

## Basic Principles of Cancer Immunotherapy

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#### Disclosures

- Consulting Fees:
  - Novartis

• I will not be discussing non-FDA approved indications during my presentation.











# The Premise of Cancer Immunotherapy

 Normally, the immune system eliminates damaged cells, including precancerous and cancer cells

• To escape, tumors evolve mechanisms to locally disable the immune system.

The goal of immunotherapy is to restore the capacity of the immune system to recognize and eliminate cancer.











# Two major mechanisms of tumor immune escape

- Render the immune response dysfunctional: cytotoxic (CD8+) T cells often become dysfunctional or exhausted during chronic stimulation (chronic viral responses or responses against tumors). To enhance T cell dysfunction, the tumor microenvironment upregulates a suite of suppressive molecules.
- Avoiding an immune response: A state in which the tumor remains invisible to the immune system. Many features of tumors can result in immune exclusion/avoidance including lack of antigens (T cells don't "see" anything on the tumor) or active immune repellents.





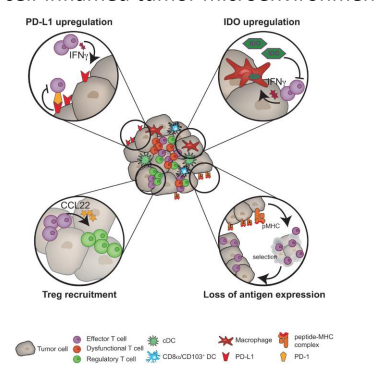




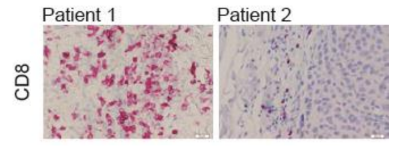


#### Immune evasion

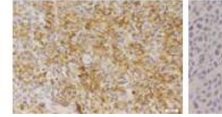
#### T cell-inflamed tumor microenvironment



#### T cells

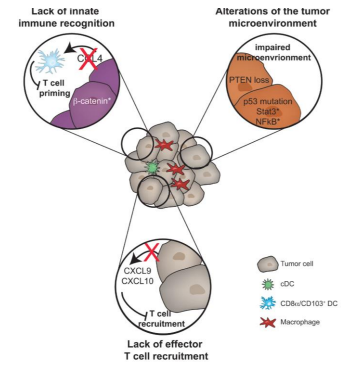


#### **Immune suppression**



PD-L1

#### Non-T cell-inflamed tumor microenvironment







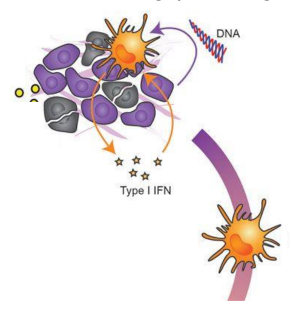






# Initiation of an anti-tumor immune response

Innate immune sensing (i.e. Sting activation)



APC maturation &

Transport to lymph node





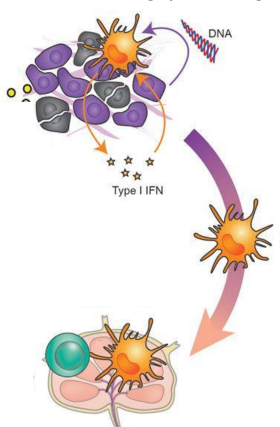






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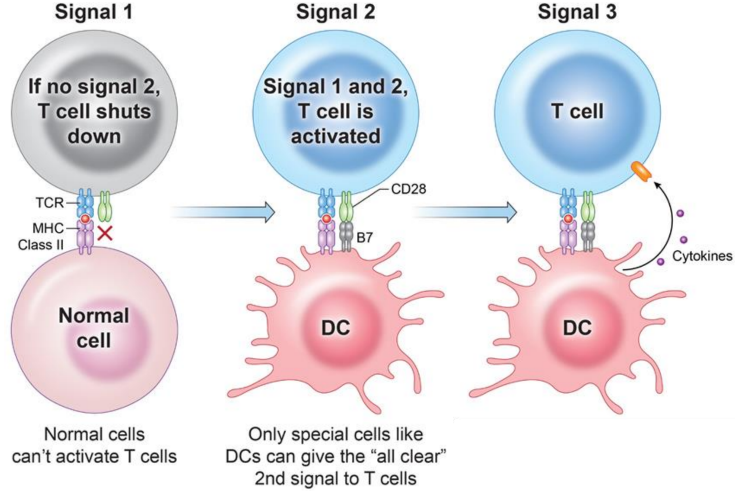








