

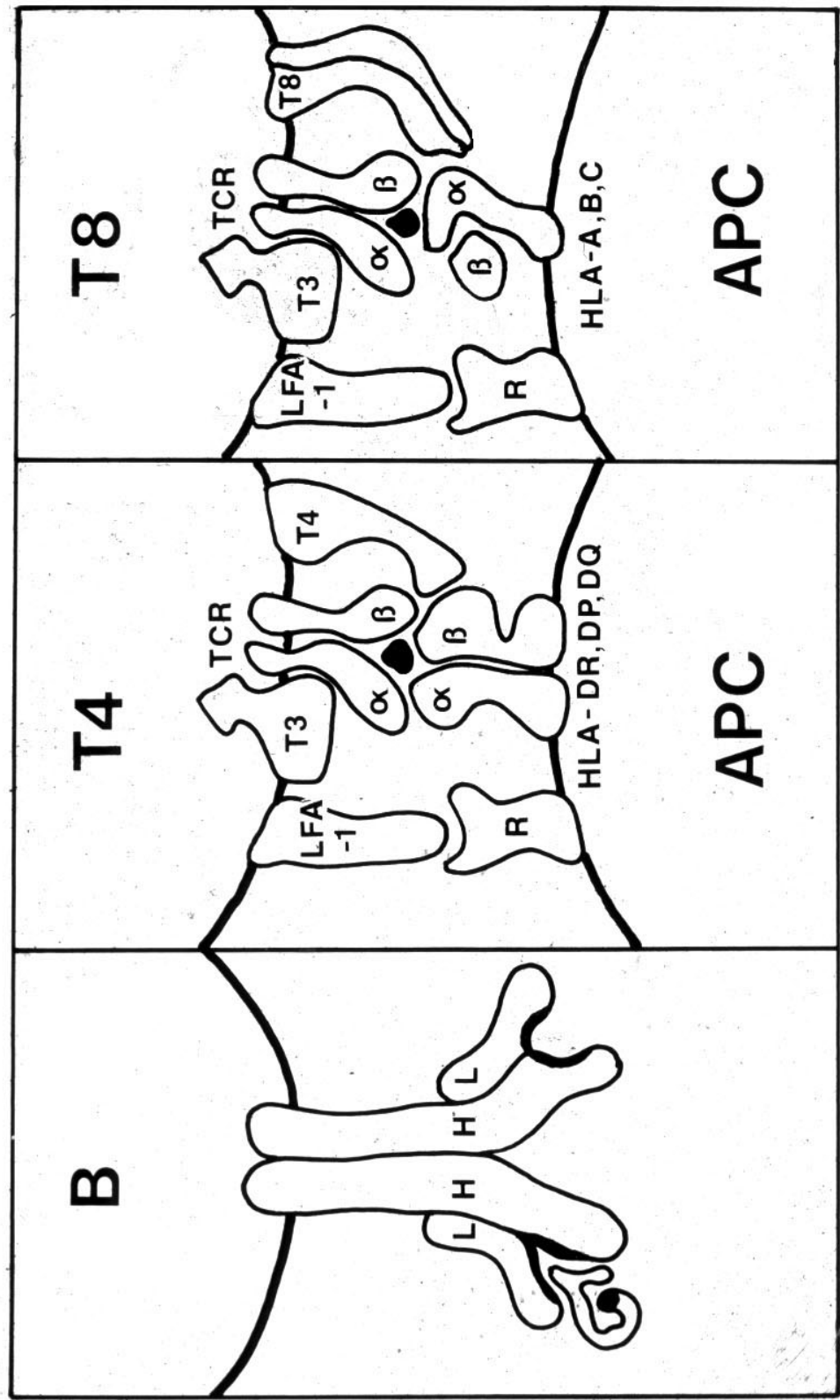
Basis of Immunology and Immunophysiopathology of Infectious Diseases

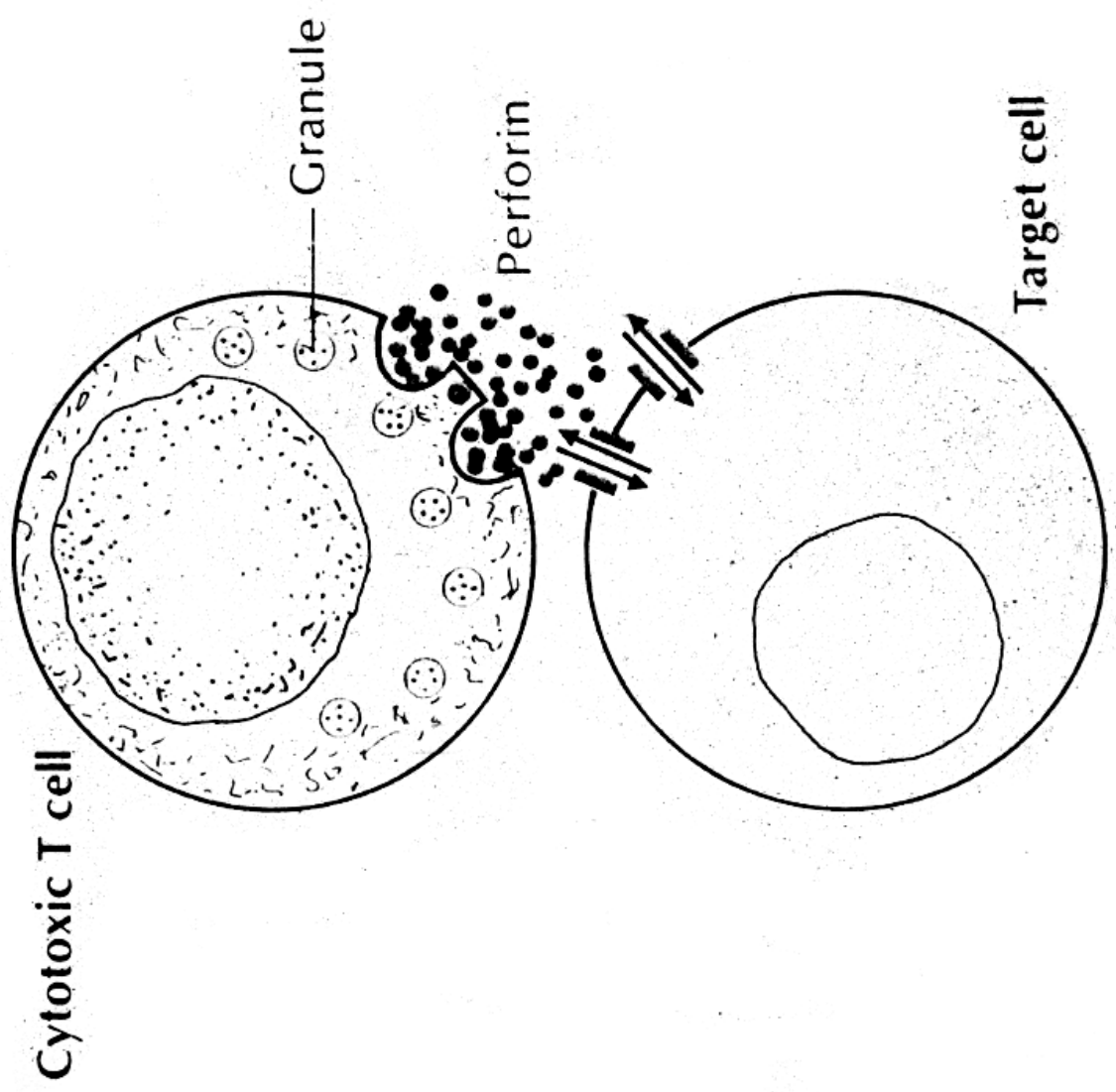
Jointly organized by
Institut Pasteur in Ho Chi Minh City and Institut Pasteur
with kind support from ANRS & Université Pierre et Marie Curie

January 24 – February 5, 2005
at the Institut Pasteur in Ho Chi Minh City, Vietnam

