

Mechanisms of Immune-Related Adverse Events

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Disclosures

- No relevant financial relationships to disclose
- I will not be discussing non-FDA approved indications during my presentation.









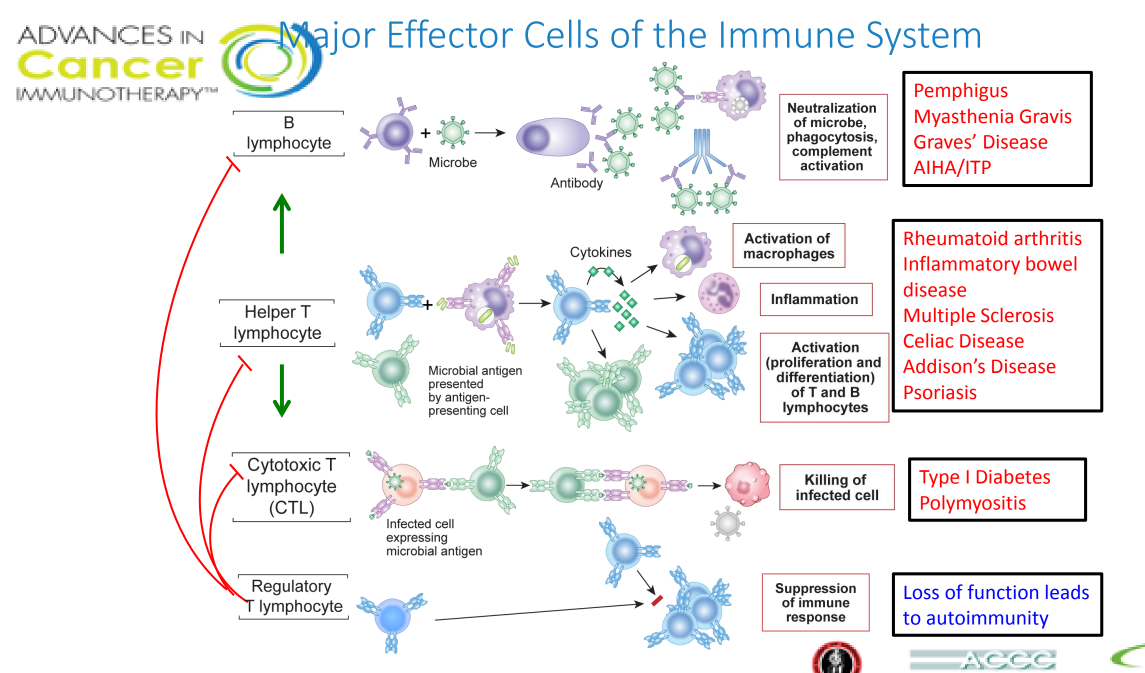
Outline

- Basic principles of immunological tolerance and autoimmunity
- Differential roles of CTLA-4 and PD-1 in maintenance of tolerance
- Mechanisms of breakdown of tolerance by checkpoint blockade











Most Autoimmune Diseases are due to Failure of T cell Tolerance

Immunologic Tolerance: unresponsiveness of immune system to self antigens

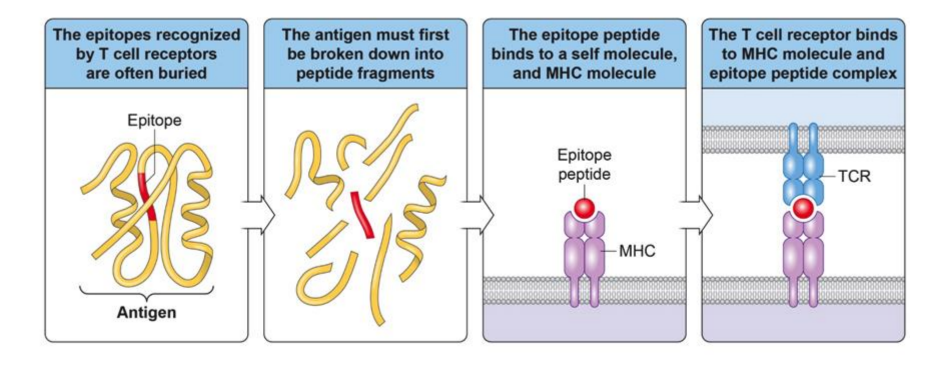








As a reminder...