### Antigen-specific T cell Activation







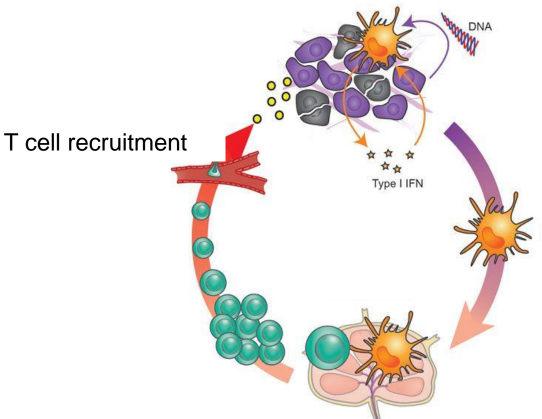






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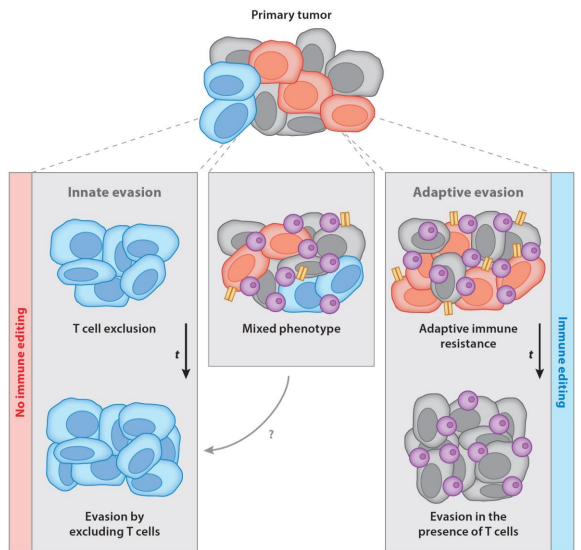


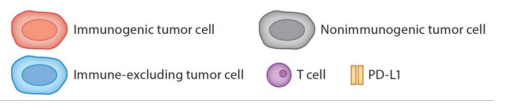






#### Immune evasion occurs over time









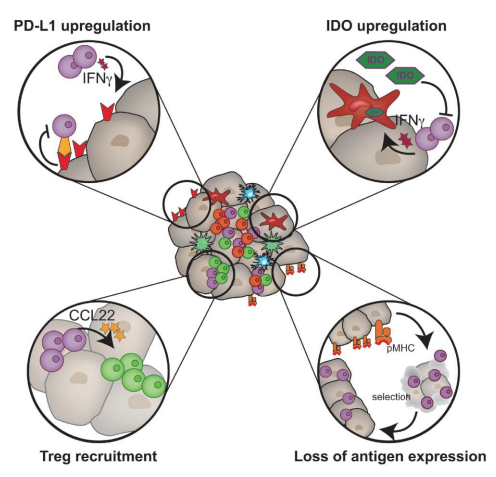






# T cell inflamed tumor microenvironment is immune suppressive

#### T cell-inflamed tumor microenvironment



T cell-inflamed tumors escape by suppressing T cell function









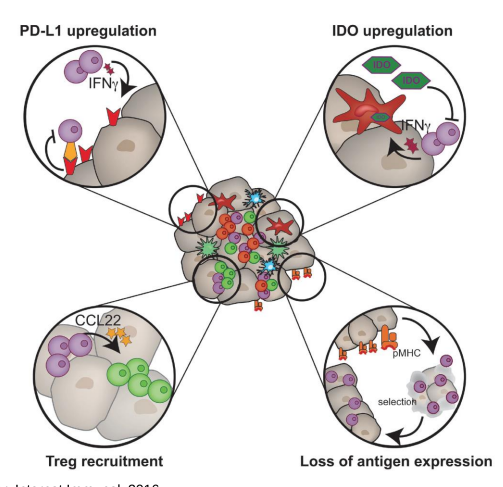


peptide-MHC



# T cell inflamed tumor microenvironment is immune suppressive

#### T cell-inflamed tumor microenvironment



T cell-inflamed tumors escape by suppressing T cell function

Non-T cell-inflamed tumors are a result of a malfunctioning cancer immune cycle













### Types of Immunotherapy

- Checkpoint blockade immunotherapy
- Cancer vaccines
- Adoptive cell transfer
- Effector antibodies
- Innate immune activation