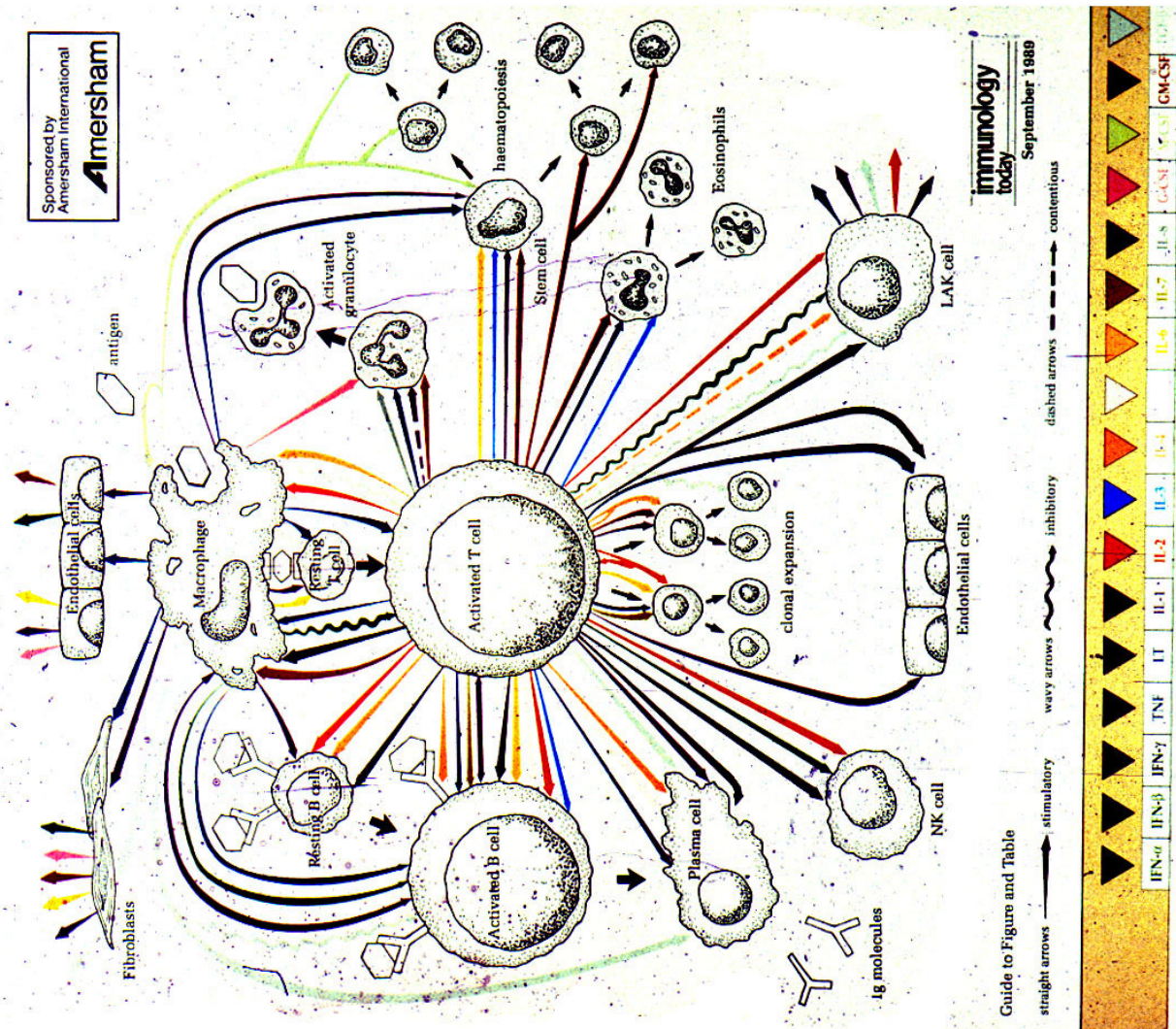
Lecture :
Activation of peripheral T lymphocytes
Prof. Jacques Louis

