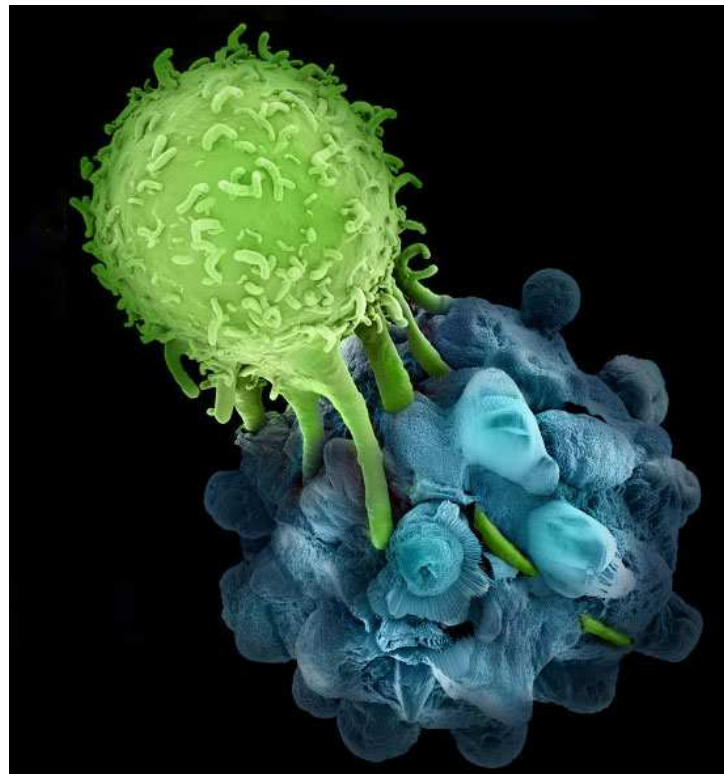


# Overview of Adaptive Immunity

Jonathan Powell  
poweljo@jhmi.edu



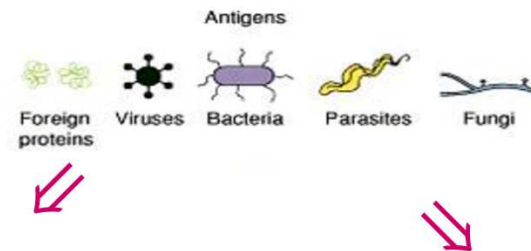
The exquisite specificity of the Adaptive Immune Response makes it a powerful modality for the treatment of cancer



2013 Cancer Immunotherapy Trials Network

# YOUR ACTIVE IMMUNE DEFENSES

1



## Innate Immunity

- invariant (generalized)
- early, limited specificity
- the first line of defense

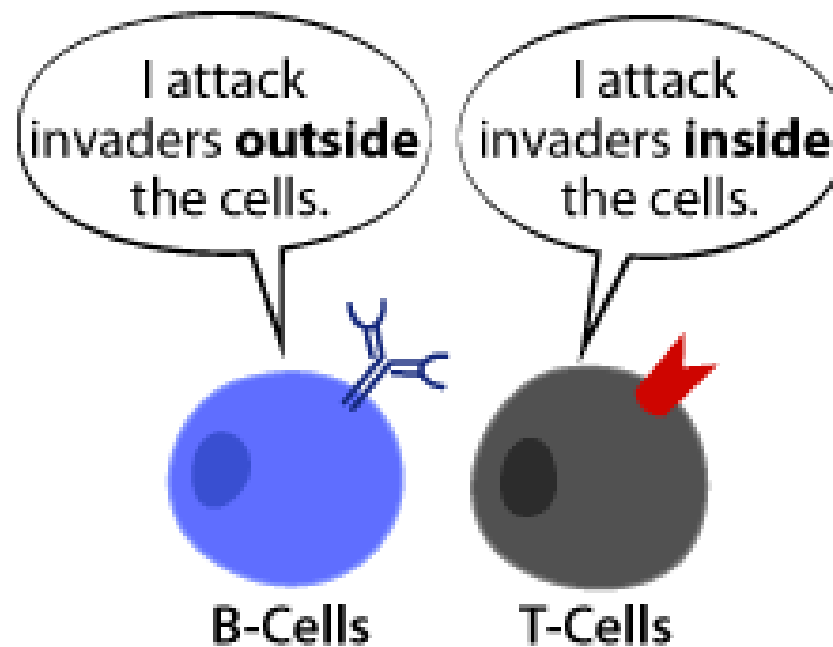
1. Barriers - skin, tears
2. Phagocytes - neutrophils, macrophages
3. NK cells and mast cells
4. Complement and other proteins

## Adaptive Immunity

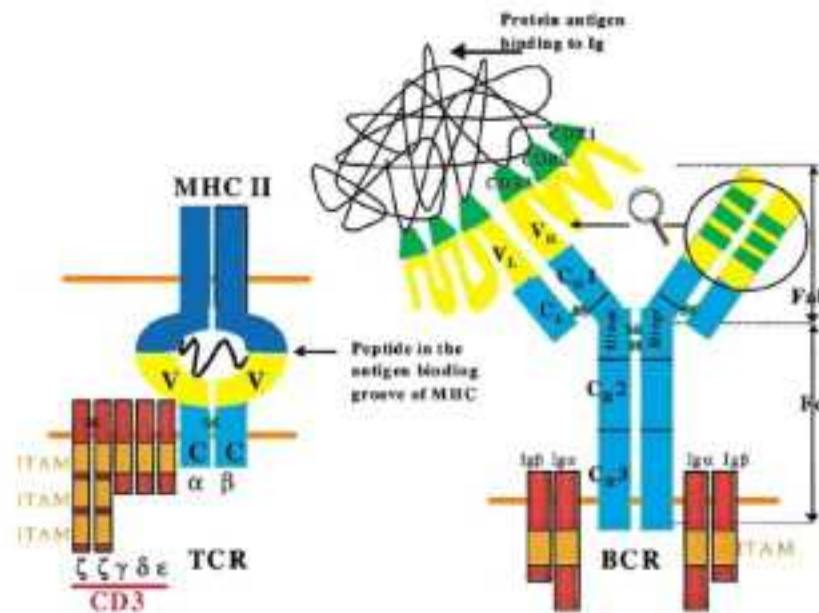
- variable (custom)
- later, highly specific
- “remembers” infection

1. APCs present Ag to T cells
2. Activated T cells provide help to B cells and kill abnormal and infected cells
3. B cells - produce antibody specific for antigen

# T Cells and B cells make up the Adaptive Immune Response



T Cell Receptors (TCR) recognize antigen presented on the surface of cells while B Cell Receptors (BCR, AKA antibodies) recognize soluble antigen