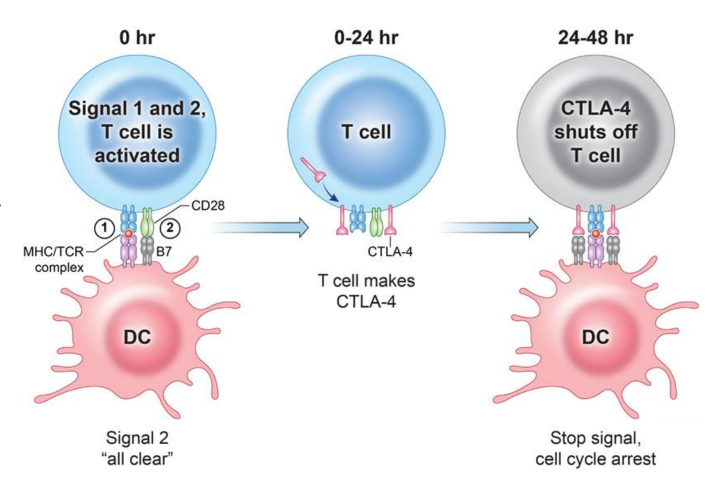


### The CTLA-4 Checkpoint

<u>Cytotoxic T-Lymphocyte</u> <u>Associated Protein 4</u>

Up-regulated in response to T cell activation

Limits positive stimulation by competition











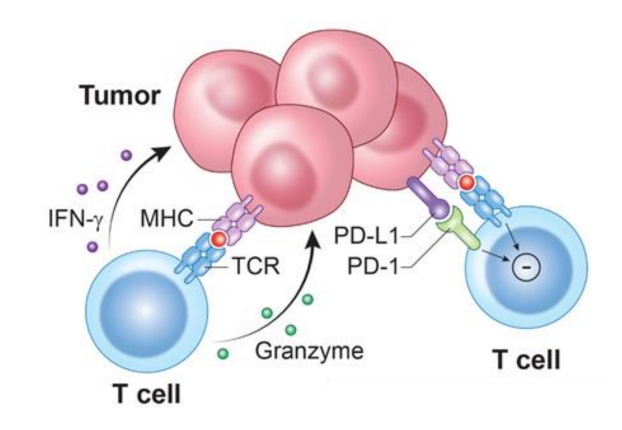


### The PD-1/PD-L1 Checkpoint

#### Programmed Death 1

Up-regulated in response to T cell activation

Ligands PD-L1 and PD-L2 are up-regulated following inflammation (IFN $\gamma$ )







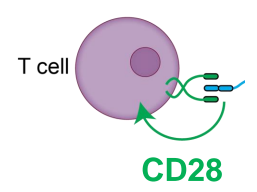






# Checkpoint blockade therapy unleashes the "brakes" on T cells

#### **Activation**



Goal: to reduce immune inhibitory signals and/or enhance stimulatory signals to allow T cells to regain effector functions.



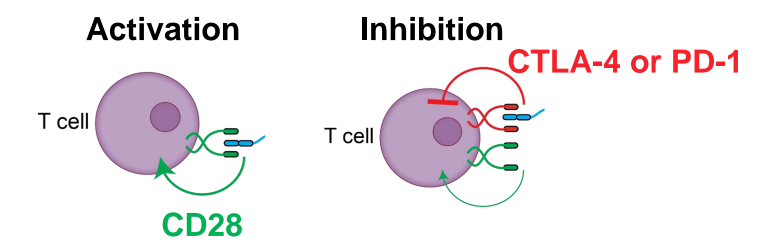








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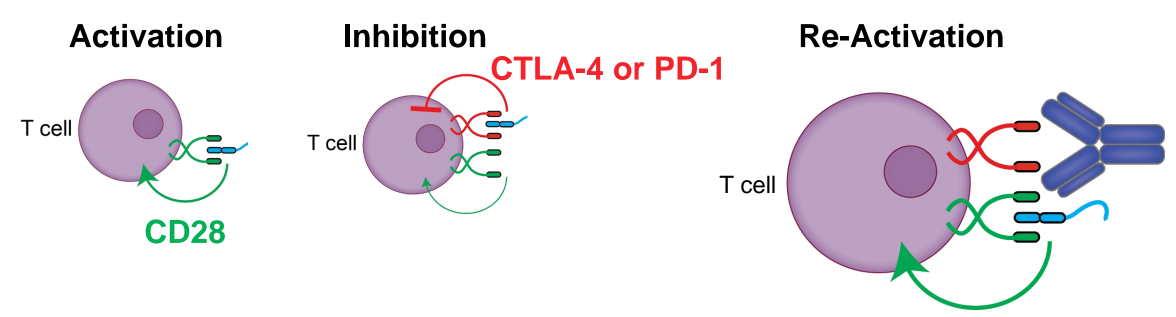








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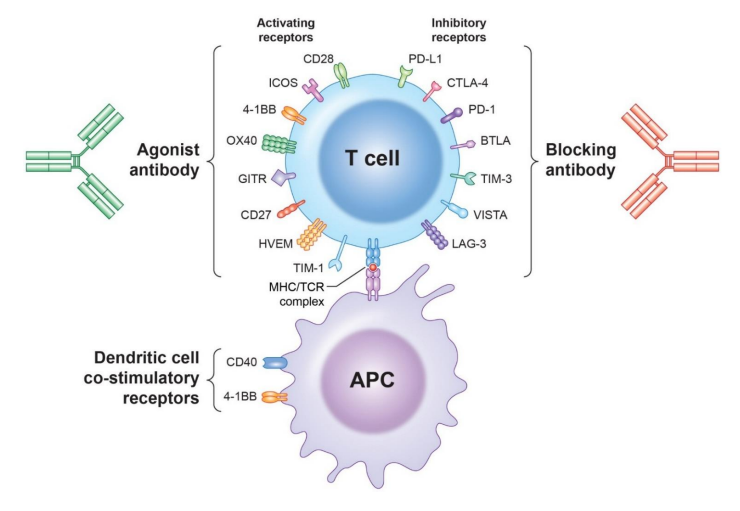