January 28, 2005



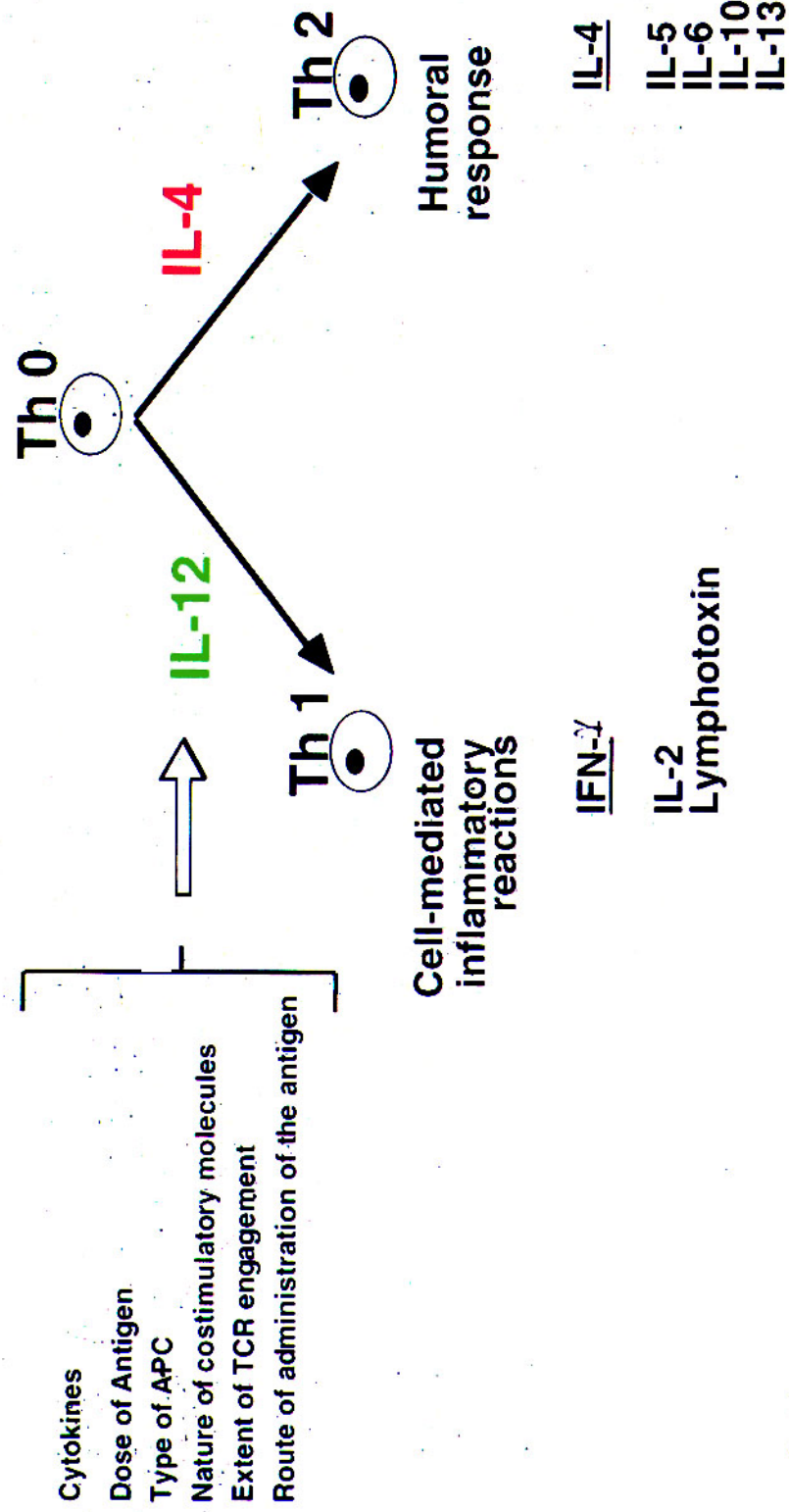


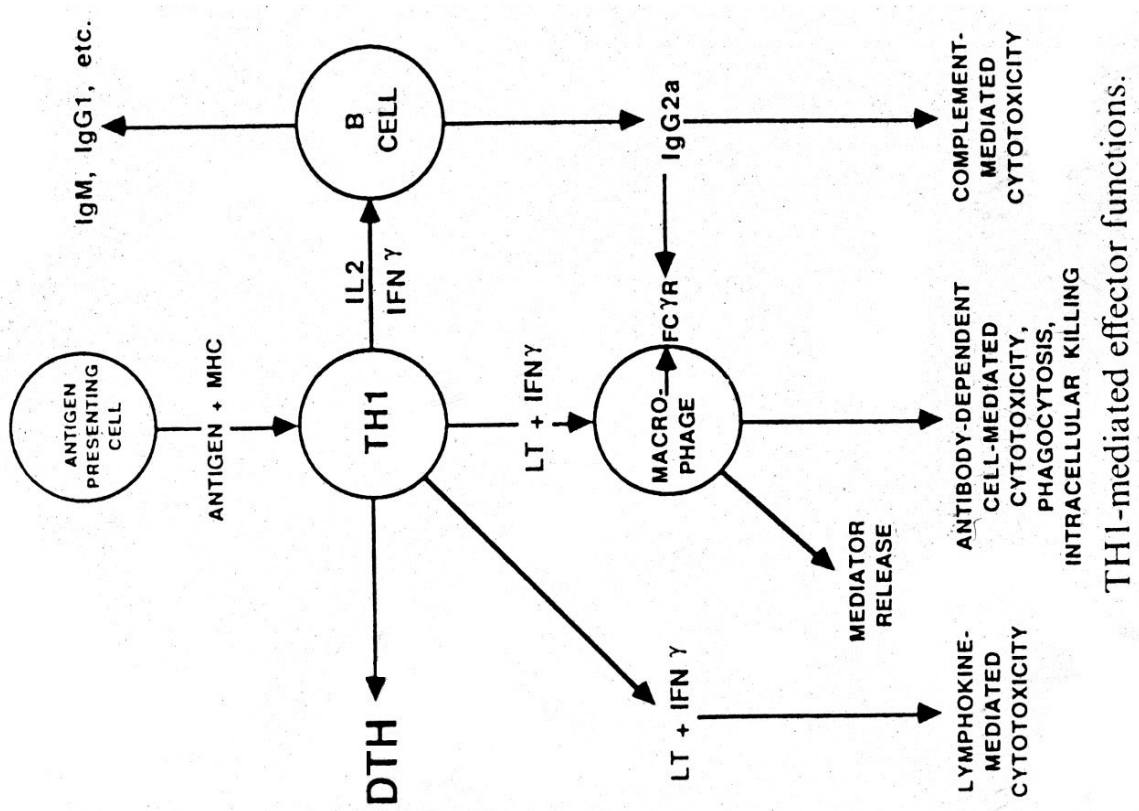
▼▼ The Cytokine Network ▲▲



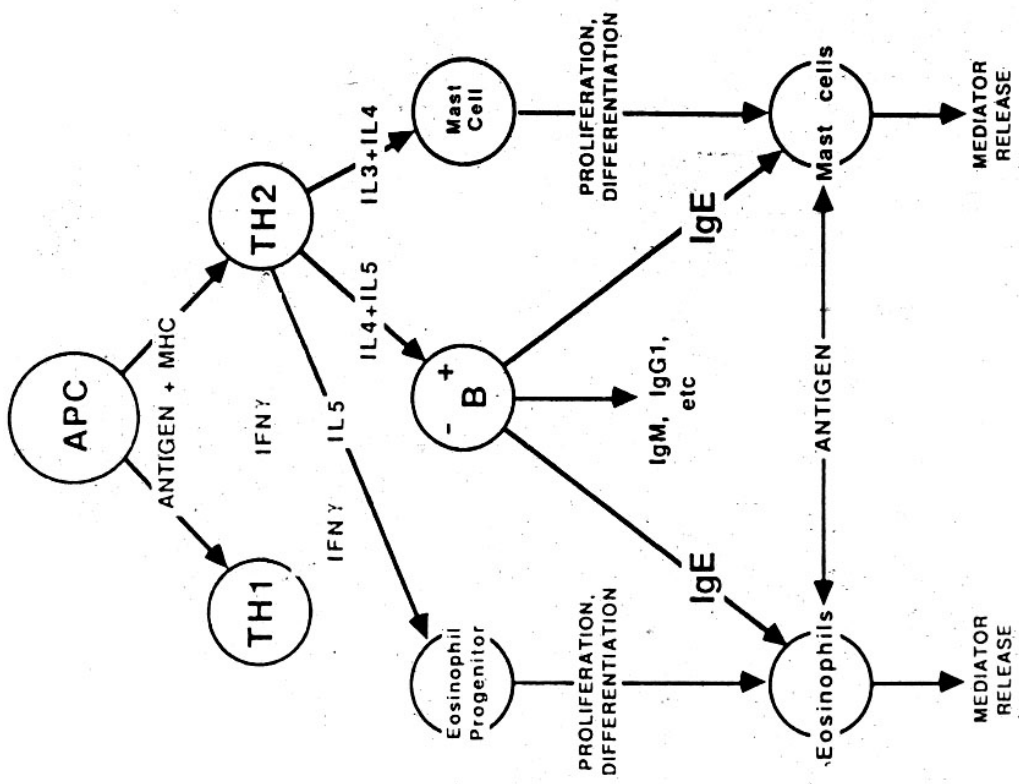
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Th cells can develop into two different subsets of effector cells





TH1-mediated effector functions.



Th1 and Th2 regulation of IgE. Stippled arrows indicate inhibitory effects, and solid arrows show stimulatory effects.

HEAVY CHAIN ISOTYPE SWITCHING INDUCED BY CYTOKINES

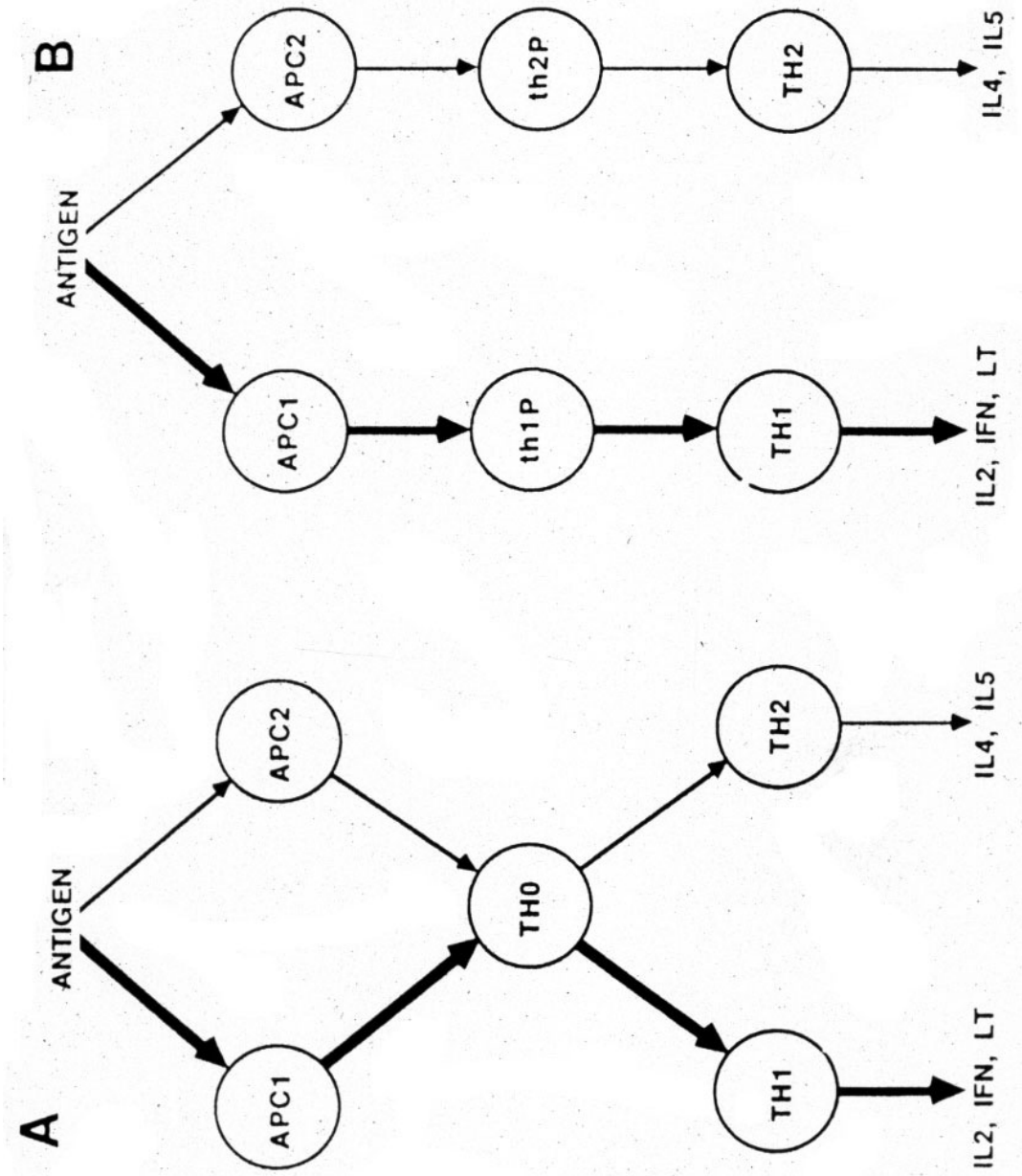
B cells cultured with:

Polyclonal activator *Cytokine*

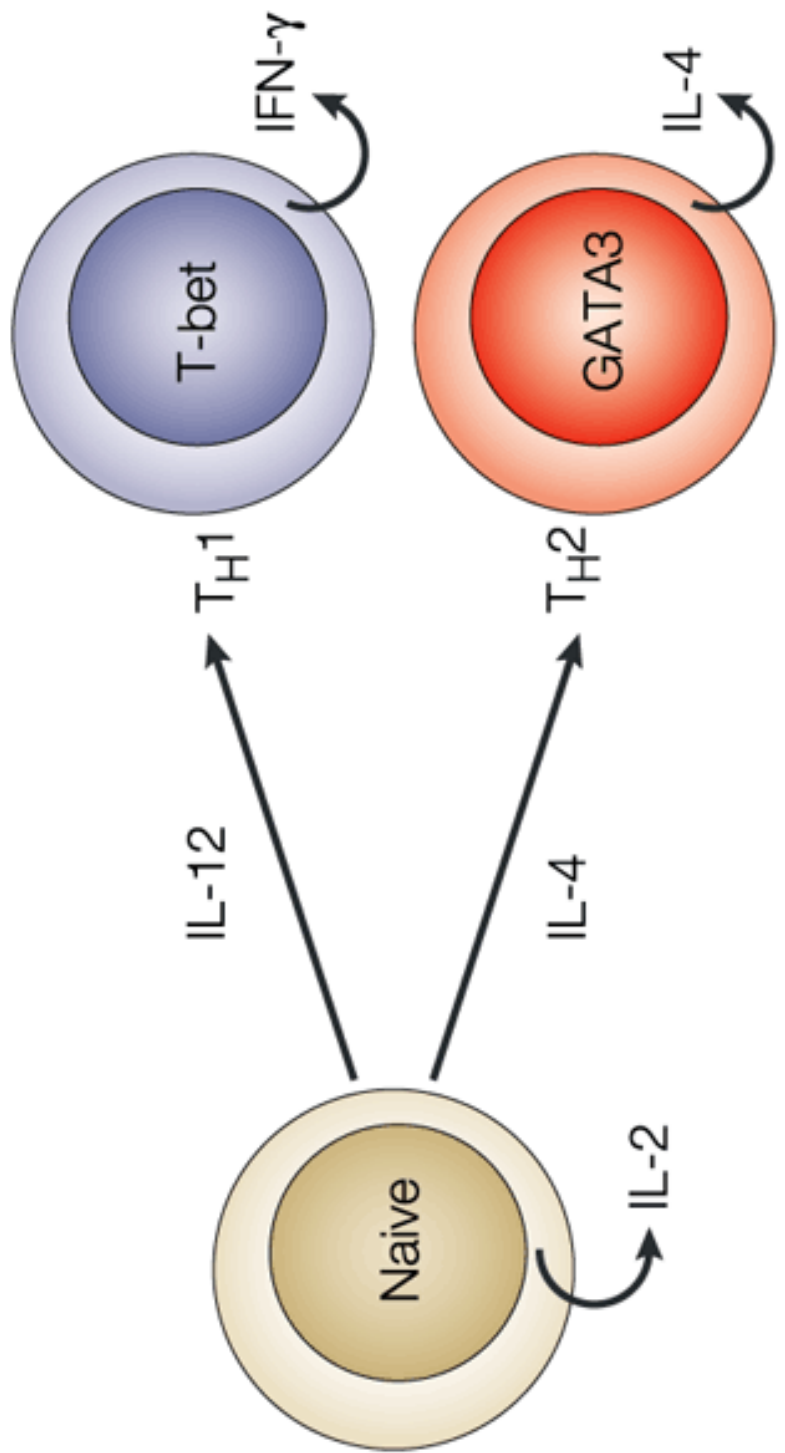
Ig isotype secreted
(% of total Ig)