## 2 key features of the adaptive immune response

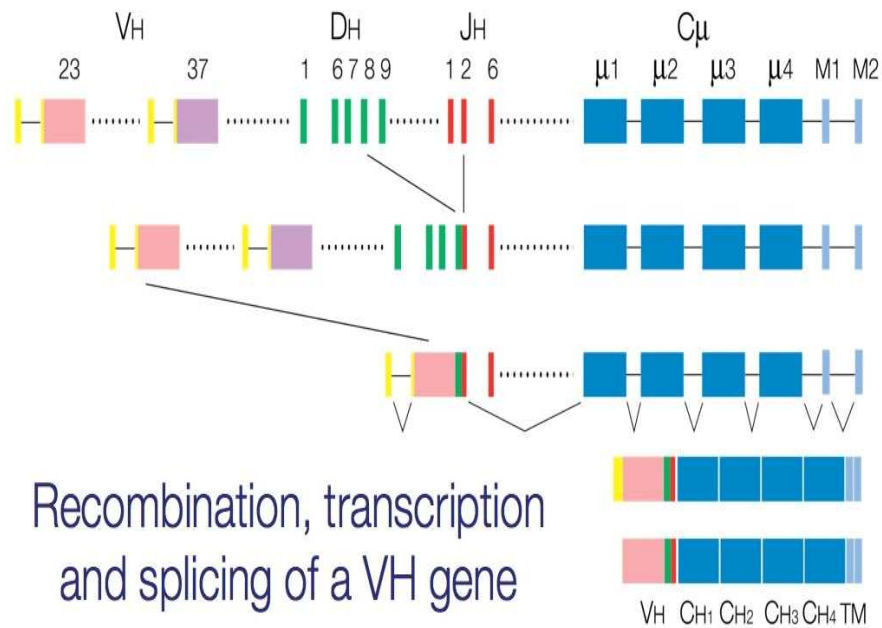
- A vast array of incredibly diverse antigen specific receptors
- Memory, that is the ability to respond rapidly upon rechallenge by the same pathogen



# Generation of Diversity

- The diverse number of antigen specific TCR and BCR are NOT encoded by germline genes
- Rather, each receptor is generated through recombining different combinations of genes

# Generation of Diversity through recombination



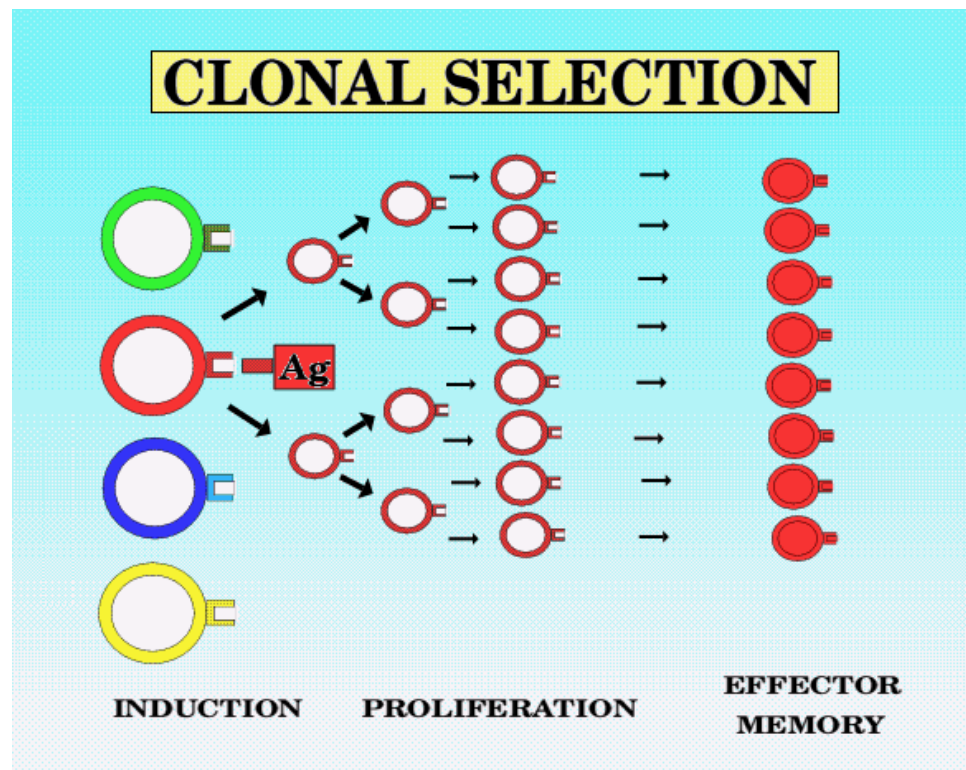


# Do the math

The [human genome](#) is presently estimated to contain 20–25 thousand genes. The number of T-cell receptors for antigen (**TCRs**) that we make is estimated at  $2.5 \times 10^7$ ; the number of different kinds of antibody molecules (**BCRs**) is probably about the same.

Antibodies (BCRs)	Gene Segments	Combinations
V $\kappa$	40	200 $\kappa$ chains
J $\kappa$	5	
V $\lambda$	31	124 $\lambda$ chains
J $\lambda$	4	
V <sub>H</sub>	51	7,650 H chains
D <sub>H</sub>	25	
J <sub>H</sub>	6	
Any H chain with any L chain (324)		$2.5 \times 10^6$

If there are so many different receptors how does an effective immune response generated?

