Regulatory T Cells & Tolerance

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Regulatory T Cells and Maintenance of Tolerance

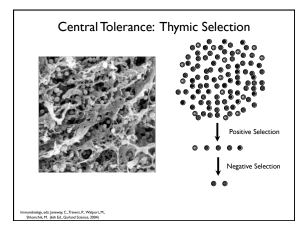
- Tolerance: *Lack* of an adaptive immune response to Ag despite exposure of lymphocytes to that Ag
 - Where is tolerance important?
 Discrimination of non-self from self
 e.g., virally infected from normal cell
 - Discrimination of harmless from dangerous
- •e.g., egg white from E. coli proteins in spoiled mayonnaise
- Ideal: Focus response on dangerous non-self
 Failure and a second and a second and a second a
- Failure: autoimmunity, allergy, transplant rejection

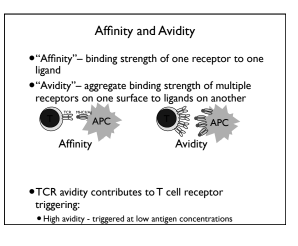
Autoimmunity: Distinguishing native tissue from foreign pathogen

- Innate System inherent in the receptors
- Directed at microbial molecules (PAMP's)
- \bullet Adaptive System \underline{not} inherent in the receptors
- Able to bind anything protein, carbohydrate, lipid
- Need safeguards to ensure non-reactivity with native (self) molecules that is, to maintain *tolerance*

Mechanisms Regulating Adaptive Response

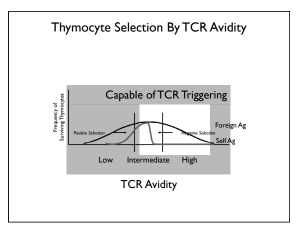
Central	Peripheral
Selection During Development	Responder Cell Intrinsic ("Recessive")
B Cells - bone marrow T cells - thymus	Ignorance Activation Induced Cell Death Anergy <u>Responder Cell Extrinsic ("Dominant")</u> Regulatory (Suppressor) T Cells





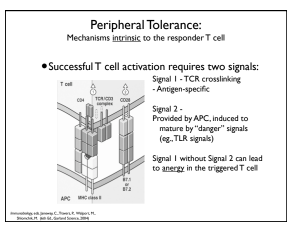
Thymocyte Avidity Profile is "Molded"

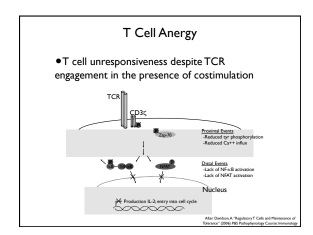
- •T cells mature from thymic precursors
- Specificities are "filtered" by positive and negative selection on self antigens
- Anti-self TCR avidities range from low to moderate
- Anti-foreign TCR avidities are "unfiltered" (range from undetectable to high)

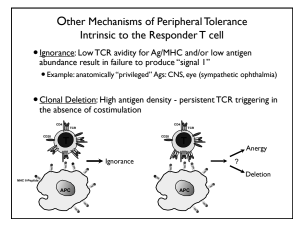


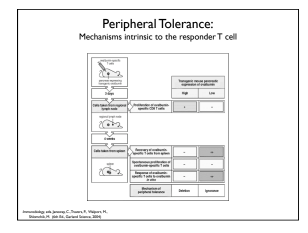
Thymic Self-Representation

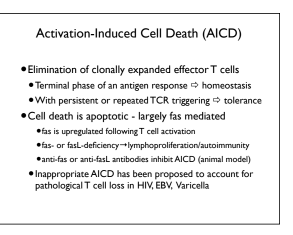
- AIRE: Autoimmune Regulator
- Gene highly expressed in thymic epithelium
- Encodes a transcriptional activator
- Induces expression of "ectopic" self proteins • Parathyroid-, retina- and ovary-specific
- APS Autoimmune Polyendocrine Syndrome • Clinical condition from mutation in AIRE
- Autoimmune attack on multiple endocrine structures (parathyroid, thyroid, adrenals, β-islets, gonads), vitiligo, alopecia