MHC = Major Histocompatibility Complex

also called the HLA (human leukocyte antigen) complex









HLA (or MHC) is the strongest genetic factor for susceptibility to autoimmune disease

HLA- and gender-associated risk for autoimmune disease			
Disease	HLA allele	Relative risk	Sex ratio (우:♂)
Ankylosing spondylitis	B27	87.4	0.3
Type 1 diabetes	DQ2 and DQ8	~25	~1
Goodpasture's syndrome	DR2	15.9	~1
Pemphigus vulgaris	DR4	14.4	~1
Autoimmune uveitis	B27	10	<0.5
Psoriasis vulgaris	CW6	7	~1
Systemic lupus erythematosus	DR3	5.8	10–20
Addison's disease	DR3	5	~13
Multiple sclerosis	DR2	4.8	10
Rheumatoid arthritis	DR4	4.2	3
Graves' disease	DR3	3.7	4–5
Hashimoto's thyroiditis	DR5	3.2	4–5
Myasthenia gravis	DR3	2.5	~1
Type I diabetes	DQ6	0.02	~1

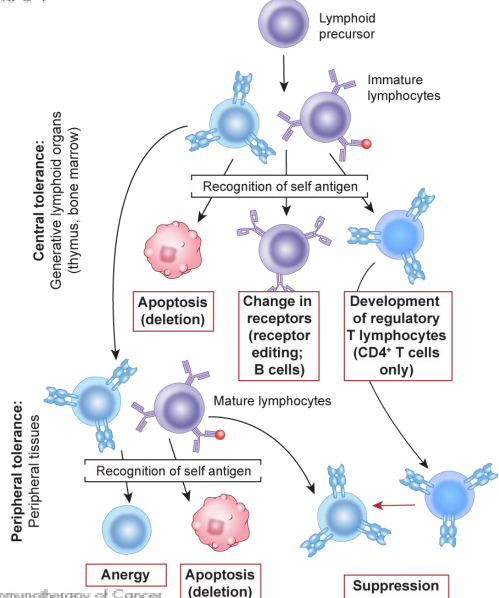






ADVANCES IN

Central and Peripheral Tolerance



Central Tolerance

- For T cells it occurs in the thymus
- Fate of high affinity, self-reactive T cells is death (deletion) and removal from T cell pool
- Some survive as regulatory (suppressor) T cells while others escape to peripheral tissues

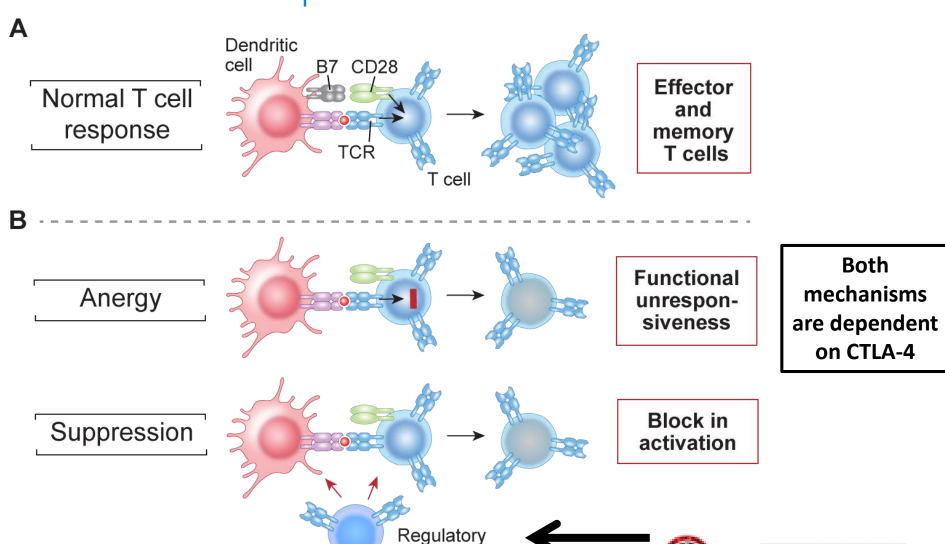
Peripheral Tolerance

- Self-reactive T cells are suppressed by regulatory T cells
- CTLA-4 and PD-1, among other molecules play a role in maintaining selfreactive T cells from becoming activated (anergic) ACCC



