

Basic Principles of Cancer Immunotherapy

Monica Thakar, MD

Medical College of Wisconsin









Disclosures

- Miltenyi Biotec, Contracted Research
- I will be discussing non-FDA approved indications during my presentation.









Why does the immune system fail to eliminate cancer?

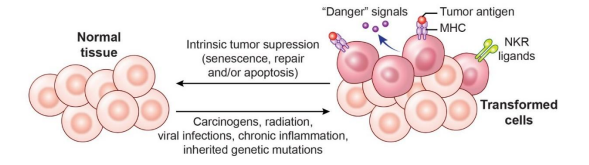
Cancer cells grow progressively in immunocompetent hosts without evidence of <u>T cell exhaustion</u> or <u>systemic anergy</u>.

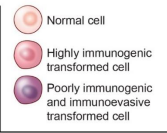










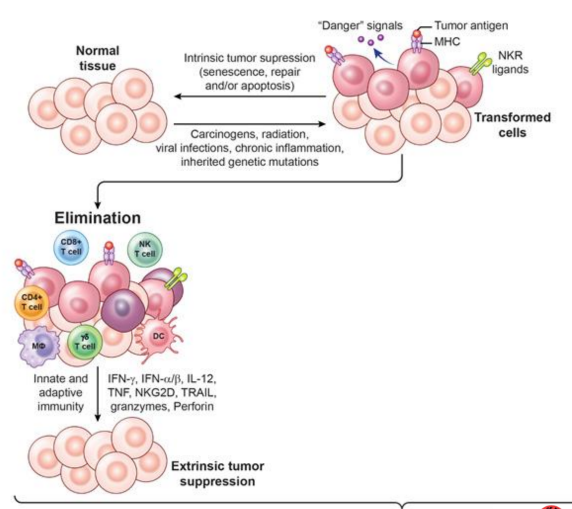


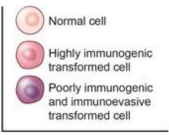










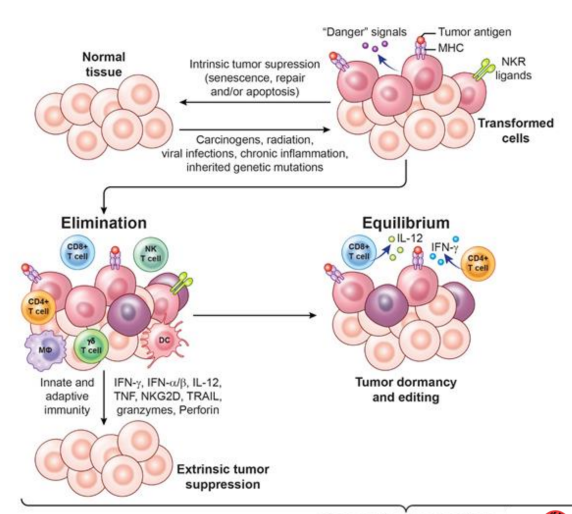


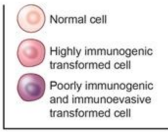










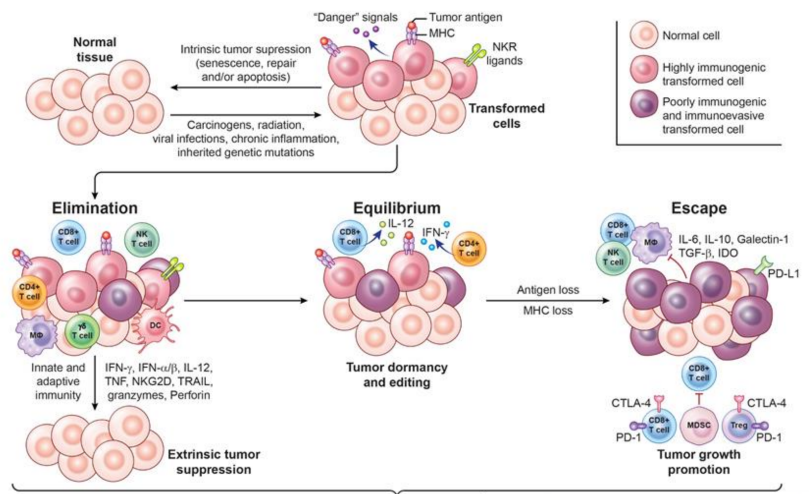












Cancer immunoediting









To exist, tumors must evolve mechanisms to locally disable and/or evade the immune system.