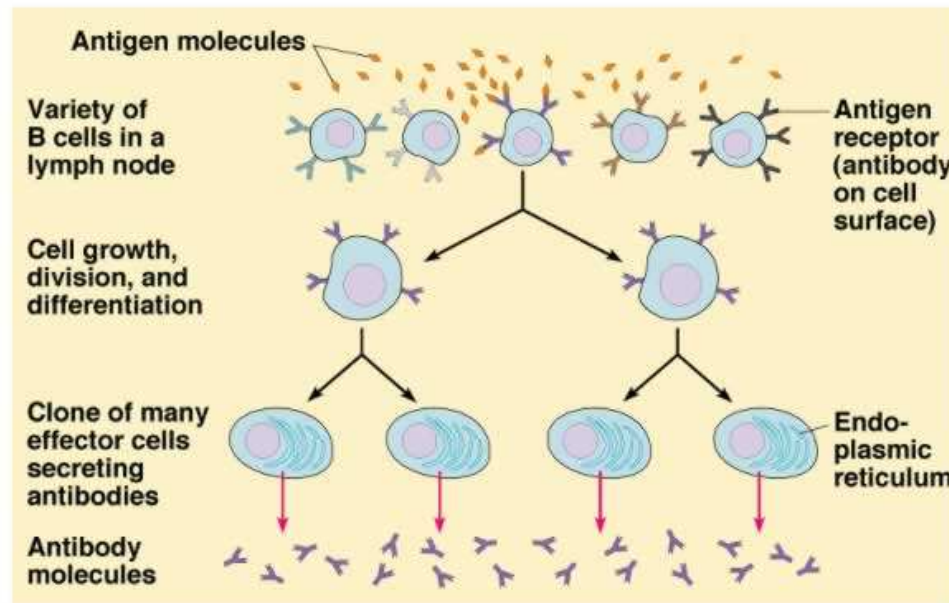


Steve Cobold

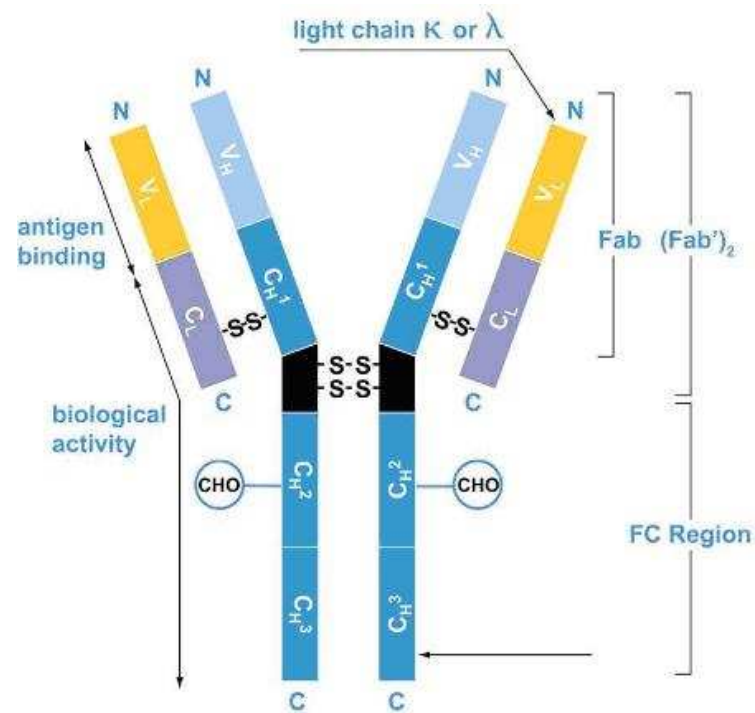
## Thus, clonal selection....

- Promotes robust, antigen specific effector responses
- Leads to the generation of memory

# B cells make antibodies



# Basic structure of antibodies



# Antibodies come in different flavors

## Antibody isotypes of mammals

Name	Types	Description
IgA	2	Found in mucosal areas, such as the gut, respiratory tract and urogenital tract, and prevents colonization by pathogens. Also found in saliva, tears, and breast milk.
IgD	1	Functions mainly as an antigen receptor on B cells that have not been exposed to antigens. It has been shown to activate basophils and mast cells to produce antimicrobial factors.
IgE	1	Binds to allergens and triggers histamine release from mast cells and basophils, and is involved in allergy. Also protects against parasitic worms.
IgG	4	In its four forms, provides the majority of antibody-based immunity against invading pathogens. The only antibody capable of crossing the placenta to give passive immunity to fetus.
IgM	1	Expressed on the surface of B cells and in a secreted form with very high avidity. Eliminates pathogens in the early stages of B cell mediated (humoral) immunity before there is sufficient IgG.

