

What Cells Do When the Blitz is On Virus infects cell Viral proteins synthesized in cytoxic Peptide fragments of visit profess in a first of the cell surface Peptide fragments of visit profess board in a first of the cell surface Peptide fragments of visit profess board in a first of the cell surface Peptide fragments of visit profess of the cell surface Pe

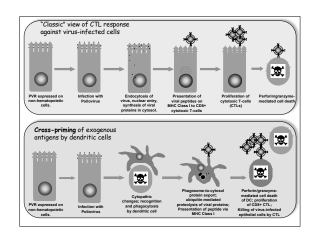
Antigen Presentation Pathways; Two Old:

MHC Class I presentation of peptides MHC Class II presentation of peptides

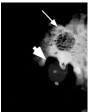
and Two New:

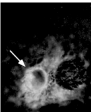
Cross-priming of exogenous peptides (MHC Class I) CD1-mediated presentation of glycolipids

Question: How do viruses that don't infect "professional APCs" such as dendritic cells elicit a primary immune response? After all, virally-infected cells normally don't traffic to 2° lymphoid organs



Cross-priming: A Dendritic Cell Engulfs a Virus-infected Macrophage

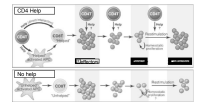




From: Albert et al., J. Exp. Med. 188:1359, 1998

Question: Does development of the cytotoxic T cell response require "help" from CD4 cells (analogous to help for B cells)?

CD8 T Cells Need Help With Their Memory*



*You will, too, in a couple of years

Memory T-cells Don't Forget

Table 1 Estimated survival of virus-specific T-cell memory after smallpox vaccination

	Voluntee	rs with CD4+ Ticell in	iemory ^a	
Vaccinations	20-30 years*	31-50 years	51-75 years	t _{1/2} of CD4* T cells
1	100% (16/16)	89% (70/79)	52% (23/44)	10.6 (0.17)
2	83% (10/12)	78% (29/37)	57% (4/7)	8.3 (0-14.1)
3-14	82% (23/28)	91% (29/32)	NDd	12.4 (0-20.5)
	Voluntes	rs with CD8+ T-cell r	nemory	
Vaccinations	20-30 years	31-50 years	51-75 years	t _{1/2} of CD8+ T cells
1	50% (8/16)	49% (39/79)	50% (22/44)	15.5(0-27.1)
2	42% (5/12)	38% (14/37)	57% (4/7)	8.1 (0-16.9)
3-14	46% (13/28)	50% (16/32)	ND	9.0 (0-18.1)

»10 IRV-18Fa" Tools per 10° 100° or 008° Tools respectively. The cutoff provides 100°s sensitivity at 1 month after vaccination or reversantation and 20°95's specificity. Bead on the vaccination or seconds in Tools because it Tools second tools respectively. The cutofficient vaccination is the second of the cutofficient vaccination. "Estimated half-life (t₁₂) in years fair 95% confidence interval in paper thesely is based on linear regression analysis using data from Figures 1 and 2, 8.0. In ord other papers.

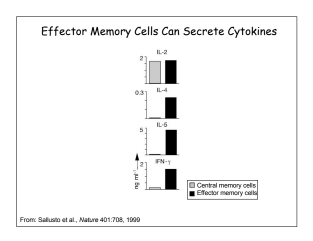
From: Hammarlund et al., Nature Med. 9:1131, 2003

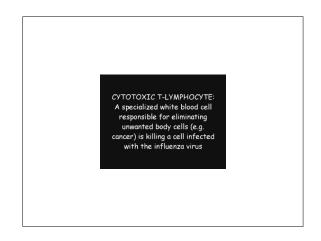
Phenotypic Differences Between Selected T Cell Subsets

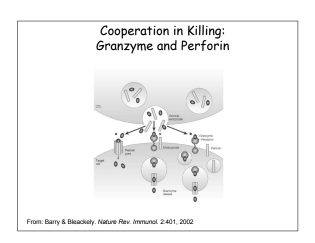
Phenotype	Naïve	
Migration	LN, spleen	
Cell cycle	-/+	
Cytokine secretion	-	
Peripheral LN homing (L-Selectin; CD62L)	+++	
Adhesion Molecules (Integrins, CD44)	+	
Chemokine Receptors (partial list)	CCR7	
IL-2 Receptor (CD25)	-	
FasL	-	

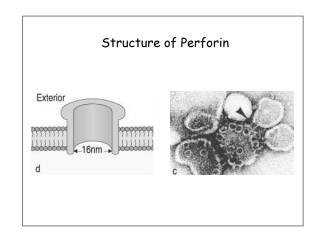
Phenotypic Differences Between Selected T Cell Subsets