The goal of immunotherapy, then, is to restore the capacity of the immune system to recognize and reject cancer.



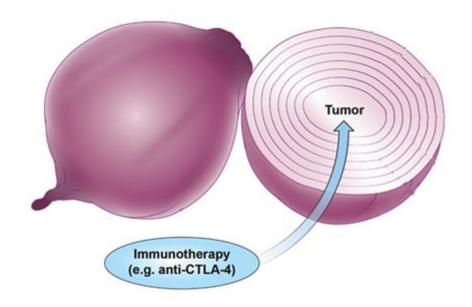






Multi-layered immunosuppression

- Tumors insulate themselves with dense layers of immunosuppressive stroma
- Overcoming the many layers of interconnected and often functionally redundant immune suppressive mechanisms represents a daunting challenge for tumorspecific T cells
- Immunotherapy can "peel back" the layers of local immune suppression, thereby restoring the capacity of T cells to eradicate the tumor



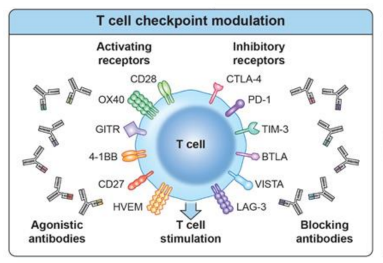


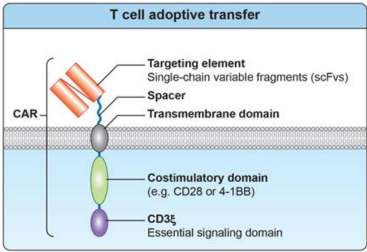


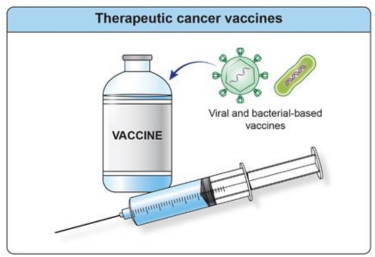