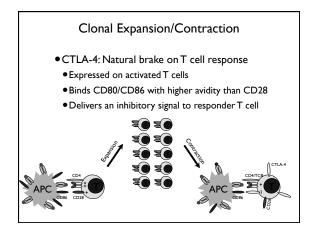


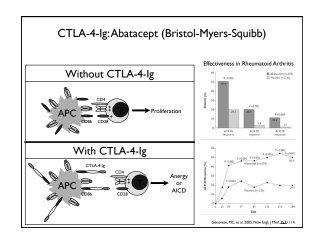


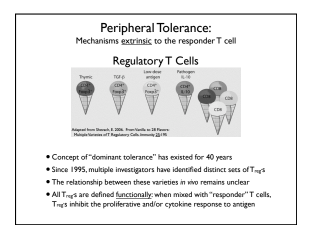


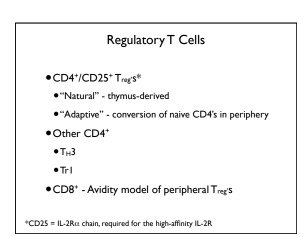












CD4⁺/CD25⁺ T_{reg}'s

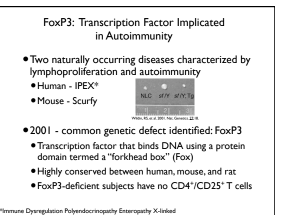
- I969 Early thymectomy (day 3-4) ⇒ autoimmune organ attack
 Prevented if mice received splenic CD4⁺ T cells from a normal adult
- 1995 Identification by Sakaguchi
 - Transfer T cells into athymic mice ⇒ normal immune function
 - Transfer CD25-depleted T cells into athymics ⇒ autoimmunity
 - thyroiditis, insulitis, gastritis, adrenalitis, arthritis, etc.
 - \bullet short time window to rescue by transfer of CD25+ fraction

CD4⁺/CD25⁺ T_{reg}'s (cont'd)

- 10% of circulating mouse CD4⁺ cells bear CD25 (<1% of CD8's)
- Upon triggering with CD3/CD28 crosslinking in vitro:
 no proliferation
- •no secretion of IL-2, IL-4, or IFN-γ
- •contact-dependent inhibition of local "responder" T cells
- Capable of self-renewal in vivo
- Constitutively express CTLA-4
- Dependent on IL-2 for maintenance of CD25 expression and regulatory phenotype
- •IL-2-deficiency and CD25-deficiency are both associated with autoimmunity

"Natural" Treg's Arise in the Thymus

- Proposed "third role" of the thymus (in addition to positive and negative selection)
- Alternative to cell death for thymocytes with significant avidity for self
 - \bullet Thymocyte TCR triggering is required for T_{reg} differentiation
 - CD28 costimulation is required for normal T_{reg} levels • Third, T_{reg}-specific signal - postulated
- T_{reg} induction is presumably driven by self recognition, however the T_{reg} repertoire is unknown

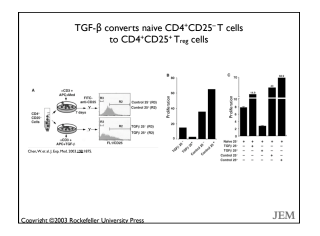


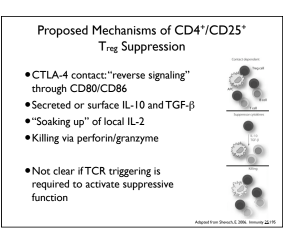
CD4⁺/CD25⁺ "Lineage Marker" FoxP3?

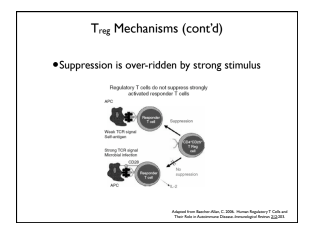
- Possible "master controller" for T_{reg} program
- FoxP3 represses TCR-triggered IL-2 transcription, induces CTLA-4 and CD25 transcription
- \bullet FoxP3+ T_{reg} function is stable transferrable by adoptive transfer from one individual to another (in mice)
- Mouse: FoxP3⁺ T cells are >90% CD4⁺/CD25⁺ and functionally suppressive

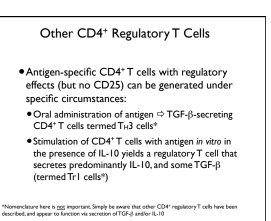
"Adaptive" Treg's Arise in the Periphery