Dhanatuna	Naïve	Effector	Memory	
Phenotype	Naive	Effector	Central	Effector
Migration	LN, spleen	Inflamed tissue	LN	Inflamed tissue
Cell cycle	-/+	++	+	++
Cytokine secretion	-	+++	-	+++
Peripheral LN homing (L-Selectin; CD62L)	+++	-	+++	-
Adhesion Molecules (Integrins, CD44)	+	+++	+++	+++
Chemokine Receptors (partial list)	CCR7	CCR5 CXCR4	CCR7	CCR5 CXCR4
IL-2 Receptor (CD25)	-	++	+	+/-
FasL	-	+++	-	+++

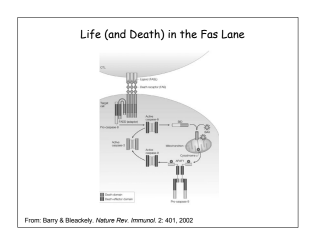


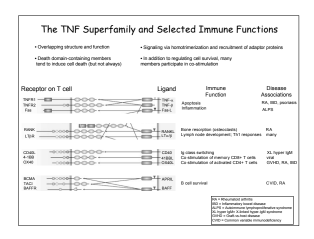


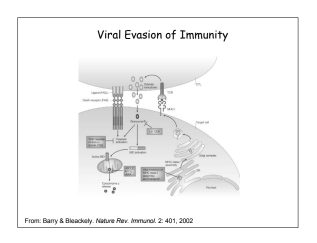


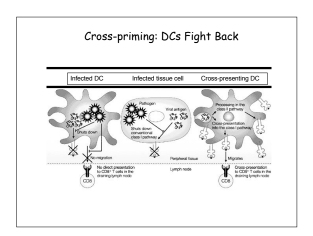


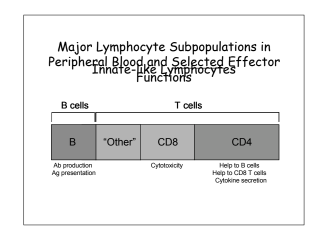
Human Diseases	Involving	Defective Granule Killing*
Disease	Gene	Clinical Manifestations
Chediak-Higashi Syndrome	CHS1	Lysosomal inclusions in all leukocytes Recurrent bacterial infections Decreased NK cell function Oculocutaneous albinism (melanosome defect) Bleeding (platelet storage granule defect)
Griscelli Syndrome	Rab27a	Partial albinism Hepatosplenomegaly (lymphohisticcytic infiltration) Decreased NK cell function
Hermansky-Pudlak Syndrome	HPS1	Oculocutaneous albinism (melanosome defect) Bleeding (Platelet storage granule defect) Pulmonary fibrosis (Type II cell surfactant body inclusions)
Familial Hemophagocytic Lymphohistiocytosis	Perforin (30% of cases)	Hepatosplenomegaly (accumulation of activated T-cell and macrophages) Decreased NK cell function Pancytopenia
*Do not memorize this list		

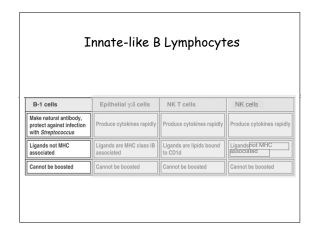


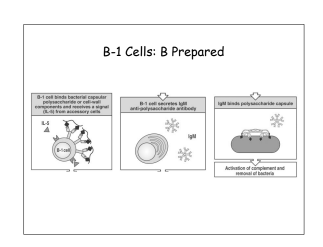












Thymus-independent Antigens are Presented to Specialized B-cells

 $\underline{\text{Marginal zone B cells}},$ like B-1 cells, respond to carbohydrate antigen and secrete mainly IgM

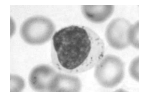


After the initial encounter of Ag, they demonstrate little memory (like trying to cram this course)

Innate-like T Lymphocytes: NK Cells

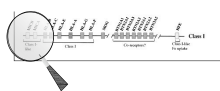
B-1 cells	Epithelial γ:δ cells	NK T cells	NK cells
Make natural antibody, protect against infection with Streptococcus	Produce cytokines rapidly	Produce cytokines rapidly	Produce cytokines rapidly
Ligands not MHC associated	Ligands are MHC class IB associated	Ligands are lipids bound to CD1d	Ligands not MHC associated
Cannot be boosted	Cannot be boosted	Cannot be boosted	Cannot be boosted

Natural Killer Cell

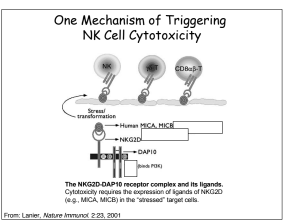


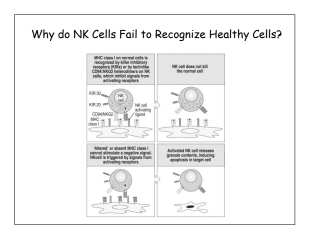
How do NK Cells Recognize Their Targets?