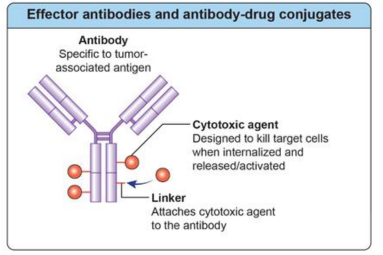


Types of immunotherapy











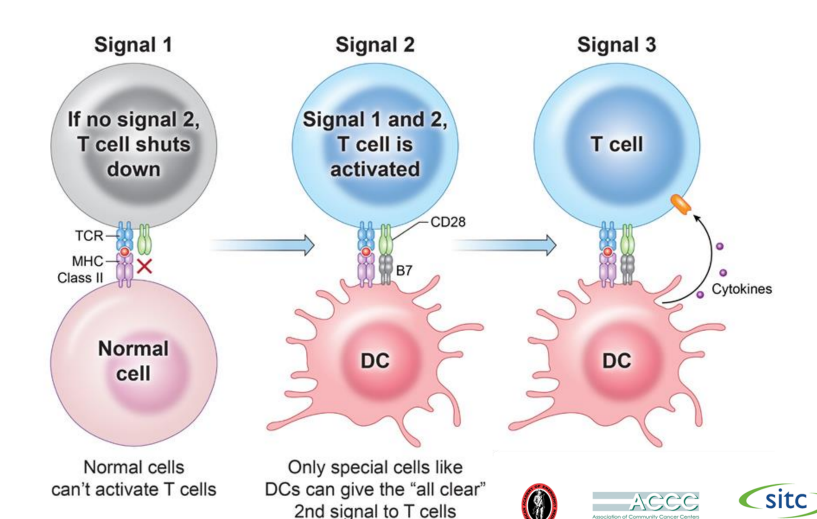






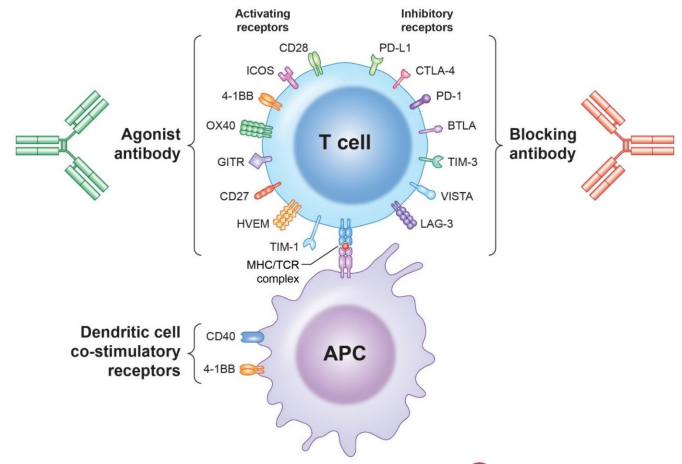
Three signals for antigen-specific T cell activation

Society for Immunotherapy of Cancer





T cell checkpoint modulation





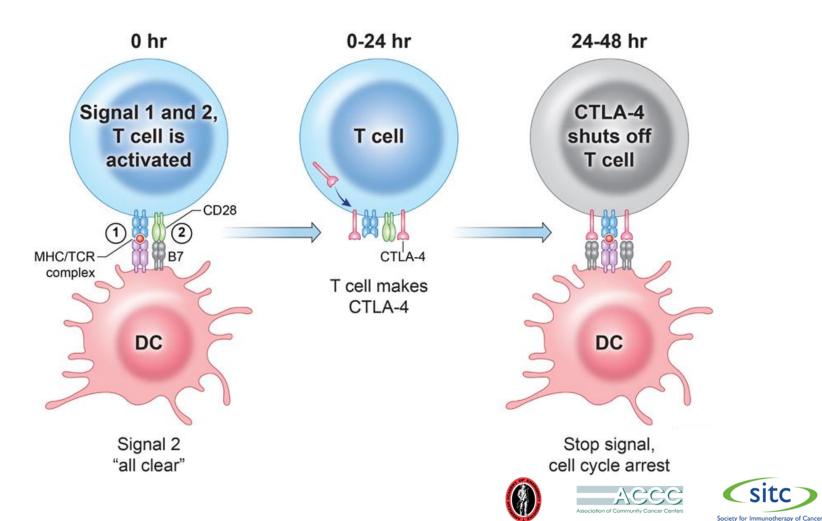






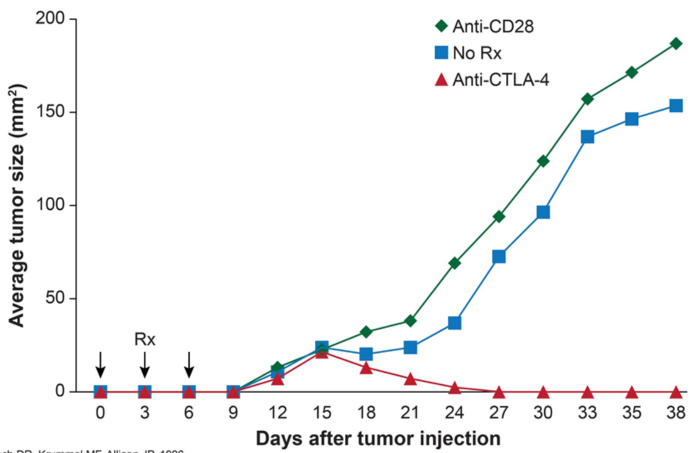
CTLA-4, a negative regulator of T cell activity limits the responsiveness of activated T cells

sitc





Anti-CTLA-4 induces regression of transplantable colon carcinoma



Leach DR, Krummel MF, Allison JP. 1996. Enhancement of antitumor immunity by CTLA-4 blockade. Science. 217(5256): 1734-6.