		<i>IgM</i>	<i>IgG1</i>	<i>IgG2a</i>	<i>IgG</i>	<i>IgA</i>
LPS	None	85	2	<1	<1	<1
LPS	IL-4	70	20	<1	5	<1
LPS	IFN- γ	80	2	10	<1	<1
LPS	TGF- β + IL-2	75	2	<1	<1	15

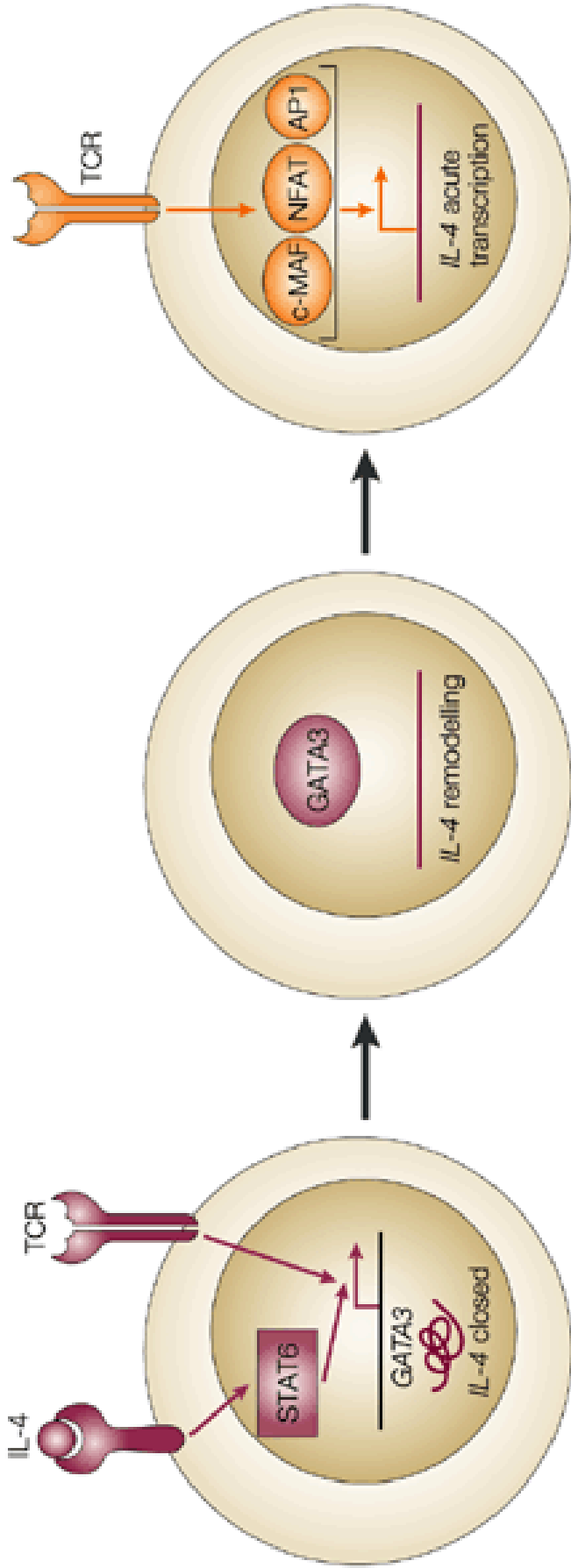
Adapted from Abbas, Lichtman, & Pober, Cellular and Molecular Immunology



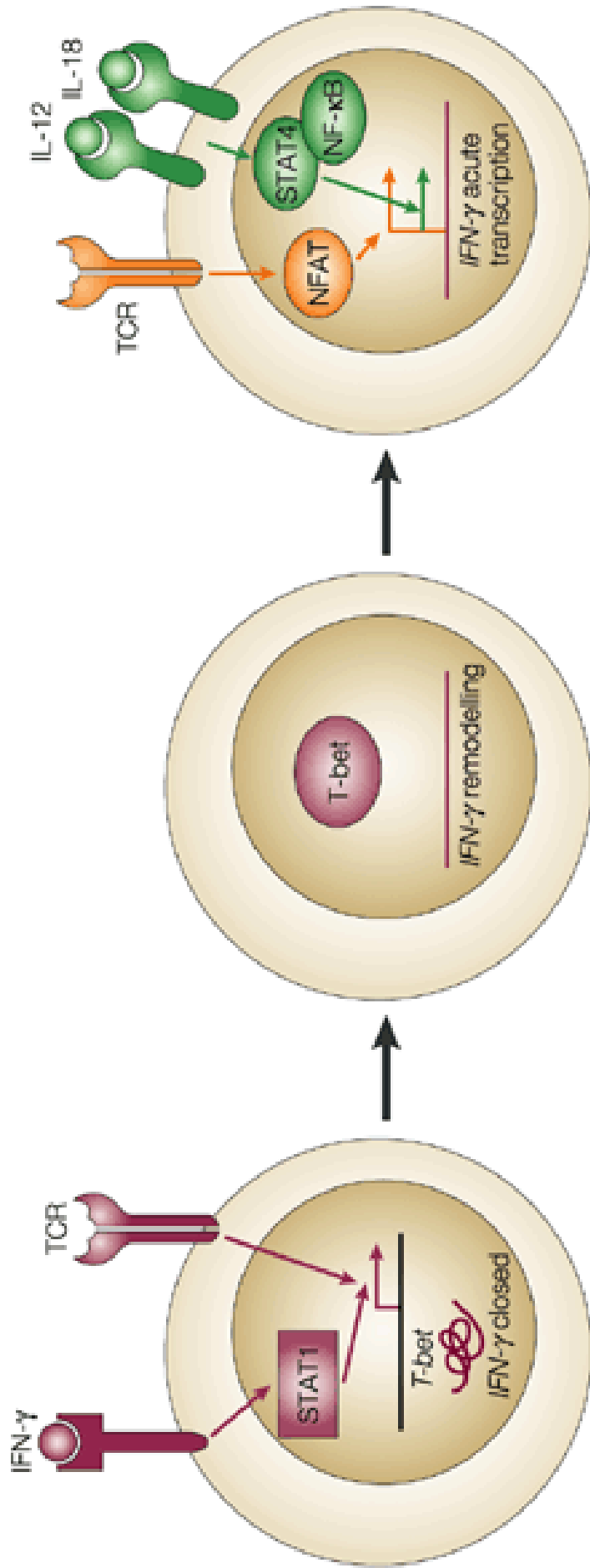
Possible TH1 and TH2 differentiation pathways.