### T Cell Checkpoint Modulation

 First generation of checkpoint modulation: blocking inhibitory checkpoints

 Second generation of checkpoint modulation: activating stimulatory checkpoints







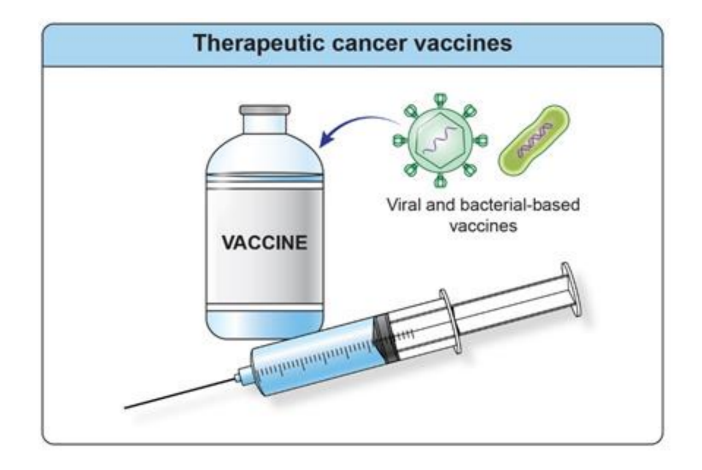






#### Therapeutic Cancer Vaccines

Goal: to increase the immunogenicity of tumor antigens in order to generate a high frequency of tumor-specific T cells.







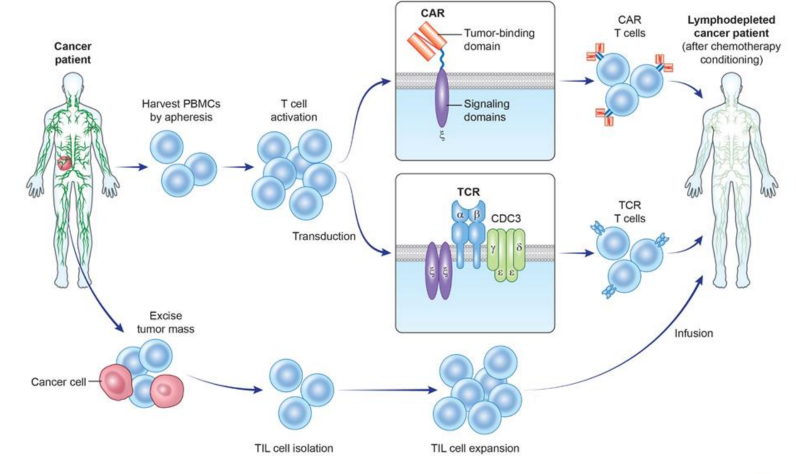






### **Adoptive Cell Therapy**

Goal: overwhelm the tumor with a higher frequency of tumor-specific immune cells and/or engineer immune cells to target cancer.







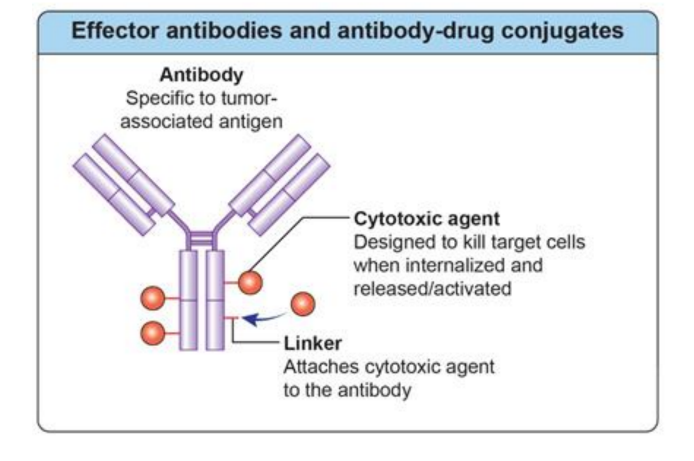






## Effector Antibodies and Antibody-Drug Conjugates (ADCs)

Goal: specifically target and kill tumor cells using innate mechanisms which are difficult to evade or suppress and/or through delivery of cytotoxic agents







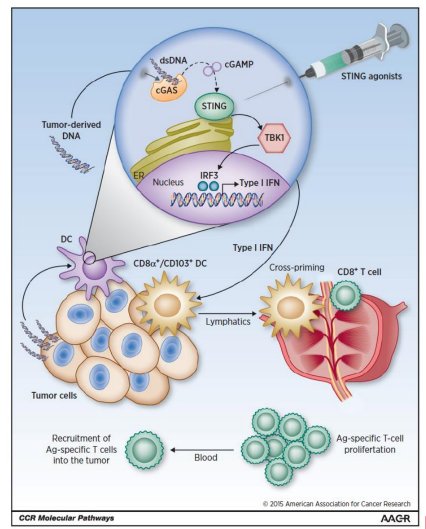






#### Innate immune activation

Goal: enhance innate immune sensing by providing stimulatory agents (frequently into the tumor itself)