- Naive CD4⁺ T cells
 ⇔ CD4⁺CD25⁺FoxP3⁺ under the following circumstances:
- TCR X-linking in presence of TGF- β (in vitro)
- Slow infusion low dose soluble antigen (in vivo)
- Adaptive Treg function
- Mouse potent suppressors of responder T cells (indistinguishable from natural Treg's)
- Human despite FoxP3 expression, suppression less consistently demonstrated; role unclear



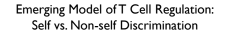






CD4⁺ T_{reg} as a Phenomenon

- Strengths
- Clear suppression of effector T cells
- Reproducible by multiple investigators
- Weaknesses
- Mechanism of "call to action" undefined
- How are inappropriate T cell responses recognized?
- Mechanism of suppressive effect remains undefined
- Many of the defining principles derived in mice • Human [?] = Mouse



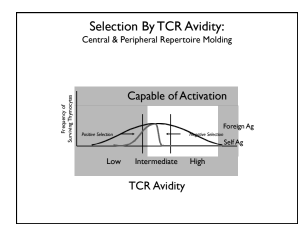
- Key Facts we know that:
- TCR avidities for foreign antigen range low to high
- <u>but</u> avidities for self are restricted (thymic selection)
- The T cell can "sense" avidity basis of thymic selection

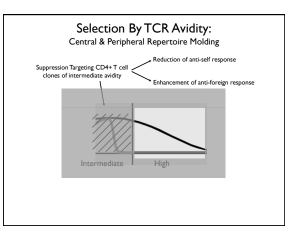
 Problem: Intermediate avidity self-reactive T cells in the periphery may encounter circumstances with high enough levels of self Ag presentation and costimulation to become activated ⇒ autoimmunity

T Cells Self-Police

- CD4+ T cells triggered following intermediate avidity interaction with MHC/peptide "flag" themselves
 - Based only on avidity completely independent of what the Ag actually is
- One of these "flags" has been identified: peptide derived from HSP-60 displayed on the cell surface in the groove of an HLA-I like molecule called HLA-E
- CD8⁺ T_{regs} recognize HSP-60⁺ CD4⁺ effector cells and target them for suppression

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Summary

- Self and non-self are *relative* terms. For the lymphocyte, "self" is best defined as those antigens present during the cell's development (in bone marrow or thymus). Thymocyte development can only be "molded" on self-antigens. Every mature T cell has a narrow range of TCR avidities for self but a wide range of avidities for foreign proteins.
- T cells emerging from the thymus may have high enough avidity for self proteins as to permit activation under circumstances of high antigen concentration and co-stimulation. Peripheral tolerance mechanisms are therefore critical to dampening the self-reactivity of T cells.
- 3. Responder-intrinsic tolerance mechanisms include ignorance (low TCR avidity, low Ag conc.); anergy ("signal I w/o signal 2" ⇒ unresponsiveness); and AICD (T cell death due to increased sensitivity to apoptosis following activation; fas-dependent). CTLA-4 is critical to maintenance of T cell homeostasis (overall cell number), and may also function in AICD. CTLA-4. Ig harnesses this regulatory function for therapeutic use. Dysfunction in native fas or CTLA-4 lead to lymphoproliferation/autoimmunity.



- 4. Responder-extrinsic tolerance mechanisms are those mediated by committed regulatory cells. The most widely studied of these are CD4*/CD25 Trags.a lineage that appears to require FoxP3 expression. These cells arise in both thymic development (natural) and in the periphery (adaptive) and require IL-2 for survival. The mechanism by which they target responder cells in unknown. Contact-dependent and -independent mechanisms of suppression have been implicated, involving CTLA-4 and TGF-β.
- 5. Other CD4* regulatory cells include T_{H3} cells (oral antigen; TGF- β -secreting); Tr1 cells (antigen stim. in presence of IL-10; IL-10-producing).
- 6. One population of regulatory T cells that expresses CD8 appears to target CD4⁺ responder cells that have been activated by a TCRantigen/MHCII interaction of *intermediate* avidity. This mechanism, so far demonstrated most definitively in mice, is the one mechanism to date that provides a rationale for self/non-self discrimination on a functional level.