IMMUNOTHERAPY™

ADVANCES Periphral tolerance occurs in the absence of CD28 dependent co-stimulation

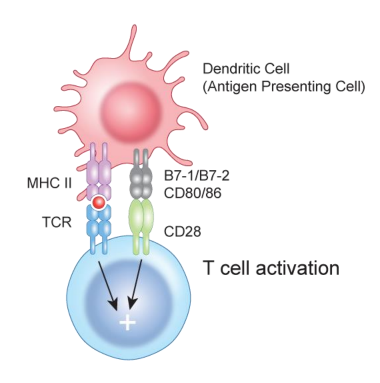


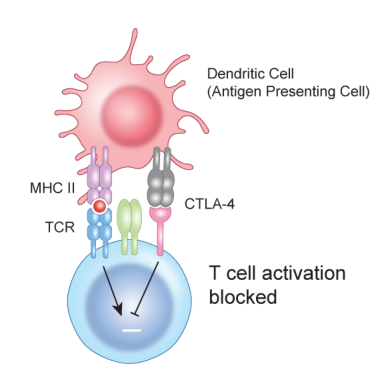
T cell





LA-4 inhibits co-stimulation by blocking interaction between CD28 and B7 molecules



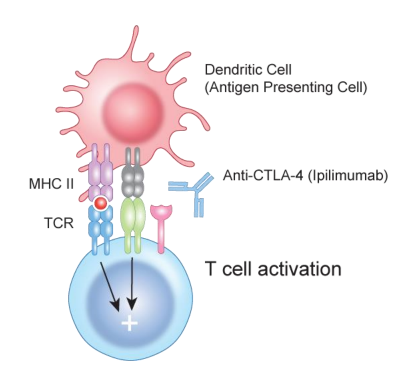


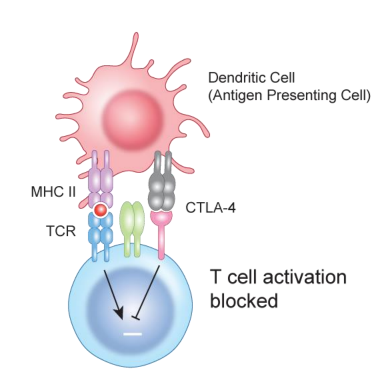






Cancer TLA-4 can lead to breakdown of peripheral tolerance by restoring co-stimulation





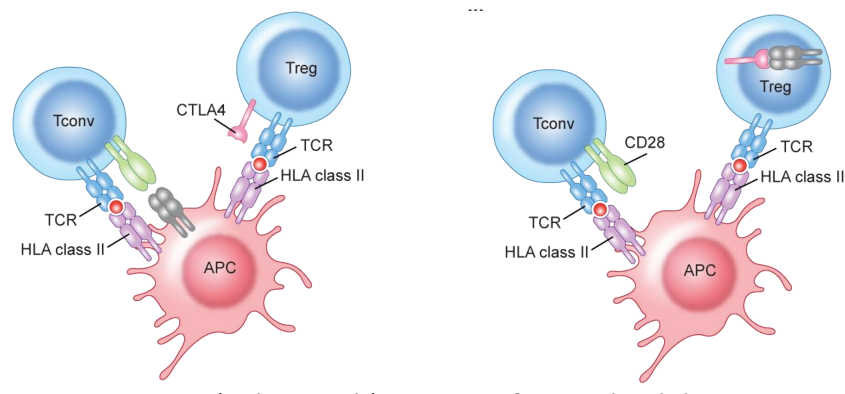
Breakdown of peripheral tolerance leading to activation of self-reactive T cells