### Antibody Complexes



Monomer  
IgD, IgE, IgG



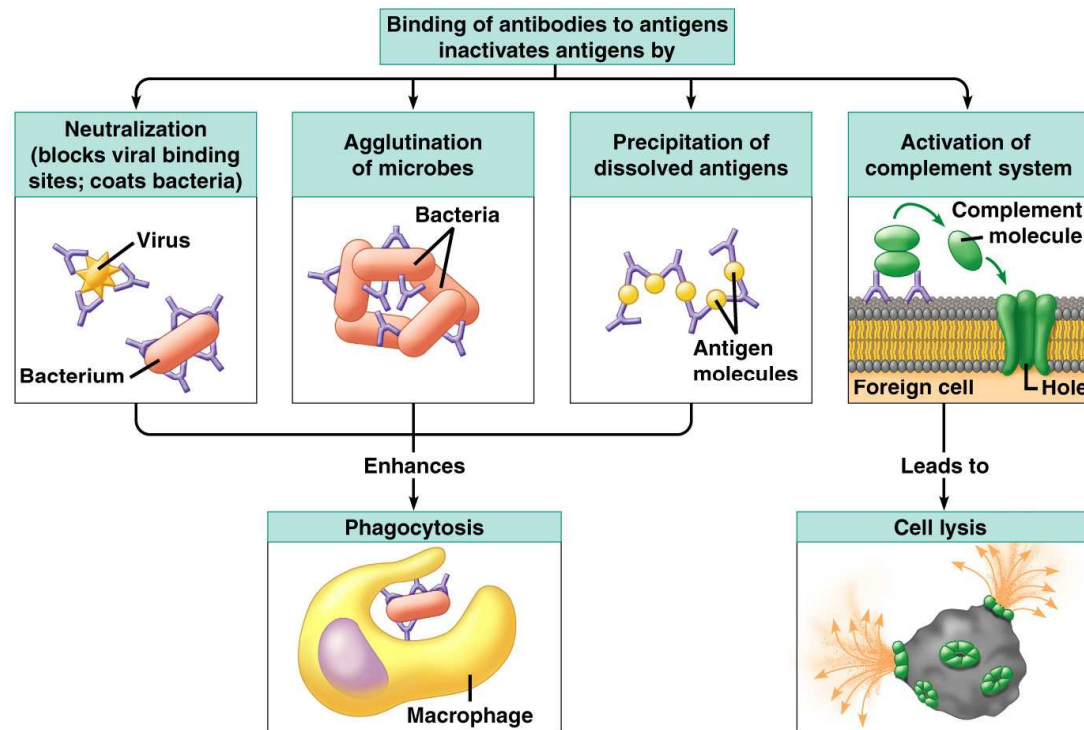
Dimer  
IgA



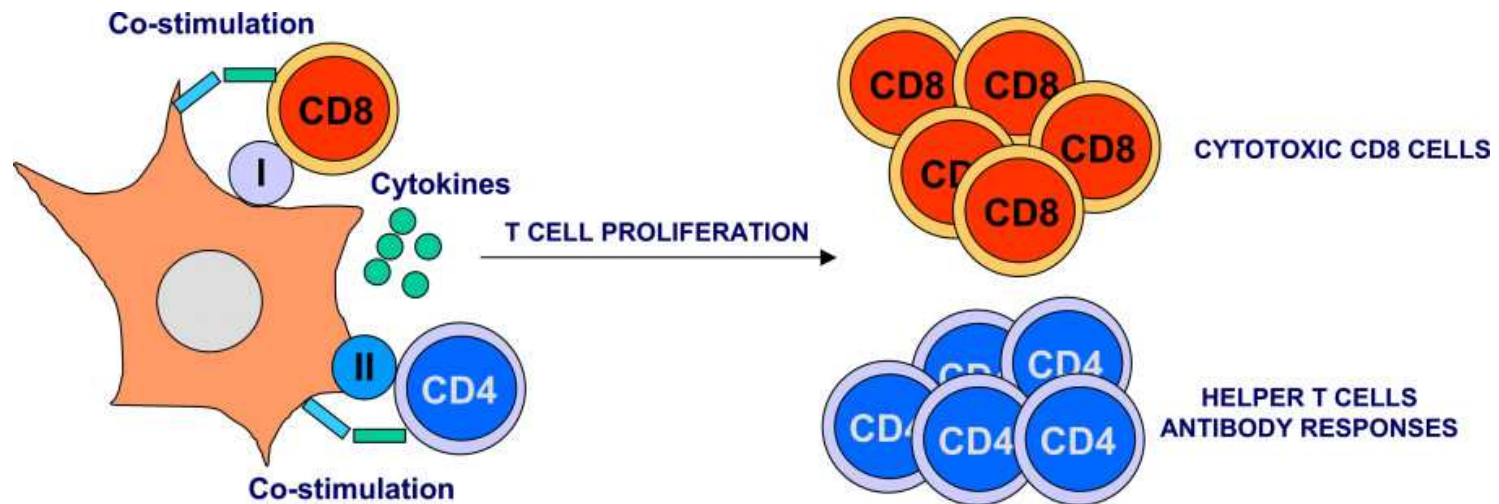
Pentamer  
IgM

*Adapted from Wikipedia, the free encyclopedia.*

# Antibody effector mechanisms

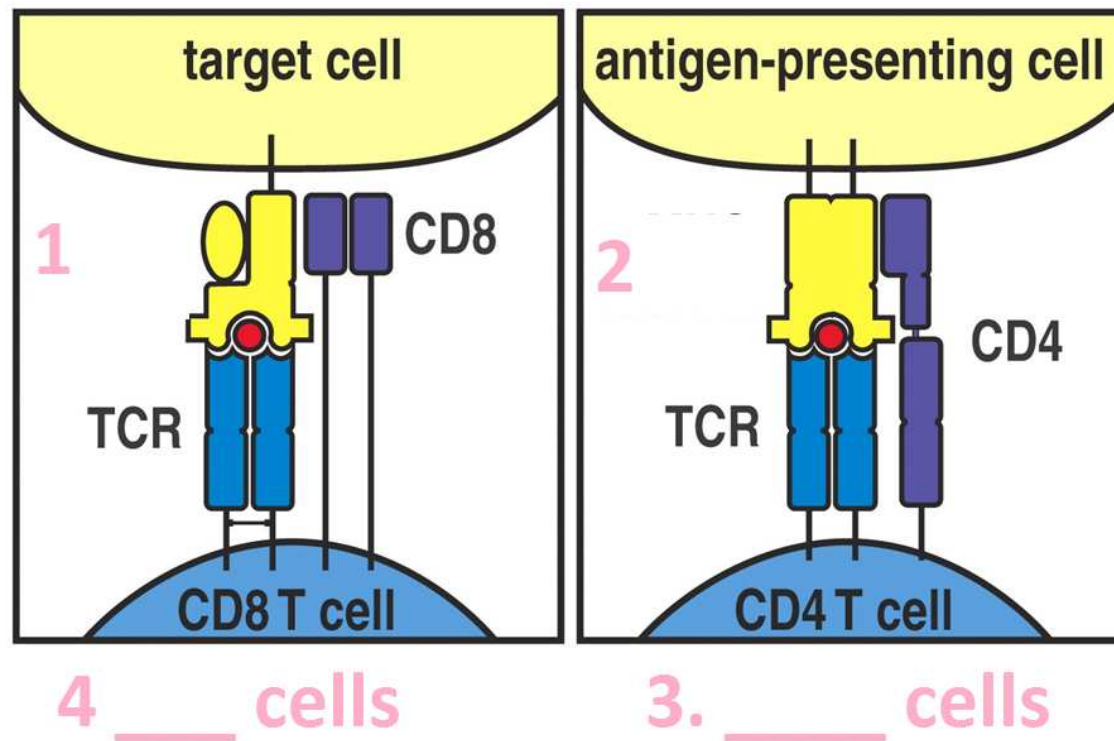


# T cells come in Two flavors





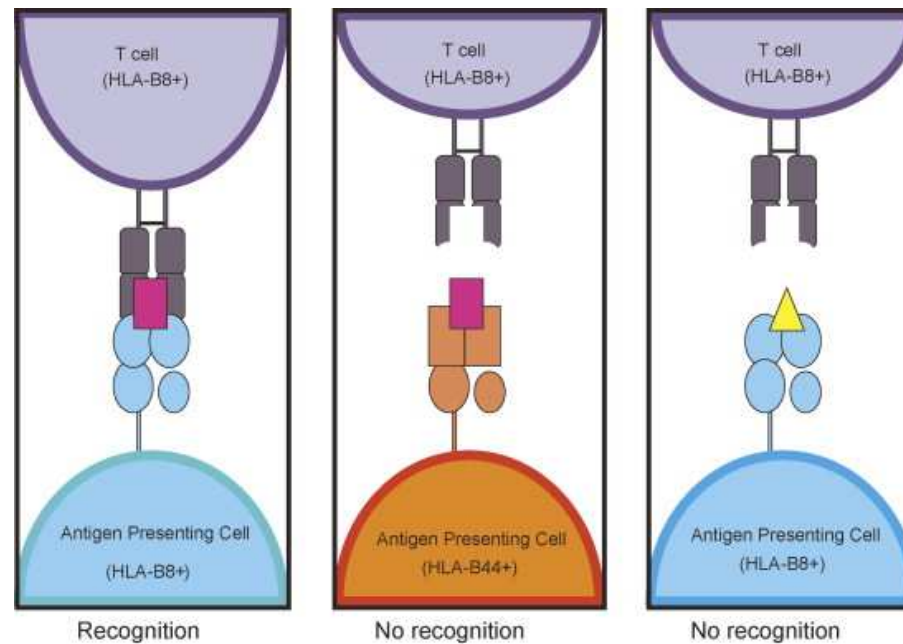
CD8 T cells recognize antigen presented by MHC class I and  
CD4 T cells recognize antigen presented by MHC class II