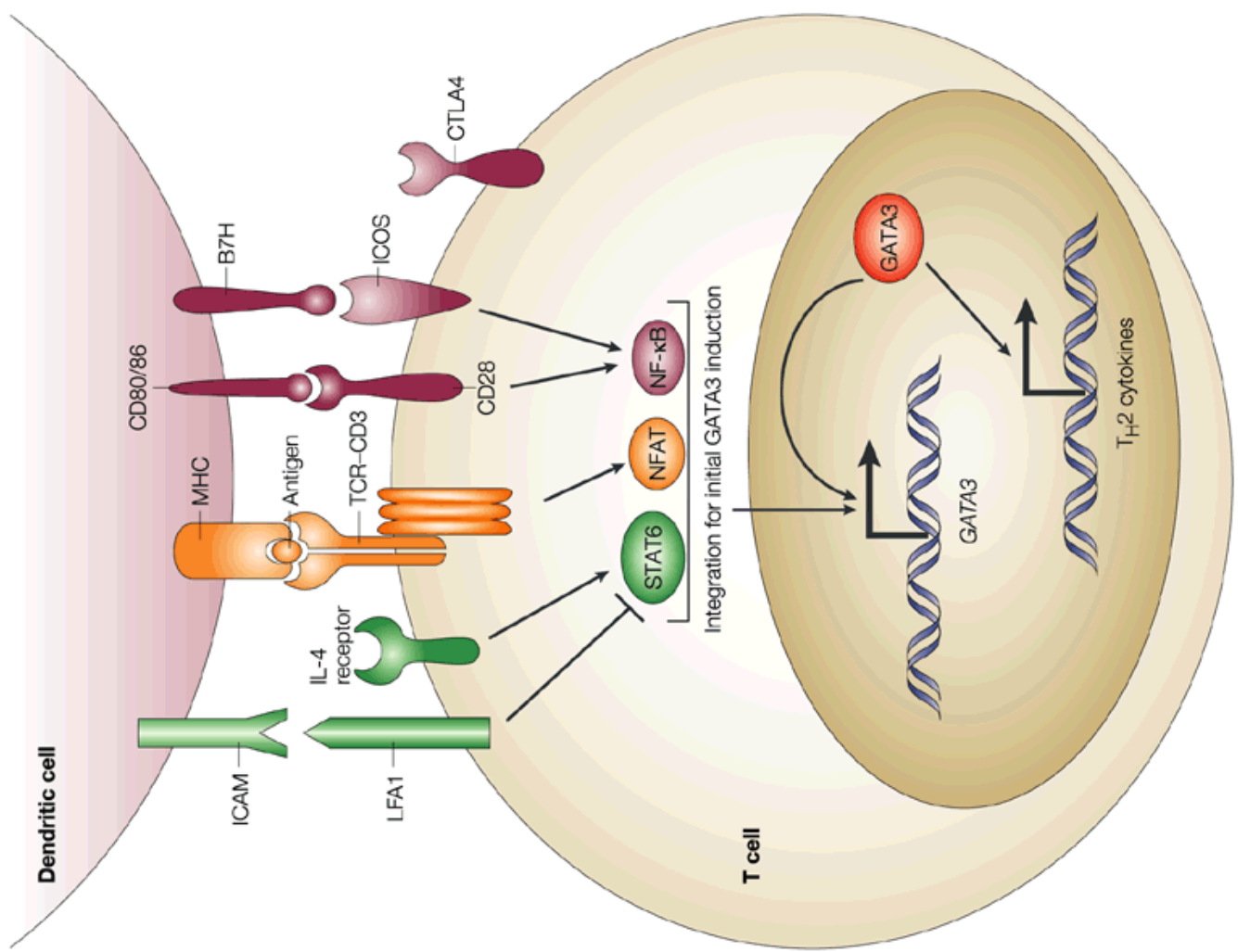


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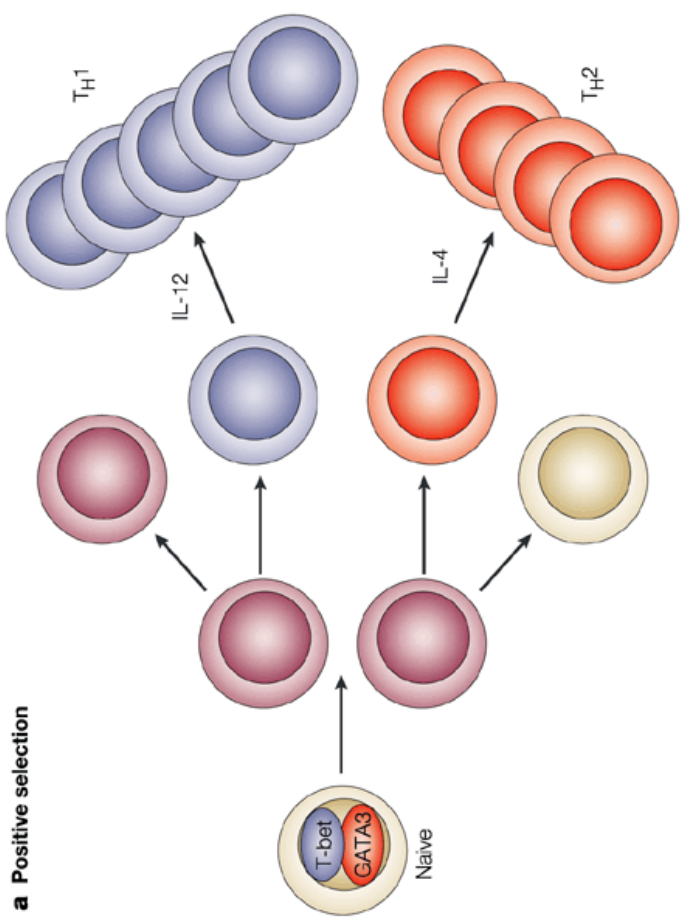


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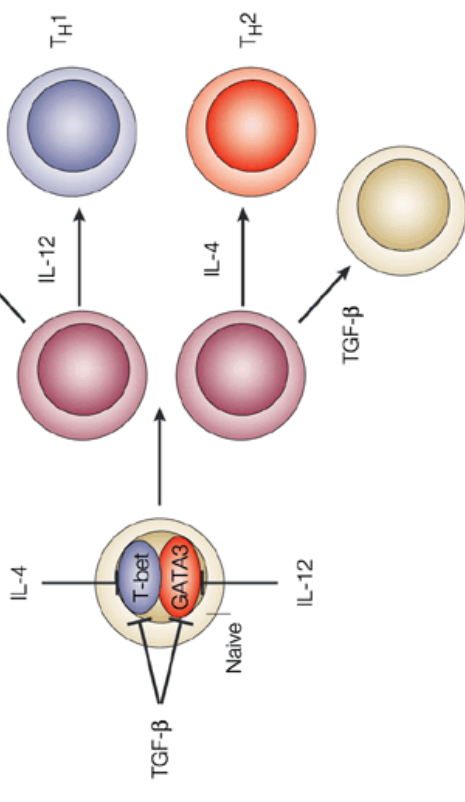