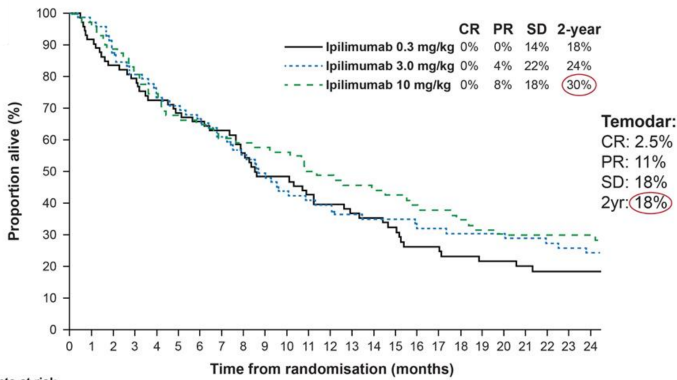








Ipilimumab (human anti-CTLA-4) was approved for the treatment of metastatic melanoma by FDA in 2010



Patients at risk

0.3 mg/kg 73 67 61 58 53 50 47 45 38 33 33 29 27 25 24 21 17 17 15 14 14 13 12 12 12 3.0 mg/kg 72 70 64 58 54 50 47 43 39 34 30 28 26 24 23 23 22 21 20 20 20 19 18 17 16 10 mg/kg 72 70 63 58 53 47 45 42 41 40 39 33 31 29 28 27 25 24 22 20 19 19 19 18 18

Wolchok et al. 2010. Lancet Oncol.

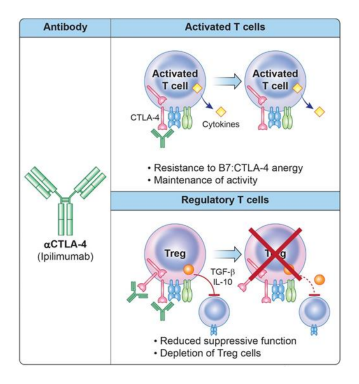








Which T cells are affected by ipilimumab (α CTLA-4)?



The efficacy and selectivity of anti-CTLA-4 therapy increase in patients who have higher percentages of activated tumor-specific T cells at the time of treatment



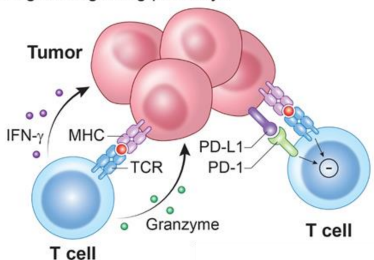


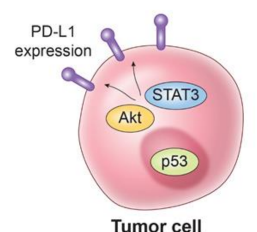




PD-1: PD-L1 inhibitory pathway

- PD-1 signaling promotes T cell tolerization by inhibiting downstream activation signals
- PD-1 expression is upregulated by activated and exhausted T cell populations
- T cell surface PD-1 receptor binds to and is activated by PD-L1 and PD-L2
- Many cells within the tumor microenvironment express PD-L1/PD-L2 allowing for the suppression of T cell activation
- Tumor PD-L1 expression is regulated via two general mechanisms:
 - 1. TIL production of IFN-γ
- 2. Oncogenic signaling pathways













To exist, tumors must evolve mechanisms to locally disable and/or evade the immune system.

The goal of T cell checkpoint blockade is to make T cell "off-switches" inaccessible to tumor cells, thus restoring tumor-specific immunity.