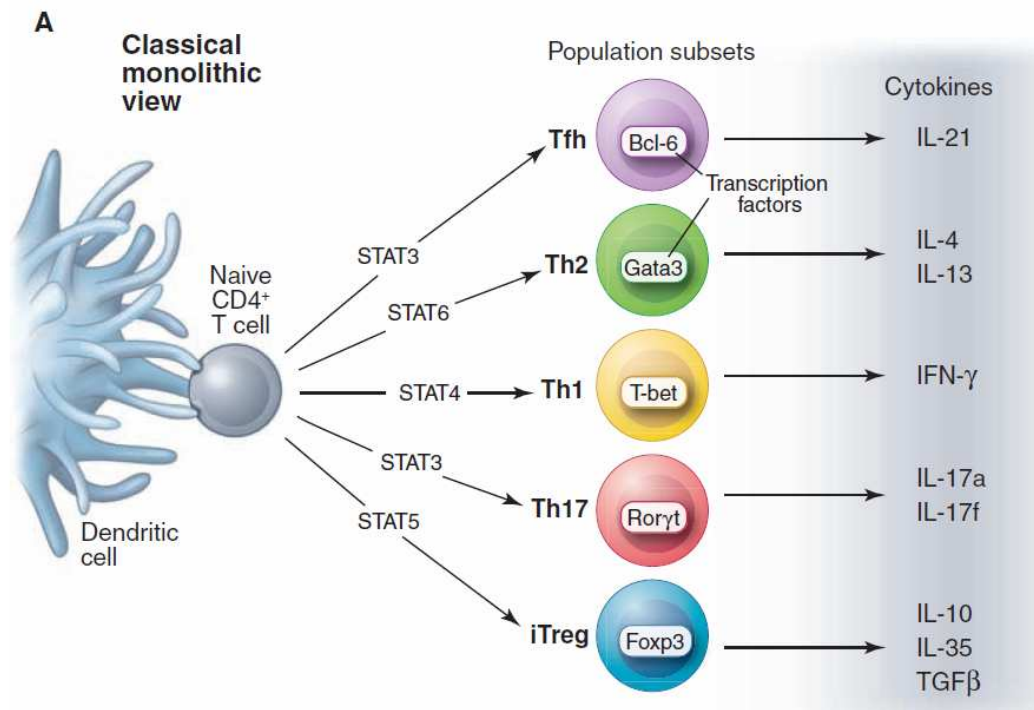
# Class I and Class II recognition relates to different functions of CD4 & CD8 cells

Table 2. Characteristics of the antigen processing pathways		
Characteristic	MHC-I pathway	MHC-II pathway
Composition of the stable peptide-MHC complex	Polymorphic chain $\alpha$ and $\beta$ 2 microglobulin, peptide bound to $\alpha$ chain	Polymorphic chains $\alpha$ and $\beta$ , peptide binds to both
Types of <a href="#">antigen presenting cells</a> (APC)	All nucleated cells	<a href="#">Dendritic cells</a> , mononuclear phagocytes, <a href="#">B lymphocytes</a> , some endothelial cells, epithelium of <a href="#">thymus</a>
T lymphocytes able to respond	<a href="#">Cytotoxic T lymphocytes</a> (CD8+)	<a href="#">Helper T lymphocytes</a> (CD4+)
Origin of antigenic proteins	<a href="#">cytosolic</a> proteins (mostly synthesized by the cell; may also enter from the extracellular medium via <a href="#">phagosomes</a> )	Proteins present in <a href="#">endosomes</a> or <a href="#">lysosomes</a> (mostly internalized from extracellular medium)

# Briefly, what is meant by MHC restriction?

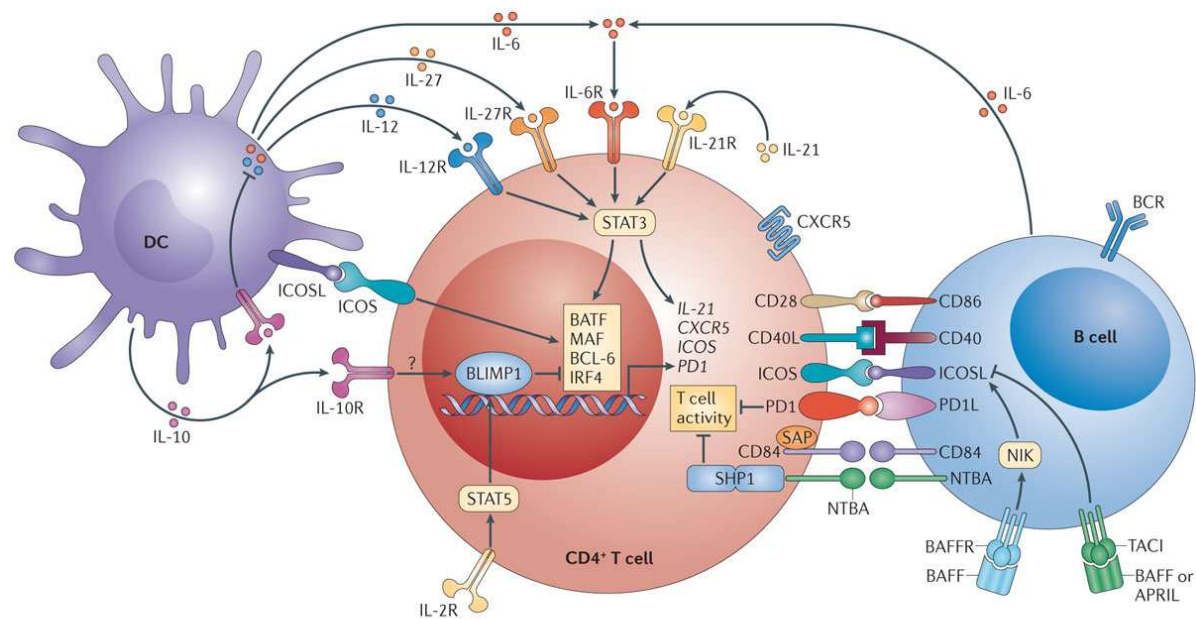


# Different Types of CD4 effector cells

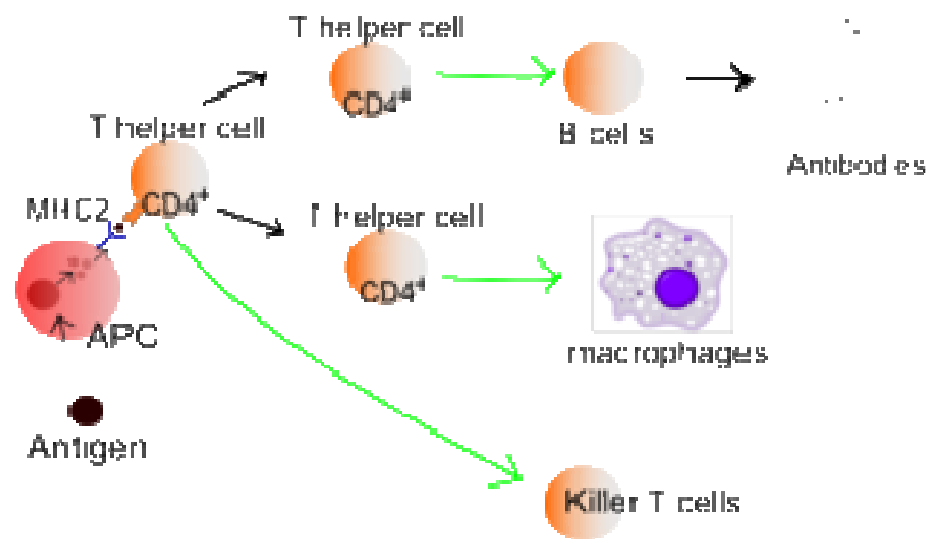


O'Shea JJ, Paul WE. Mechanisms underlying lineage commitment and plasticity of helper CD4<sup>+</sup> T cells. *Science*. 2010 Feb 26;327(5969):1098-102.