Agents:
Sting agonists
TLR agonists
Immunogenic RNA



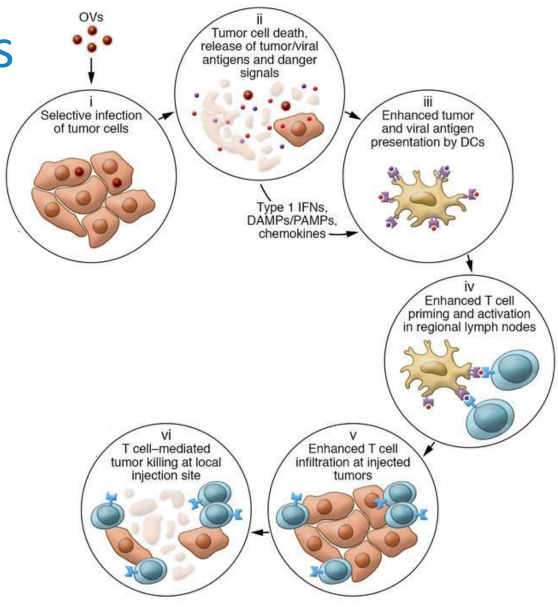






Oncolytic Viruses

Goal: specifically target and kill tumor cells through viral replication AND release innate immune activators and tumor antigens







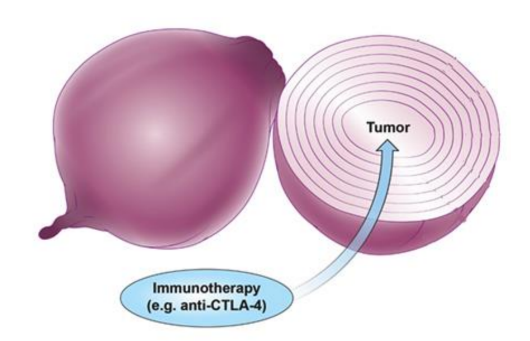






### Multi-layered Immunosuppression

- Tumors insulate themselves with dense layers of immune-suppression
- Overcoming the many layers of interconnected and often functionally redundant immune suppressive mechanisms represents a daunting challenge for tumor-specific T cells
- Immunotherapy can "peel back" the layers of local immune suppression
- Combination therapy might be needed to overcome all layers