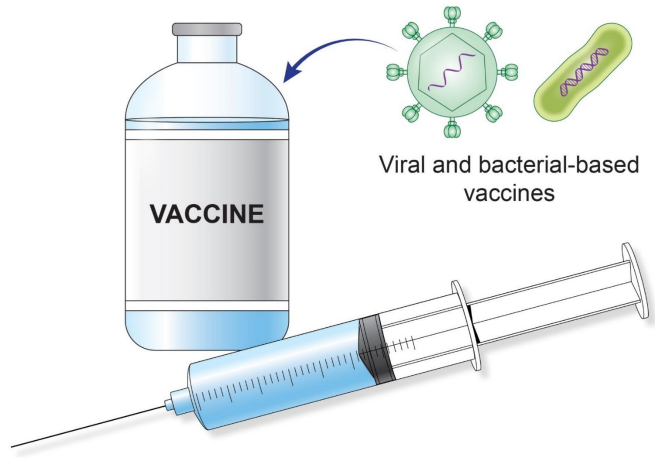








Therapeutic cancer vaccines



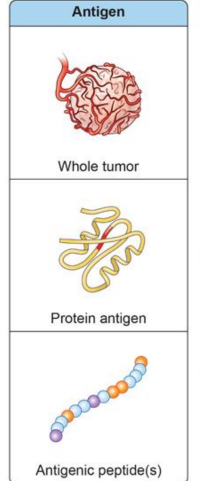


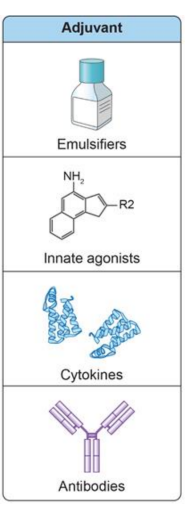


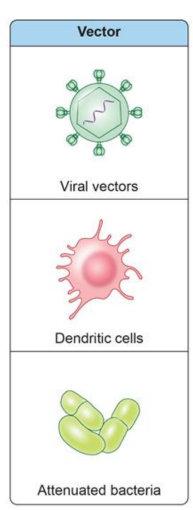


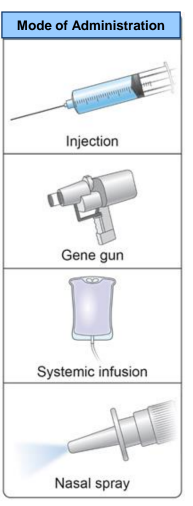


Components of a cancer vaccine











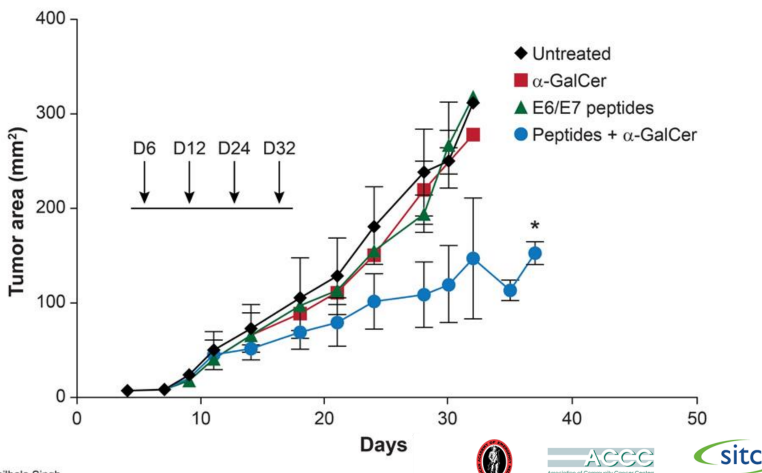






An intra-nasal HPV E6/E7: α -GalCer vaccine slows growth of TC-1 tumors

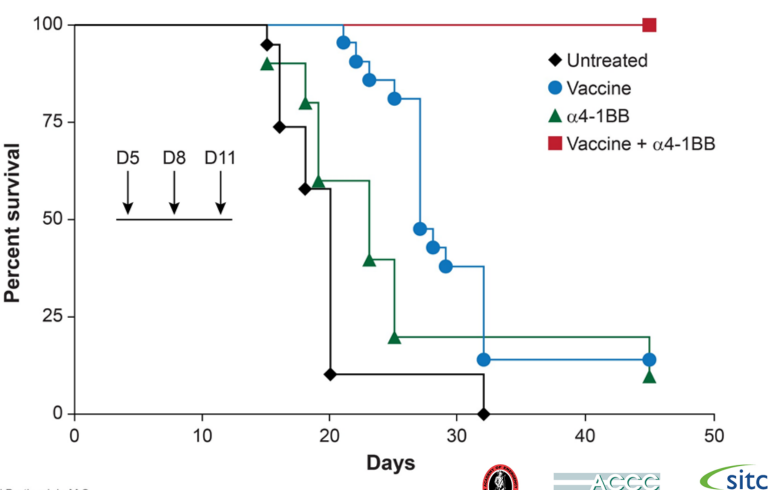
Society for Immunotherapy of Cancer





4-1BB agonist antibody and HPV E6/E7 vaccine synergize in curing TC-1 tumors

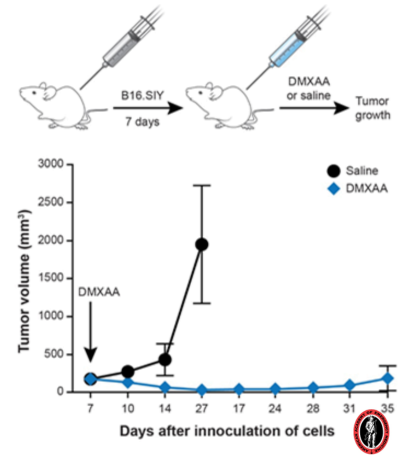
Society for Immunotherapy of Cancer





Intratumoral injection of innate immune agonists: The direct vaccination approach

Intratumoral DMXAA (mouse STING agonist) triggers rejection of B16 melanoma









To exist, tumors must evolve mechanisms to locally disable and/or evade the immune system.