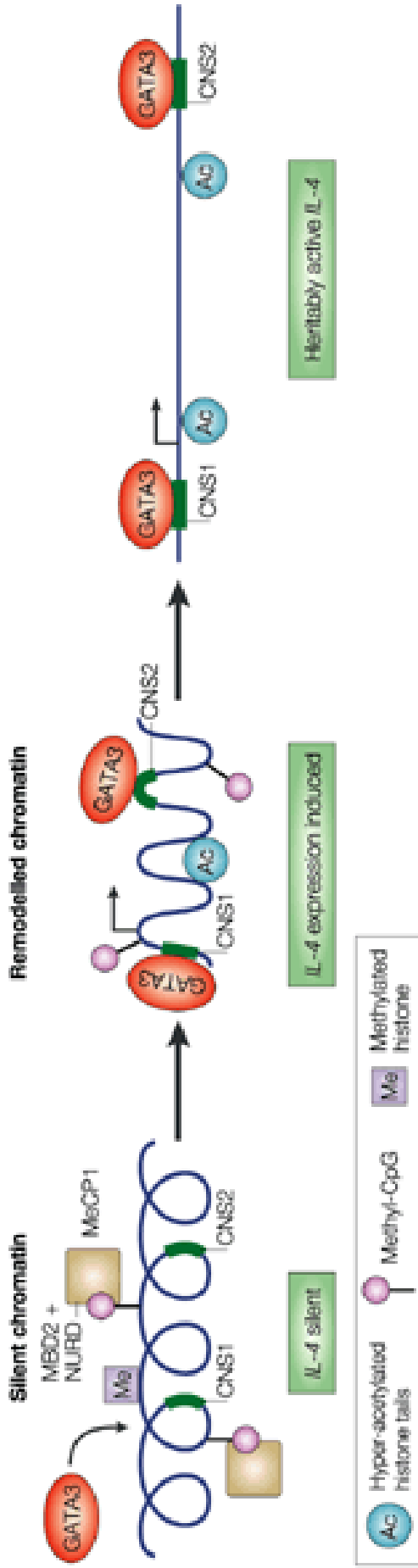


a Positive selection

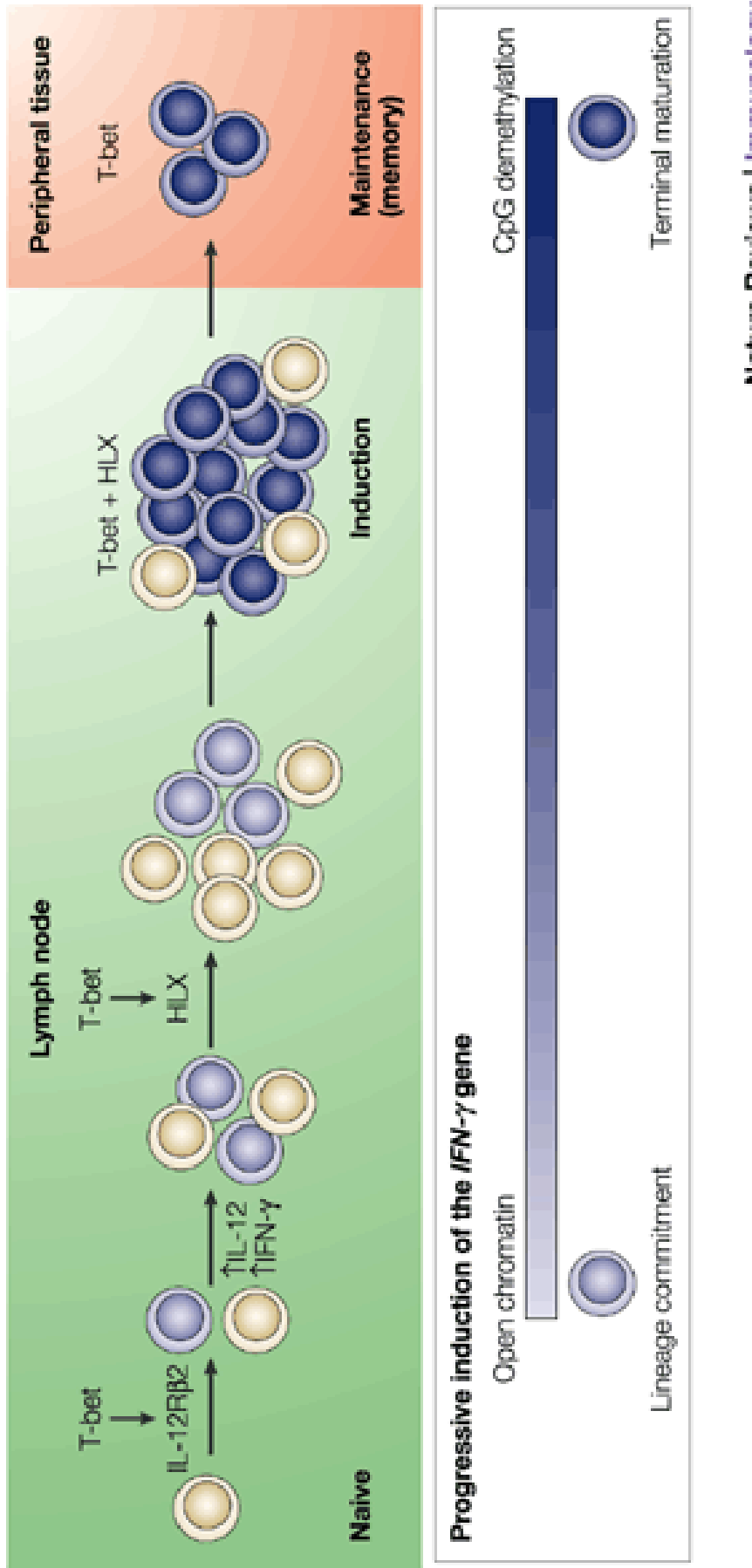


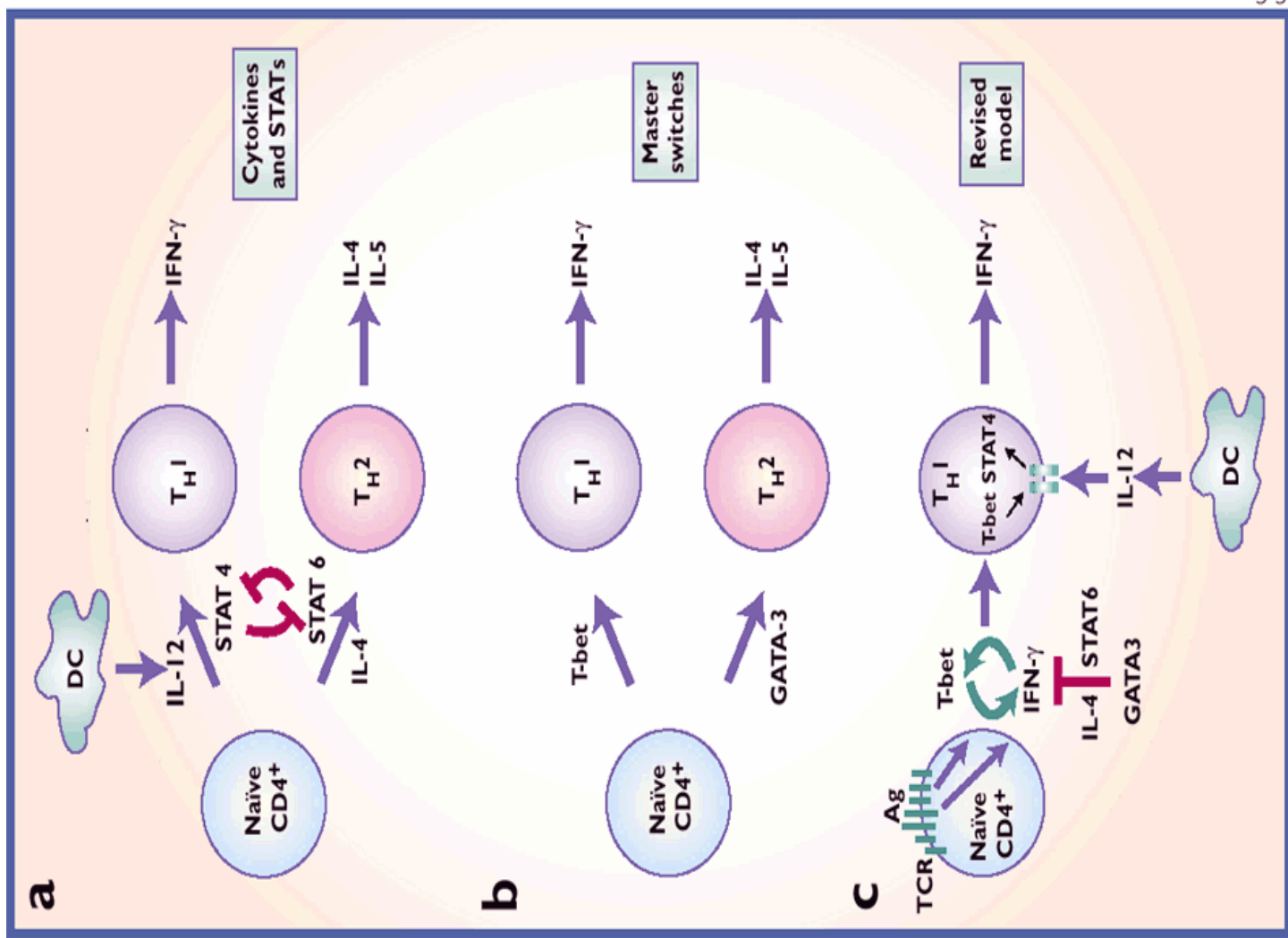
b Negative selection

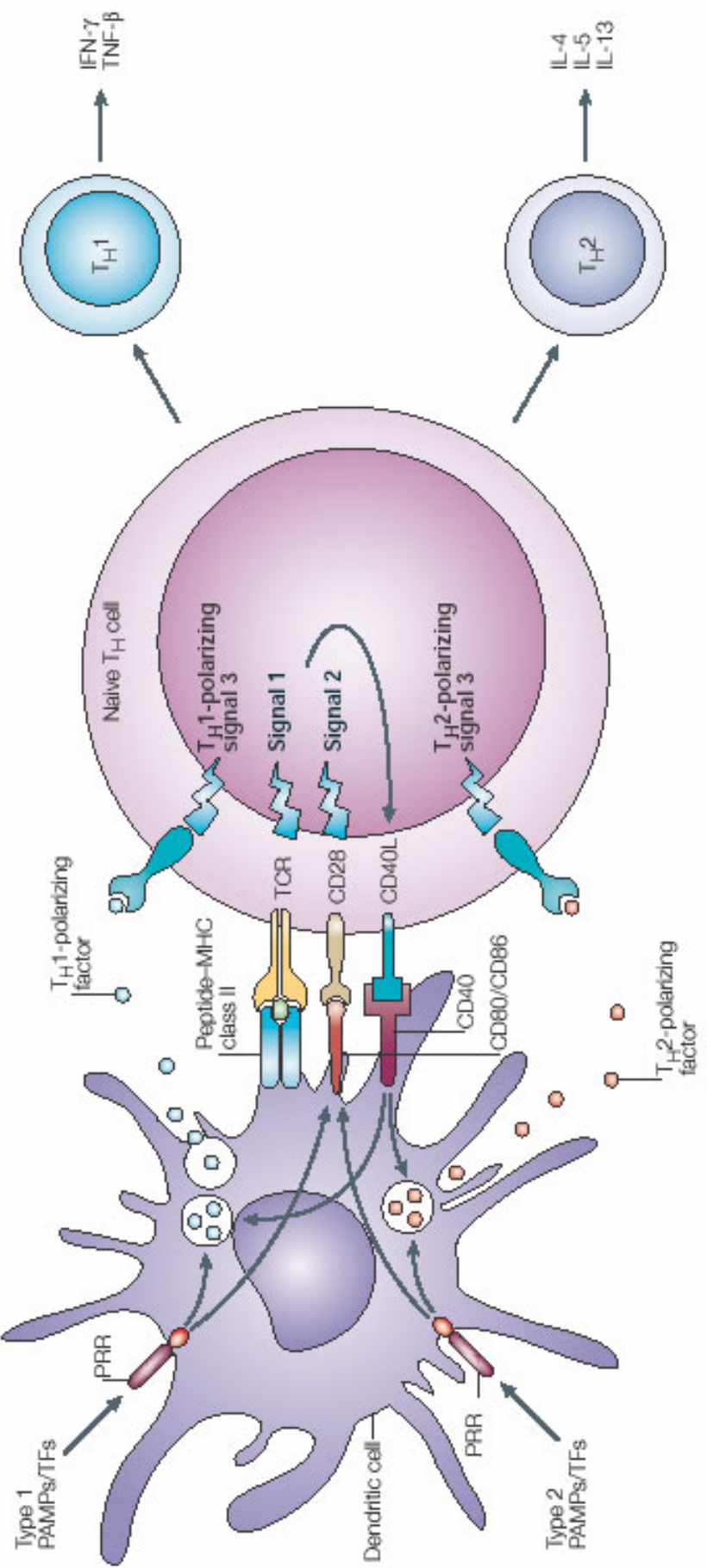




