libraries cloned into VSV can be used to induce antigen-specific immune responses against established tumours without eliciting autoimmunity. The authors further point out that the cDNA libraries can be readily constructed for off-the-shelf use and can be easily and systemically delivered via vectors that are amenable to production at a clinical grade.

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ORIGINAL RESEARCH PAPER Kottke, T. et al. Broad antigenic coverage induced by vaccination with virus-based cDNA libraries cures established tumours. Nature Med. **17**, 854–859 (2011)