ADVANCES in Regulatory T cells (Tregs) use CTLA-4 to remove B7 Cancer Molecules from surface of antigen presenting cells to prevent activation of self reactive T cells

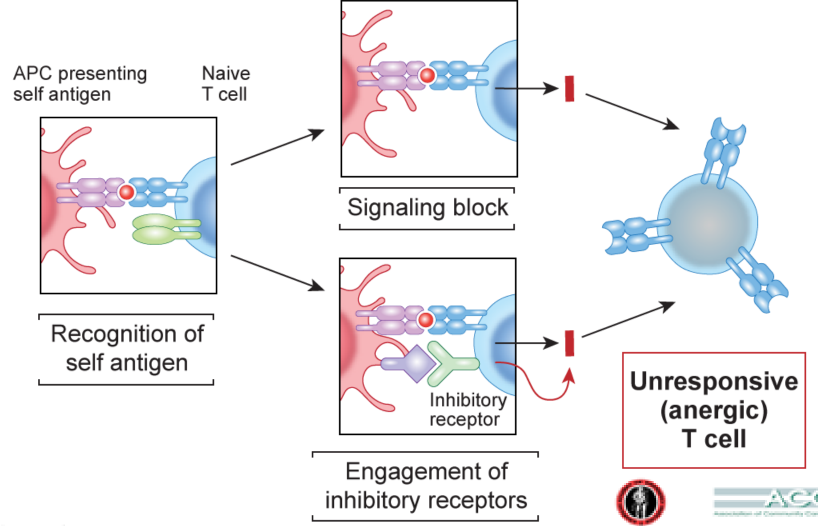


Anti-CTLA-4 (ipilimumab) may interfere with inhibitory function of Tregs





nhibitory receptors provide a second mechanism for maintenance of tolerance

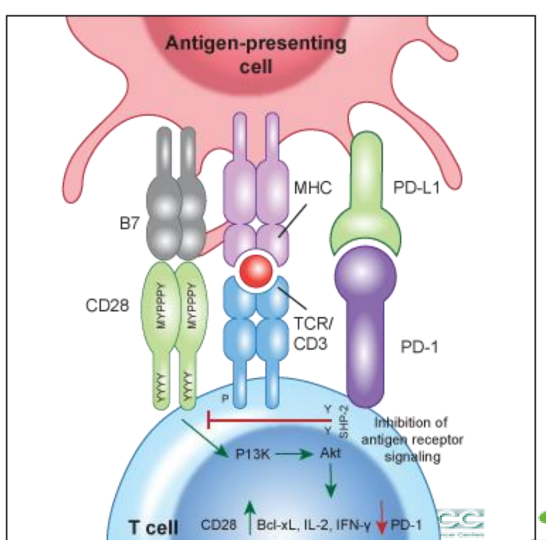






ADVANCES IN Internation of PD-1 with its ligands, PD-L1/PD-L2 inhibits CD28 signaling in T cells

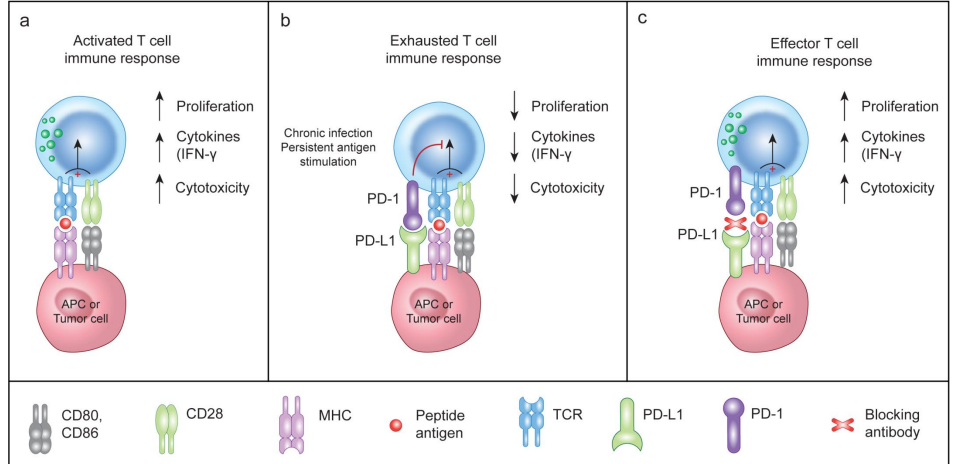
- PD-1 is upregulated on T cells after activation
- PD-L1 is found on both immune and non-immune cells in peripheral tissues
- PD-L2 is mostly found on immune cells in response to inflammatory stimuli
- In contrast, CTLA-4 and its ligands are only found on immune cells
- Mice deficient in PD-1 have delayed development of autoimmune disease compared to CTLA-4 deficient ones