# CD4+ T cells “Help” B cells



CD4+ T cells also “Help” macrophages and CD8+ CTL's



# CD8+ T cells kill infected cells

## But first they must be activated

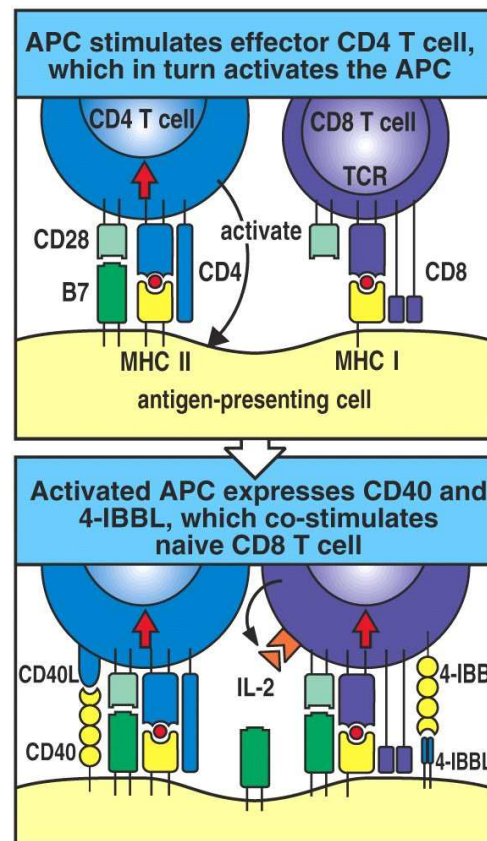


Figure 8-26 Immunobiology, 6/e. (© Garland Science 2005)

# Effector CD8+ T cells expand, kill their targets, then die

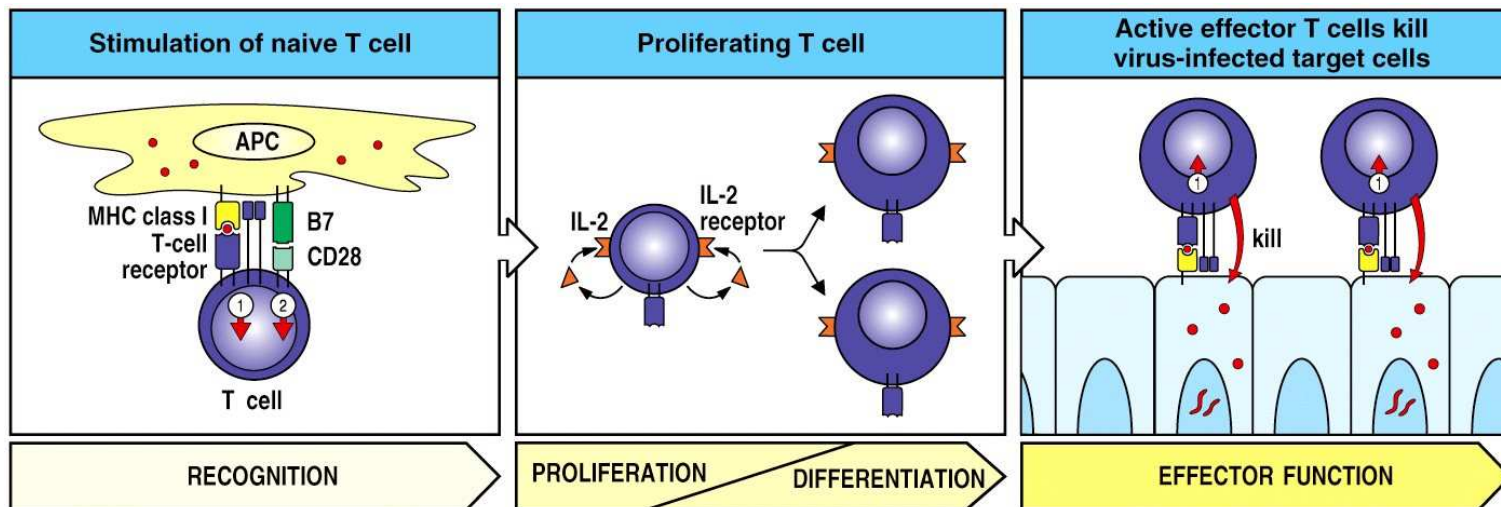
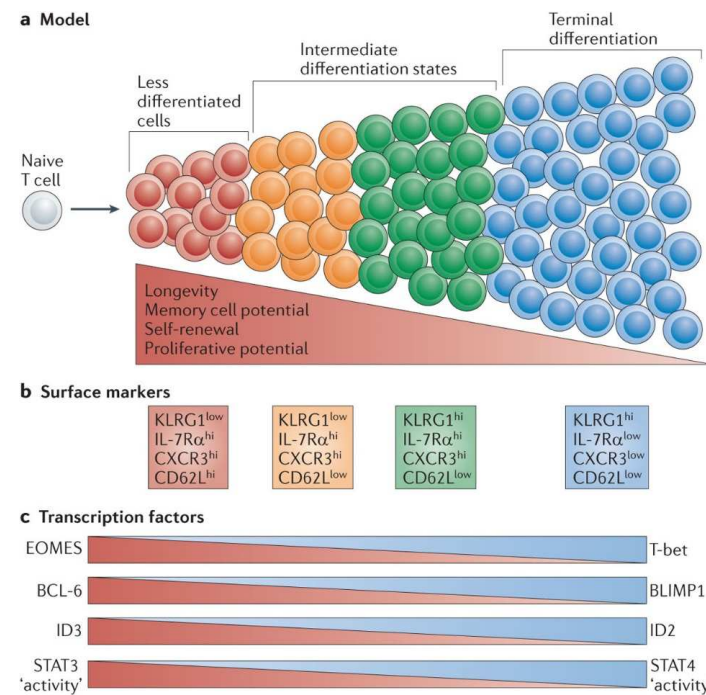