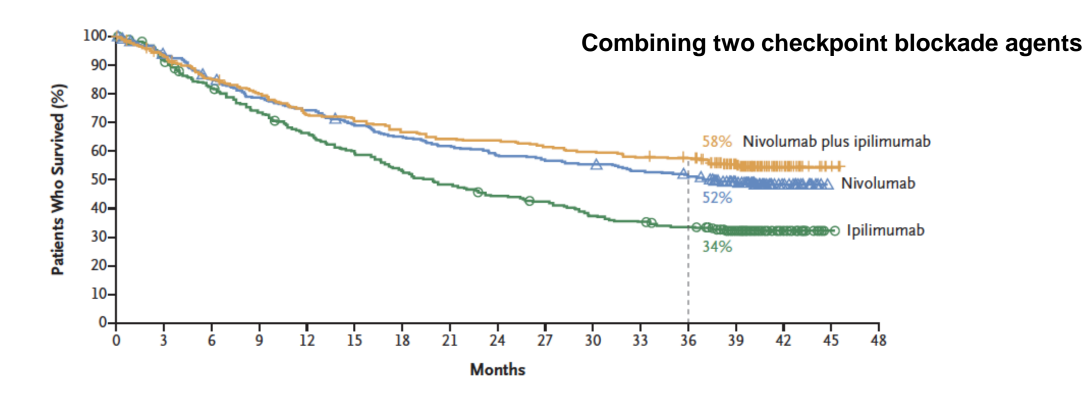






### **Combination Immunotherapies**

Dual CTLA-4 and PD-1 inhibition





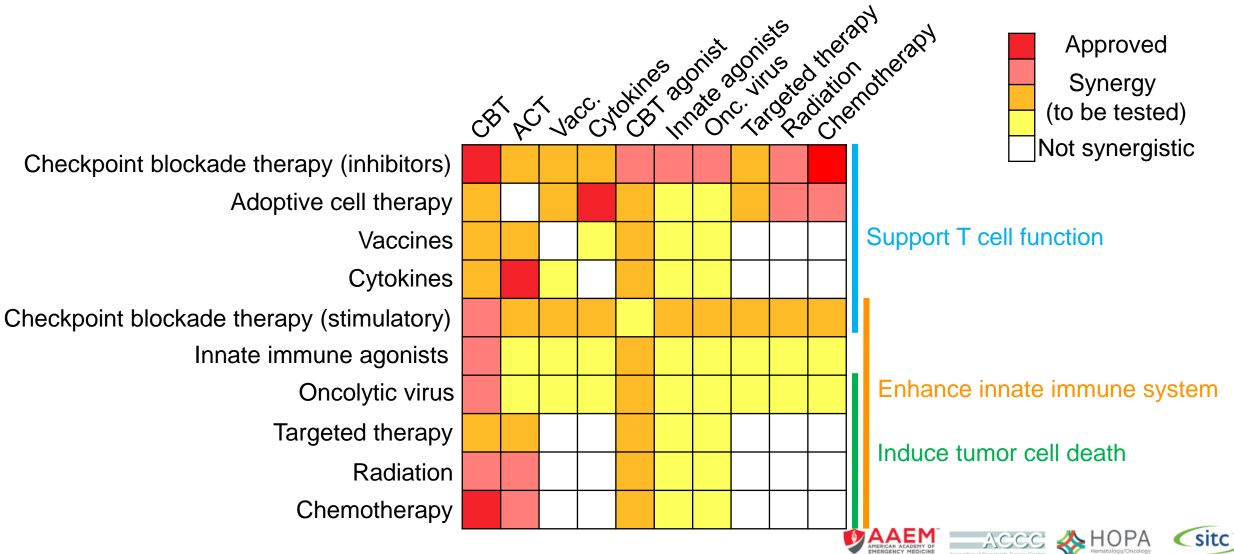






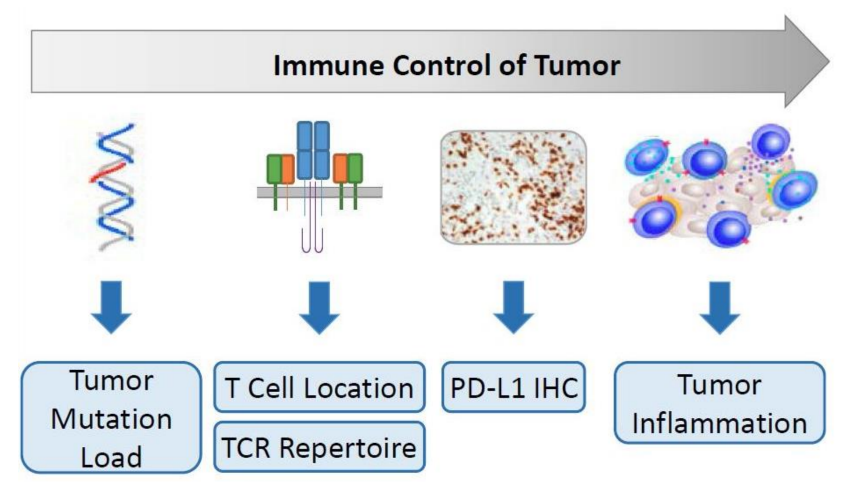


### Combination Immunotherapies





## **Immunotherapy Biomarkers**





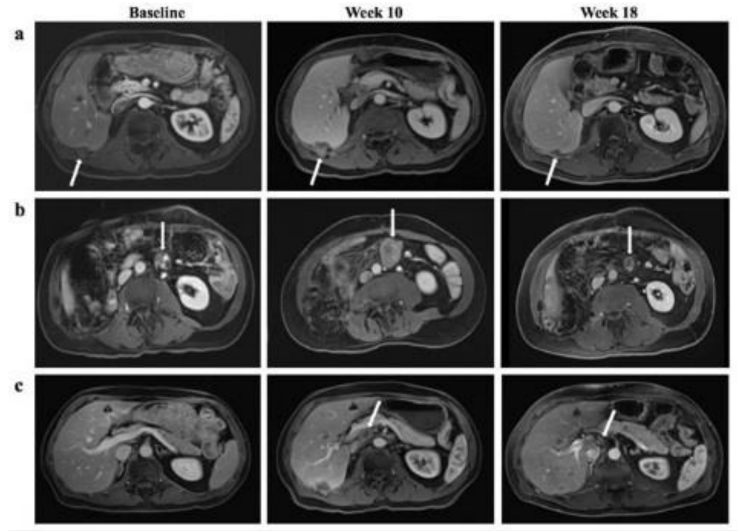








### Assessment of response





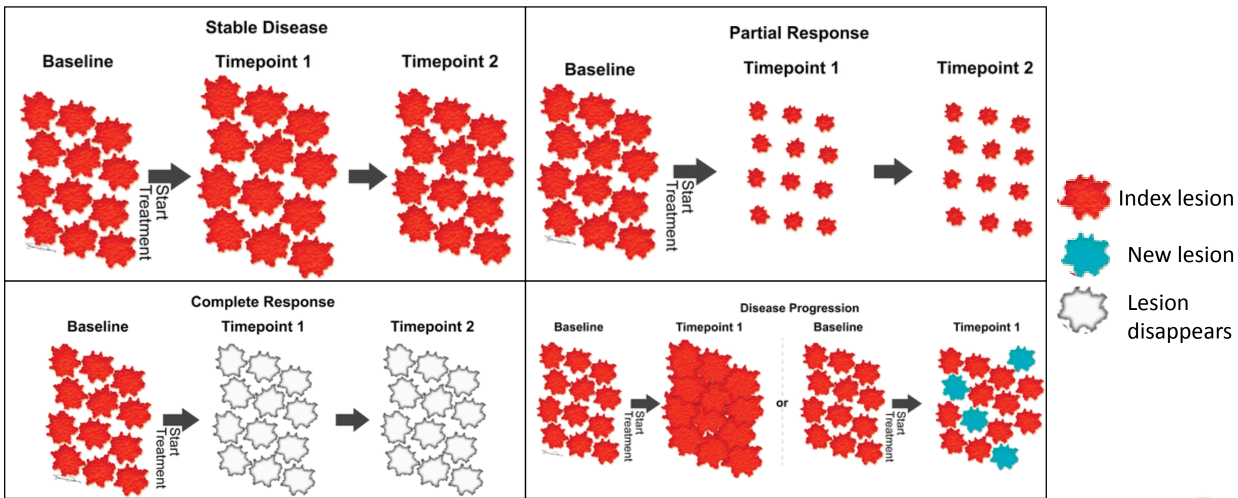








## Many possible imaging findings





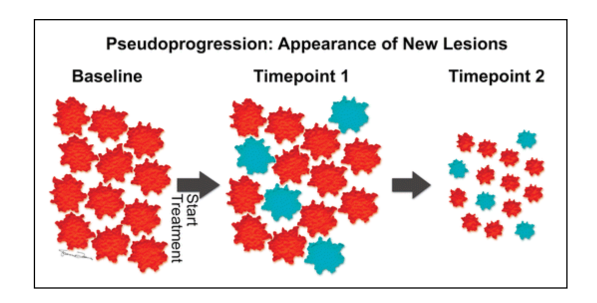


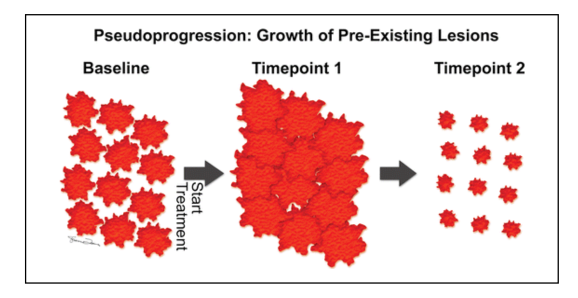






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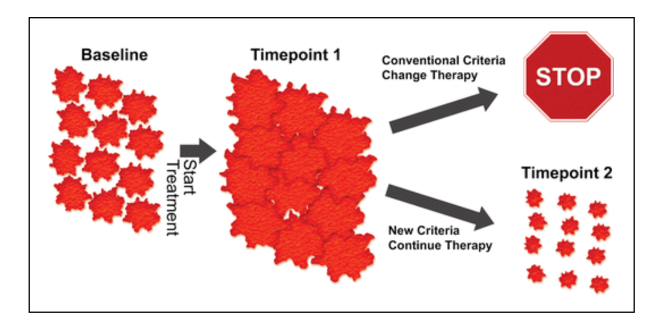


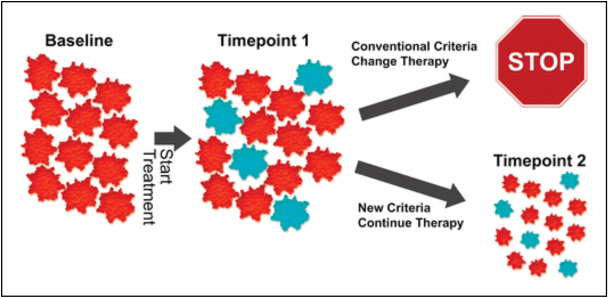






# Assessment of response – unique considerations for immunotherapy















## Comparison of disease progression by conventional and immune-related criteria

Treatment Response	RECIST 1.1	irRC	
Progressive disease	≥20% increase in lesion sum* (absolute size increase ≥5 mm) or 1+ new lesions at any single observation	≥25% increase in tumor burden+ versus nadir in two consecutive observations ≥4 weeks apart	
New measurable lesions#	Always represent progressive disease	Incorporated into disease burden	
New non-measurable lesions	Considered equivocal; followed at future examinations to clarify whether it is truly new disease	Does not define progression but precludes complete response	

Wang, RadioGraphics 2017.









<sup>\*</sup>Sum of lesion diameters: sum of the longest diameter in the plane of measurement for non-nodal target lesions and short-axis diameter for target nodal lesions.

<sup>&</sup>lt;sup>+</sup>Based on the sum of the products of the two largest perpendicular diameters of all index lesions.