Blocking PD-1/PD-L1 Pathway Reactivates T cells

<u>PD-1</u> is the receptor on T cells – its ligand <u>PD-L1</u> is on immune cells or tumor cells









Autoimmune Disease	Polymorphism
Thyroiditis, Graves' disease, Hashimoto's disease	CTLA-4
Diabetes mellitus	CTLA-4
Celiac disease	CTLA-4
Myasthenia gravis	CTLA-4
Lupus	CTLA-4; PD-1
Rheumatoid Arthritis	CTLA-4; PD-1
Addison's disease	CTLA-4

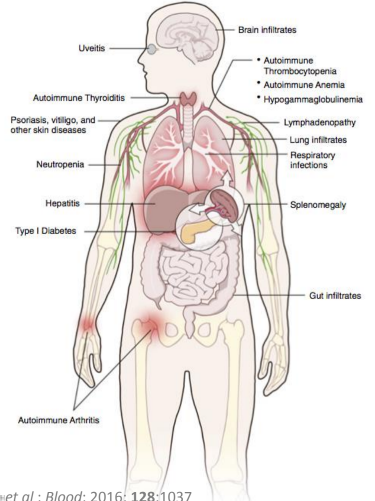




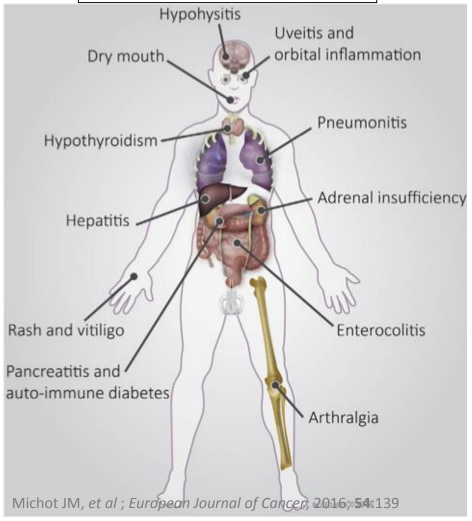


ADVANCES IN People with CTLA-4 haploinsufficiency develop a Canceppetrum of autoimmune diseases similar to the irAEs observed with ipilimumab

CHAI/LATAIE Phenotype



Ipilimumab irAEs







late irAEs may occur by distinct mechanisms

Early and common

Mucosal

Colitis

Rash

Pneumonitis

Global Regulatory T cell dysfunction

Activation of Effector T cells (Th₁₇)

Recruitment of inflammatory cells (neutrophils)

Late and rare

Specific organ
Hypophysitis
(other endocrine)
Myocarditis; Neurologic
Arthritis; Vitiligo

Breakdown of organ specific tolerance

Activation of tumor specific T cells that recognize antigen shared between tumor and healthy tissue: vitiligo, myocarditis

Activation of tissue specific anergic T cells that recognize antigen distinct from the tumor

T cell or probable mediated tissuestruction





- CTLA-4 expression on effector and regulatory T cells prevents co-stimulation through CD28 and maintains T cell anergy and peripheral tolerance
- Activation of PD-1 on activated T cells by its ligands renders them non-functional
- PD-1 activates regulatory T cells to maintain peripheral tolerance
- Humans with CTLA-4 haploinsufficiency develop a spectrum of autoimmune manifestations similar to irAEs seen after treatment with Ipilimumab