Major Genes in the MHC Class I Region

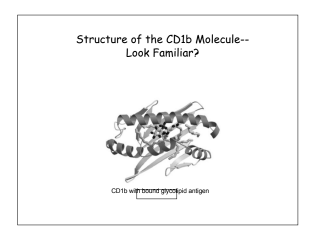


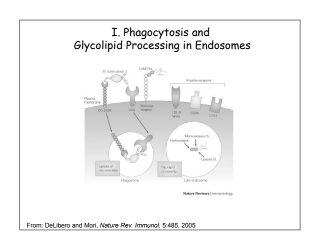
dale., Immunity. 15:363, 2001

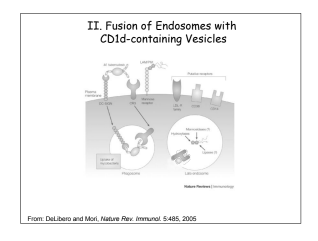


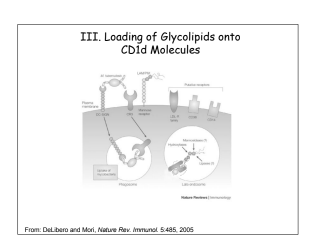


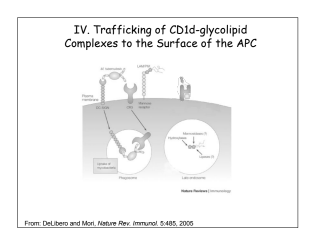
A Digression... Antigen Presentation of Non-peptide Antigens: Recognition of Glycolipid Antigens

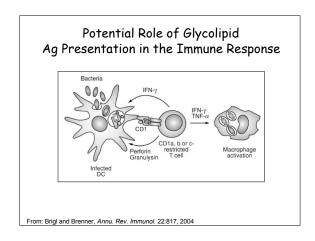


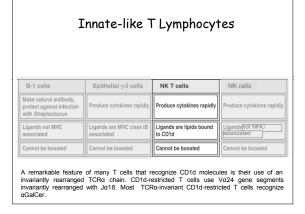


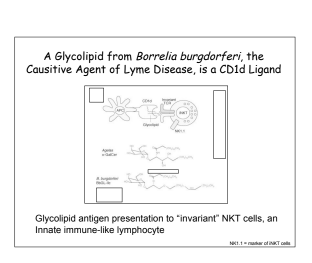


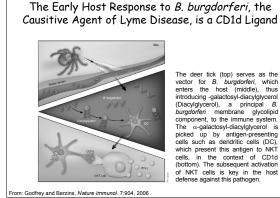












The deer tick (top) serves as the vector for *B. burgdorferi*, which enters the host (middle), then the tirducing -galactosyl-diacylglycerol (Diacylglycerol), a principal *B. burgdorferi* membrane glycollipid component, to the immune system. The *α*-galactosyl-diacylglycerol is picked up by antigen-presenting cells such as dendritic cells (DC), which present this antigen to NKT cells, in the context of CD1d (bottom). The subsequent activation of NKT cells is key in the host defense against this pathogen.

Innate-like T Lymphocytes B-1 cells Epithelial γ:δ cells NK cells NK T cells Ligands not MHC Ligands are MHC class IB Ligands not MHC Cannot be boosted Cannot be boosted

Question: Do lymphocytes of the acquired immune system even care about lymphocytes of the innate immune system?

Innate Immune Lymphocytes Trigger Dendritic Cell Maturation Antigen Dendritic Cell Maturation Small phospioor amnoantigens Viral proteins Wiral proteins

Summary

- 1. For cytotoxic CD8 T-cells, ligation of the TCR by MHC l/peptide + co-stimulation results in release of granzymes and perforin and/or FasL, leading to apoptosis of the target cells.
- 2. Viruses evade host defense, in part, by down-regulating MHC Class I. Uninfected dendritic cells circumvent this by "cross-priming": phagocytosis of virus-infected cell and presentation of "exogenous" viral antigens on MHC Class I.
- 3. CD8 T cells can function without CD4 help, but need CD4 help to develop into effective memory cells. CD4 memory cells live for years; central memory cells home to lymph nodes and effector memory cells home to inflamed tissue.
- 4. NK cells lack TCRs, but instead express both activating and inhibitory (e.g., KIRs) receptors at their surfaces. The relative expression and ligation of these receptors determines the outcome (i.e., killing or not) of the NK effector response.
- 5. Innate immune B-cells (e.g., B-1 cells and marginal zone B cells) recognize carbohydrate antigens, secrete lgM, and are not long-lived.
- 6. Innate immune T-cells (γ 8 T-cells, and NK T cells) recognize non-peptide antigens in non-classical MHC-like molecules. They mediate cytotoxicity, rapid cytokine secretion, and trigger maturation of DCs (and therefore initiate acquired immunity).