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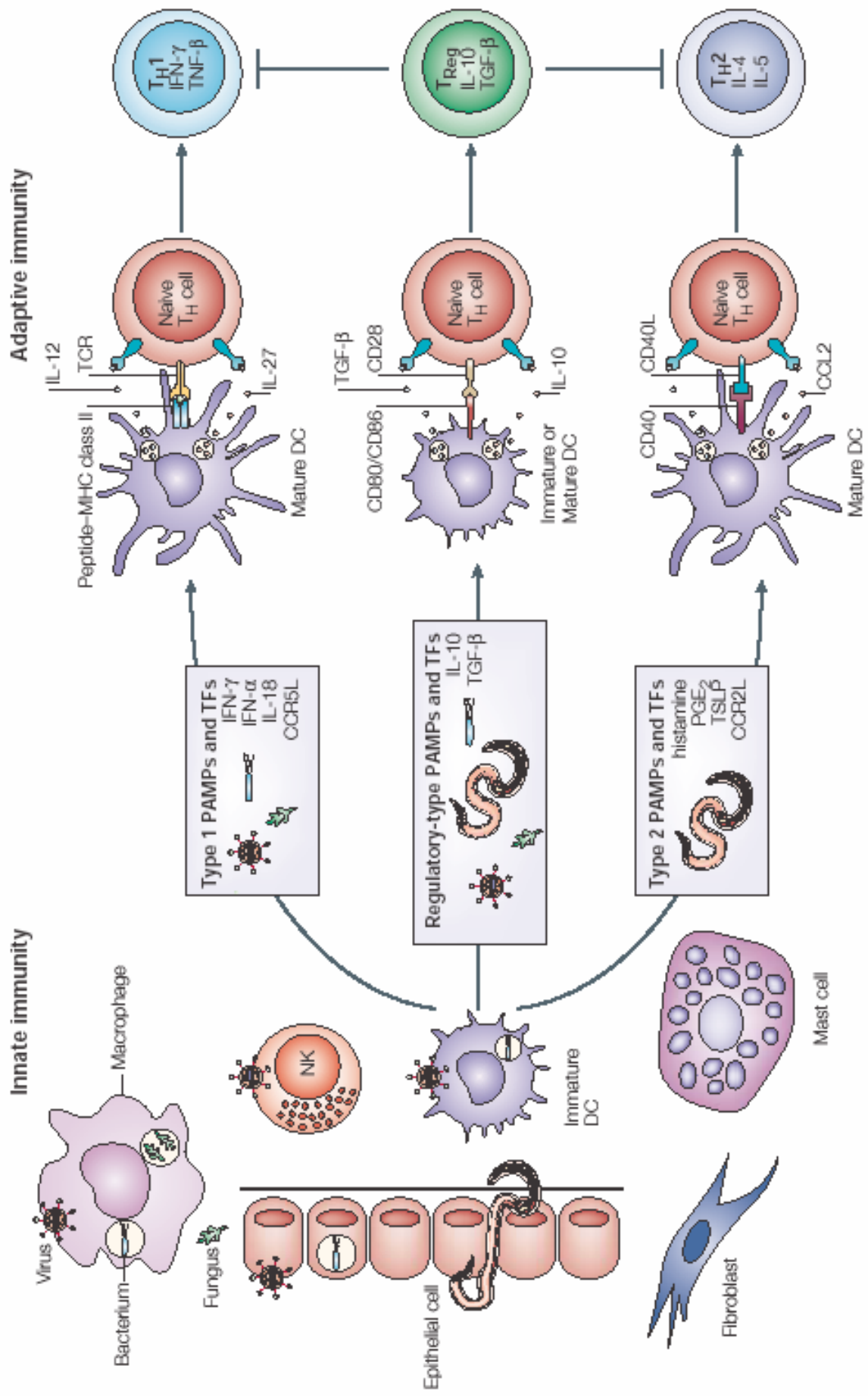
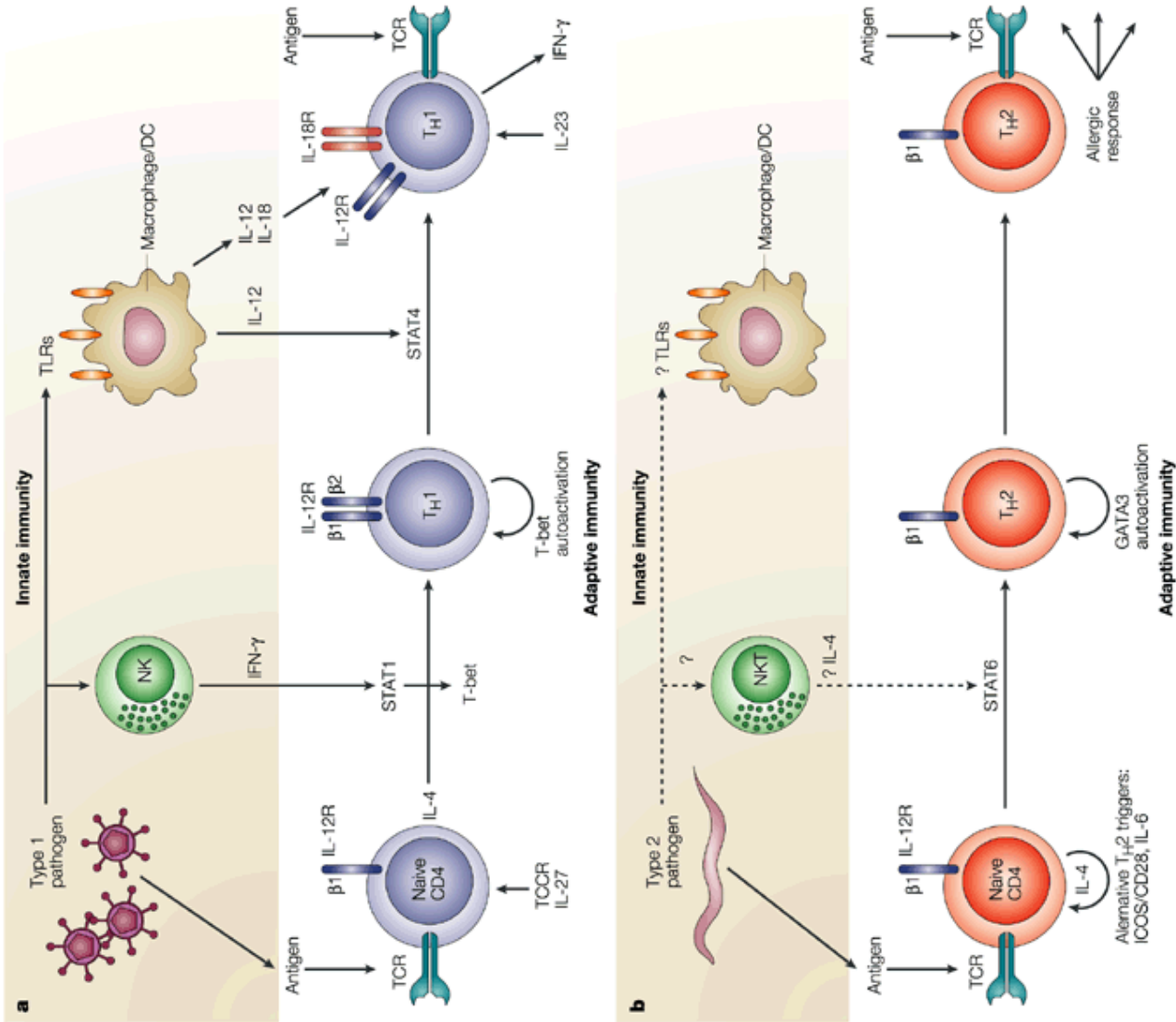


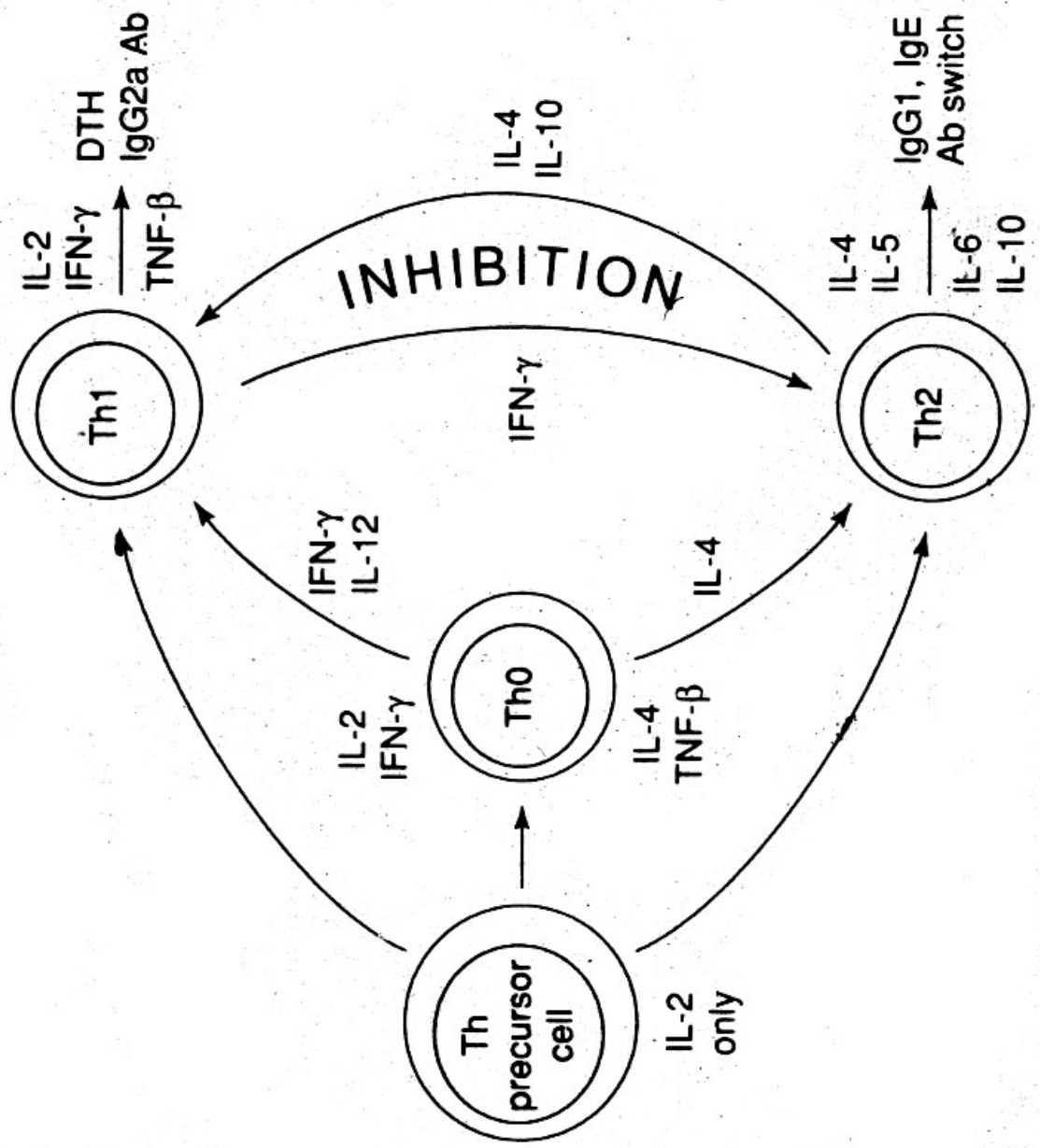
Table 1 | Type 1, type 2 and regulatory-type tissue factors