The goal of therapeutic cancer vaccination is to increase the immunogenicity of tumor antigens which are poorly presented by the tumor in order to generate a high frequency of tumor-specific T cells.

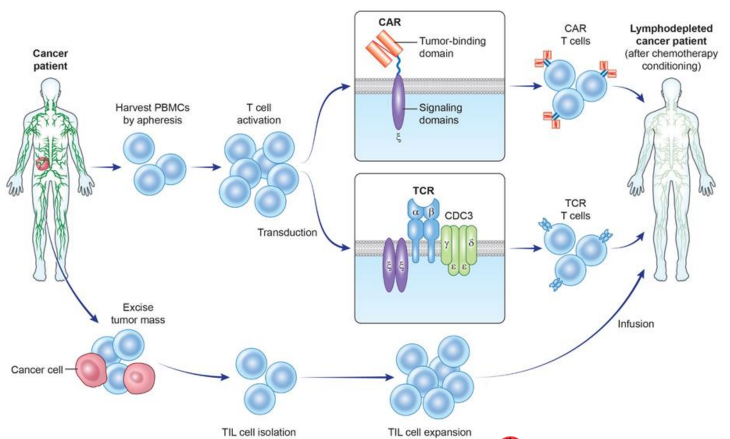








Adoptive T cell therapy can involve engineered (CAR, TCAR) or patient-derived (TIL, PBMC) T cells



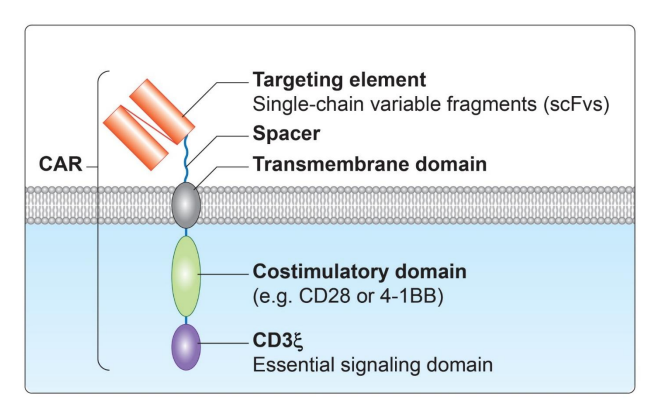








T cell adoptive transfer



CARs, TIL, TCR, PBMCs

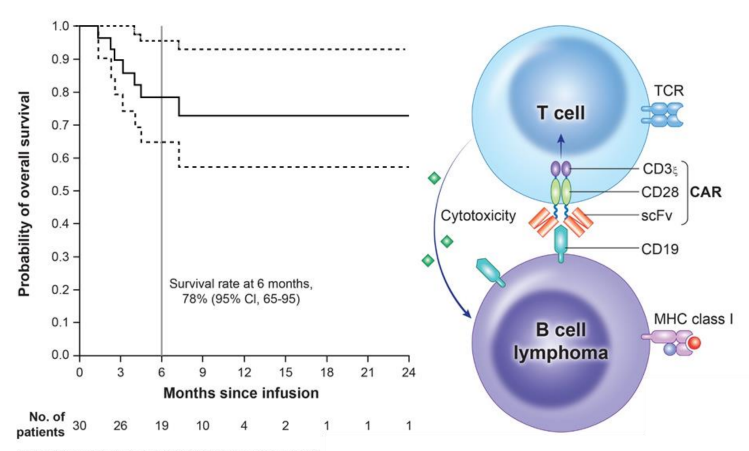








Effective treatment of relapsed B cell ALL with CD19 CAR T cell therapy



Maude S, Frey N, Shaw P, Aplenc R, Barrett D, Bunin N, Chew A, Gonzalez V, Zheng Z, Lacey S, et al. 2014. Chimeric antigen receptor T cells for sustained remissions in leukemia. The New England Journal of Medicine. 374(10): 988.