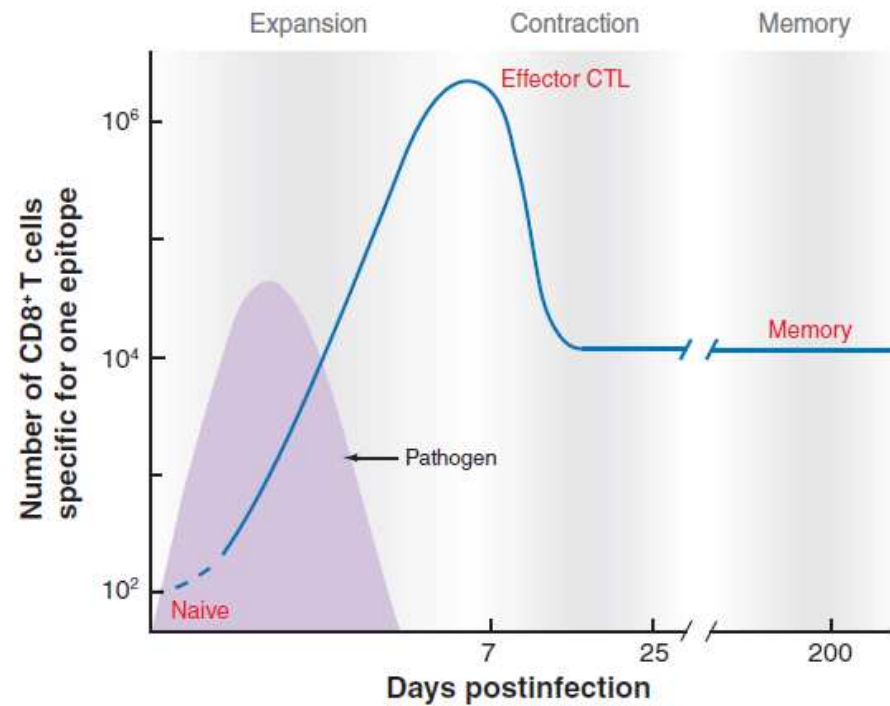
Figure 8-22 Immunobiology, 6/e. (© Garland Science 2005)



# Some of the activated CD8<sup>+</sup> T cells become long lived memory cells



# Generalized CD8+ T cell response



# Summary of T cell functions







	CD8 cytotoxic T cells	CD4 T <sub>H</sub> 1 cells	CD4 T <sub>H</sub> 2 cells	CD4 T <sub>H</sub> 17 cells	T <sub>FH</sub> cells	CD4 regulatory T cells (various types)
Types of effector T cell						
Main functions in adaptive immune response	Kill virus-infected cells	Activate infected macrophages Provide help to B cells for antibody production	Provide help to B cells for antibody production, especially switching to IgE	Enhance neutrophil response Promote barrier integrity (skin, intestine)	B-cell help Isotype switching Antibody production	Suppress T-cell responses
Pathogens targeted	Viruses (e.g. influenza, rabies, vaccinia) Some intracellular bacteria	Microbes that persist in macrophage vesicles (e.g. mycobacteria, <i>Listeria</i> , <i>Leishmania donovani</i> , <i>Pneumocystis carinii</i> ) Extracellular bacteria	Helminth parasites	<i>Klebsiella pneumoniae</i> Fungi ( <i>Candida albicans</i> )	All types	

Figure 9.1 Janeway's Immunobiology, 8ed. (© Garland Science 2012)

**Since the generation of diversity is a stochastic post-germline encoded event, a critical component of the immune response must subsequently be the ability to discern between harmful and innocuous antigens in a tolerance process.**



- On the one hand in order to deal with any and all potential pathogens the immune system generates at random a diverse array of receptors.
- On the other hand a mechanism (tolerance) must be in place in order to prevent these receptors from recognizing and destroying the organism it is designed to protect.