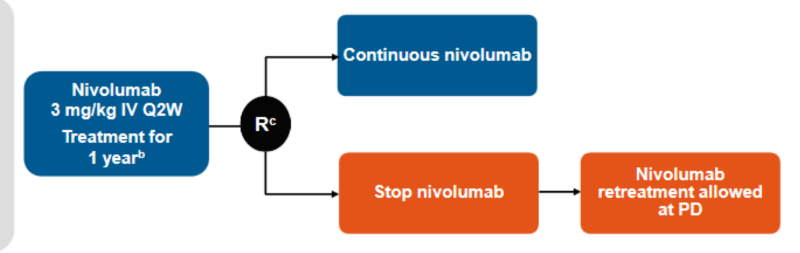
<sup>\*</sup>Measurable lesion for RECIST1.1 is ≥10mm at CT; irRC is ≥10x10mm at CT. Smaller lesions are considered non-measurable.



# When to stop immunotherapy: Checkmate 153

#### Key eligibility criteria:

- Advanced/ metastatic NSCLC
- ≥1 prior systemic therapy<sup>a</sup>
- ECOG PS 0-2
- Treated CNS metastases allowed



Exploratory endpoints<sup>d</sup>: Safety/efficacy<sup>e</sup> with continuous vs 1-year treatment, efficacy, other (eg, biomarkers, PK)



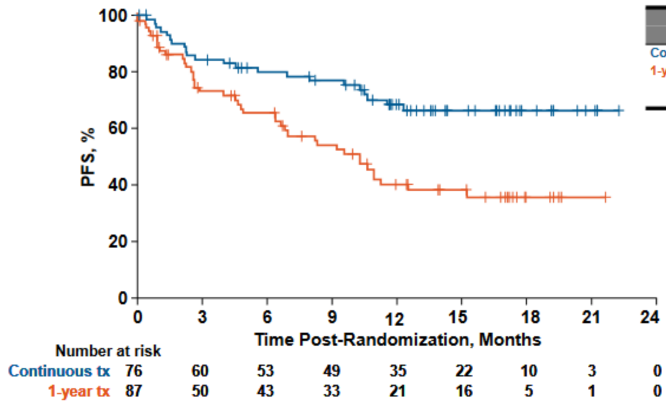








# When to stop immunotherapy: Checkmate 153



	Median, Months	PFS Rate, %	
	(95% CI)	6-Month	1-Year
Continuous tx	NR (NR)	80	65
1-year txb	10.3 (6.4, 15.2)	69	40
н	IR: 0.42 (95%	CI: 0.25.	0.71)

Conclusion: >1 year of treatment may be necessary











# When to stop immunotherapy: KEYNOTE-006

- Pembrolizumab 10 mg/kg Q2W or Q3W or ipilimumab 3 mg/kg Q3W for 4 doses
- Could stay on pembrolizumab for up to 2 years
- Of patients who completed 2 y pembro treatment, 86% did not progress after 20 months follow-up
- More responders with pembrolizumab, but duration of response was similar for pembrolizumab and ipilimumab





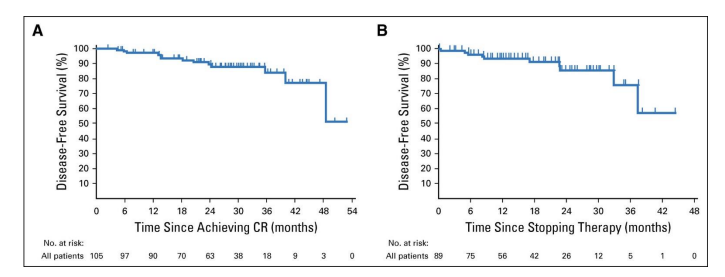






# When to stop immunotherapy: KEYNOTE-001

- 16% of patients achieved complete response
- Disease-free survival at 24 months after complete response:
  - In all CR patients: 90.9%
  - In patients who discontinued cancer therapy: 89.9%













# When to stop immunotherapy: clinical measures

- PET-based metabolic response
  - Metabolic response may precede anatomical changes on CT or MRI
- Achievement of CR



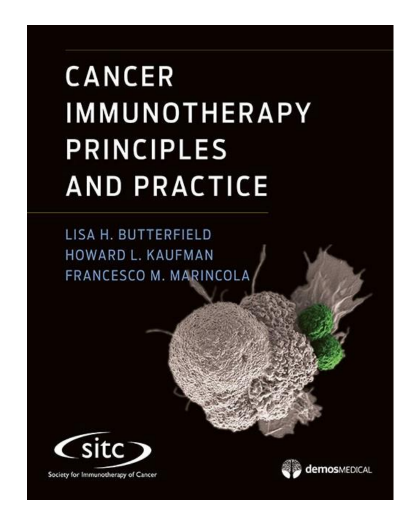








#### Further Resources



#### **SOCIETY FOR IMMUNOTHERAPY OF CANCER**