To exist, tumors must evolve mechanisms to locally disable and/or evade the immune system.

The goal of T cell adoptive transfer is to win the numbers game and overwhelm the tumor with a higher frequency of tumor-specific T cells than it is capable of suppressing.





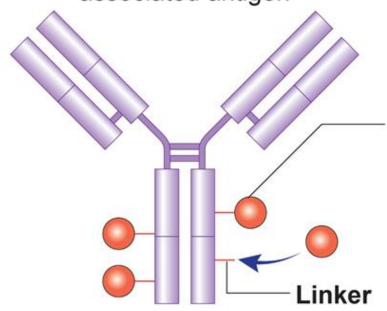




Effector antibodies and antibody-drug conjugates (ADCs)

Antibody

Specific to tumorassociated antigen



Radiolabeled or Cytotoxic agent

Designed to kill target cells when internalized and released or activated

Attaches cytotoxic agent to the antibody





Key ADC / antibody principles

- Specificity: The more tumor specific the target antigen is, the higher the agent can be dosed without limiting toxicity
- Internalization: The target tumor surface protein must internalize to deliver the toxin - it should do so frequently and to a suitable endosomal compartment.
- Stability: The toxin must remain inert and tethered to the antibody until it is delivered to its target cell.

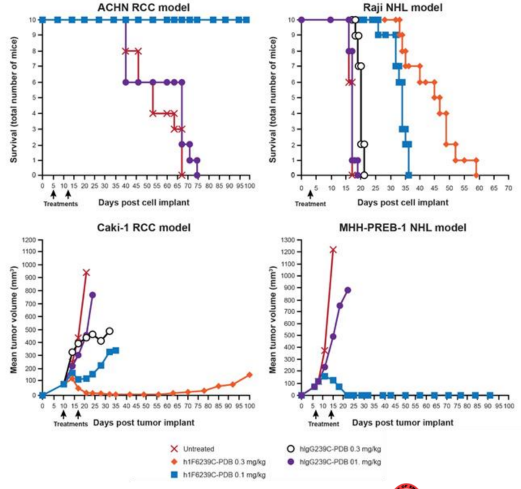








SGN-70A in the clinic for NHL and RCC



Jeffrey SC et al. 2013. Bioconjug Chem. 24(7): 1256-63









To exist, tumors must evolve mechanisms to locally disable and/or evade the immune system.

The goal of effector antibodies is to utilize the exquisite sensitivity of antibodies to specifically target and kill tumor cells using mechanisms which are difficult to evade or suppress.









of tumor and mobilization of antitumor effectors

Augmentation of tumor-specific T cells

Removal of barriers to immune rejection



- Chemotherapy
- Radiation
- ACT (CARs, TCR transfer)
- Co-inhibitory blockade
- Co-stimulatory activation
- Activation of APCs
 - Innate immune recognition
- Tumor vascular resistance
- Desmoplastic stroma
- Hypoxic microenvironments