**Does the immune system recognize Good versus Evil or Self versus Non-self?**



# Shifting Paradigms

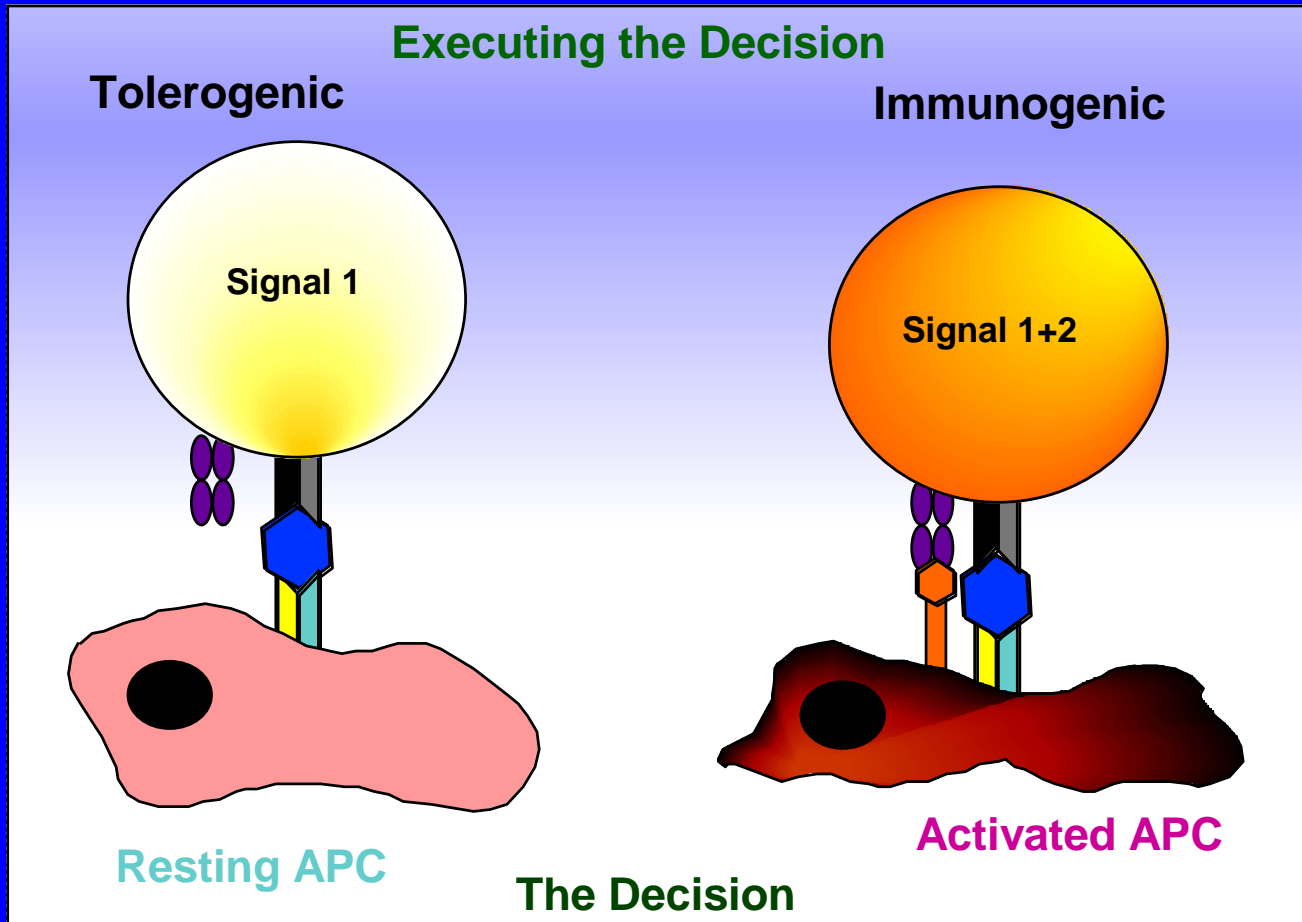
## Old

- Self-Non self
- Antigen Driven
- Developmentally Determined

## New

- Context Models
- Circumstance Driven
- Determined by milieu

# Executing the Will of APC's

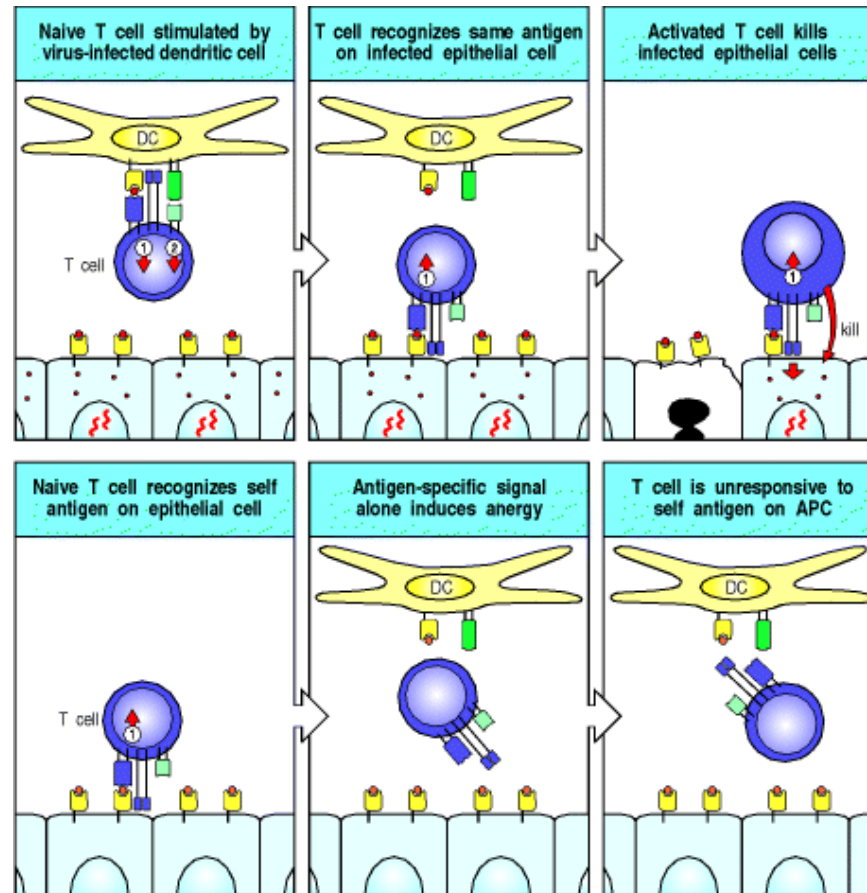


## **Activated APCs express costimulatory molecules (Signal 2)**

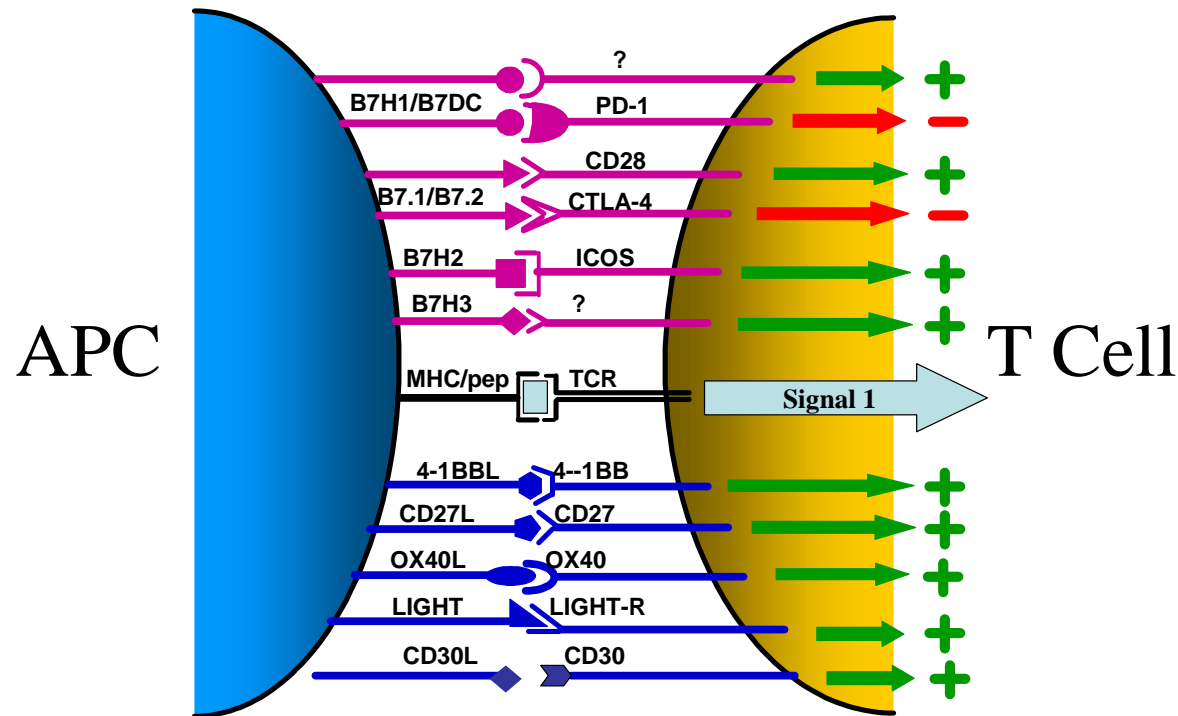
- **Signal 1: Refers to T cell receptor (TCR) recognition of antigen. Signal 1 alone leads to anergy or deletion.**
- **Signal 2: Refers to costimulatory molecule engagement (example CD28).**
- **Signal 1+2 leads to full T cell activation.**



## Consequences of Signal 1 versus Signal 1+2



In reality Signal 2 is the sum of both costimulatory signals AND co-inhibitory signals

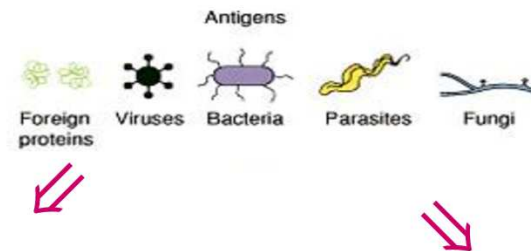


## **Based on what I told you how can we exploit the Adaptive Immune Response to Kill tumors?**

- Vaccine to induce clonal selection and memory response
- Passively give antibodies that are directed against tumors
- Grow up large quantities of tumor-specific T cells and then give them back to patient.
- Promote the activation of APC's (Rev up Signal 2)
- Inhibit/ destroy regulatory cells
- Block co-inhibitory signals (Checkpoint Blockade)

# YOUR ACTIVE IMMUNE DEFENSES

1



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