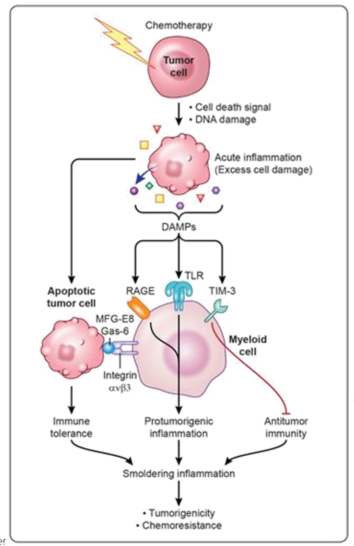






different perspective on chemotherapy

Immunogenic versus non-immunogenic cell death





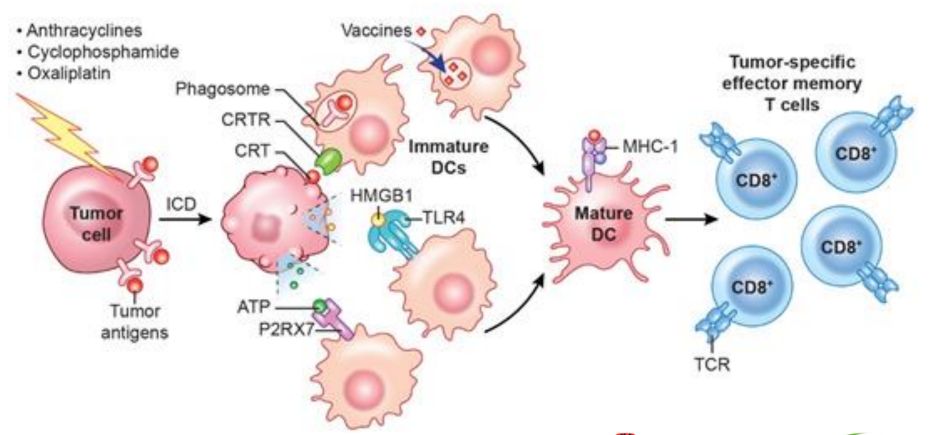




ADVANCES IN Cancer

different perspective on chemotherapy

Immunogenic versus non-immunogenic cell death





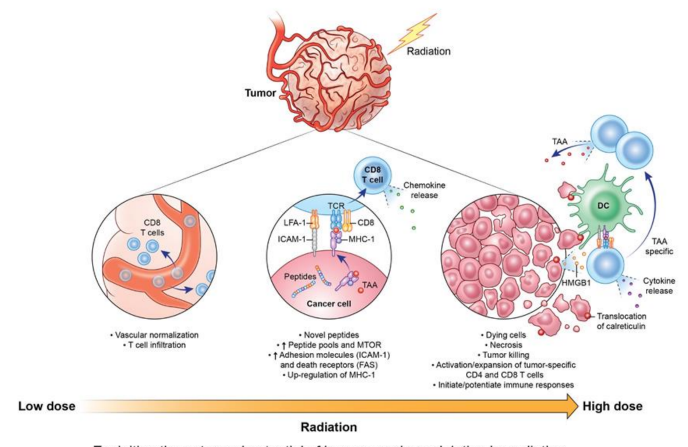






Radiation Therapy:

A potent adjuvant for tumor immunity



Exploiting the untapped potential of immunogenic modulation by radiation in combination with immunotherapy for the treatment of cancer

http://www.ncbi.nlm.nih.gov/pubmed/18777956









Abscopal effect

- Localized treatment of a tumor causes/evokes reduction of distant ones
- Immunotherapy + radiotherapy with targeted immunomodulators and immune checkpoint blockade is intended to elicit the abscopal effect.

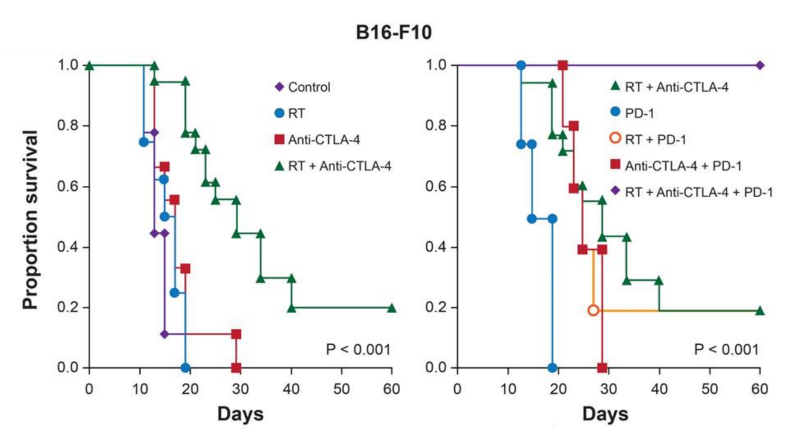








Radiotherapy synergizes with blockade of CTLA-4 and PD-1 to cure melanoma lung metastases



Victor CT, Rech A, Maity A, Rengan R, Pauken K, Stelekati E, Benci J, Xu B, Dada H, Odorizzi P, et al. 2015. Radiation and dual checkpoint blockade activate non-redundant immune mechanisms in cancer. Nature. 520: 373-377.







Why combination immunotherapy is the future?

More consistent benefit for a larger percentage of patients with a wide range of cancer types