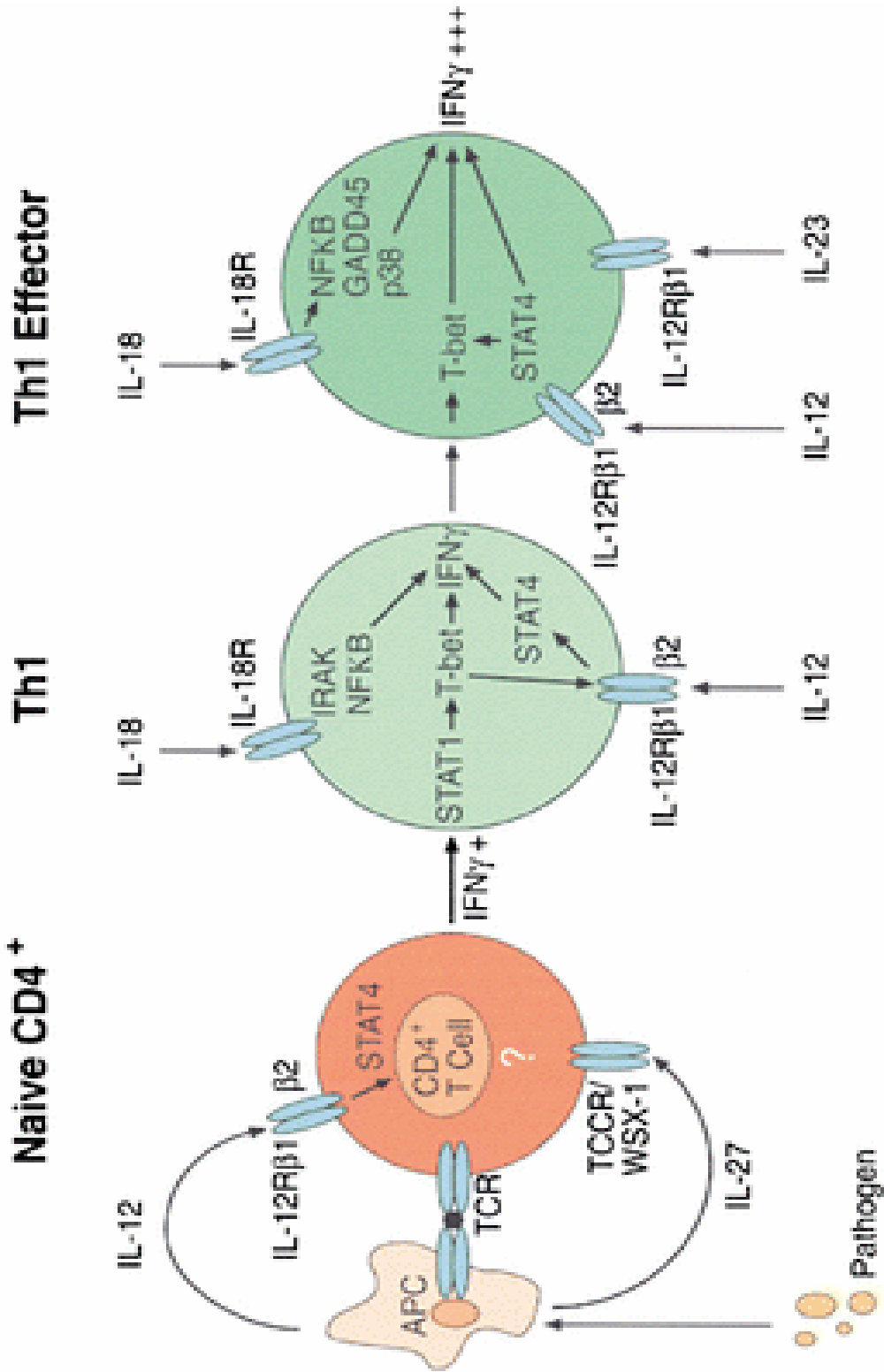
Tissue factors	Type 1	Type 2	Regulatory
Cytokines	*IFN- γ ⁸⁸ , *IFN- α/β ¹²⁰ , IL-12p70, TNF- β , IL-18 (REF. 89), *IL-27	IL-4, IL-5, IL-9, IL-13, IL-25, *TSLP ⁹²	*IL-10 (REF. 94), *TGF- β ⁹⁵ , TSP1
Chemokines	CxCL9, CXCL10, CCL21	*CCL2, *CCL7, *CCL8, *CCL13 (REF. 93), CCL17	
Co-stimulatory factors	ICAM1	OX40L	PD1/PD2L, CTLA4, GITRL, CD47, SIRP- α
Eicosanoids		*PGE ₂ (REF. 91)	
Others		*Histamine ⁹⁰	

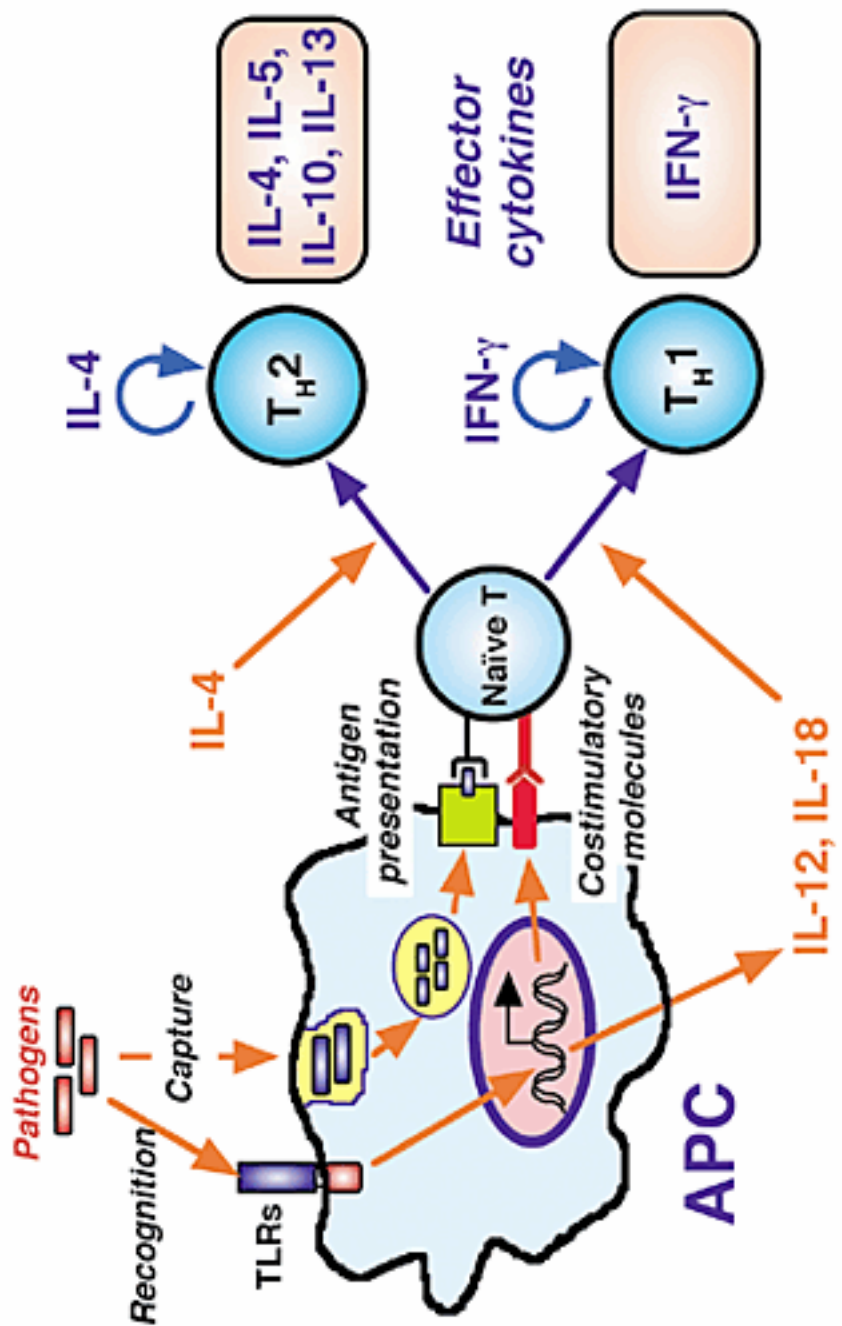


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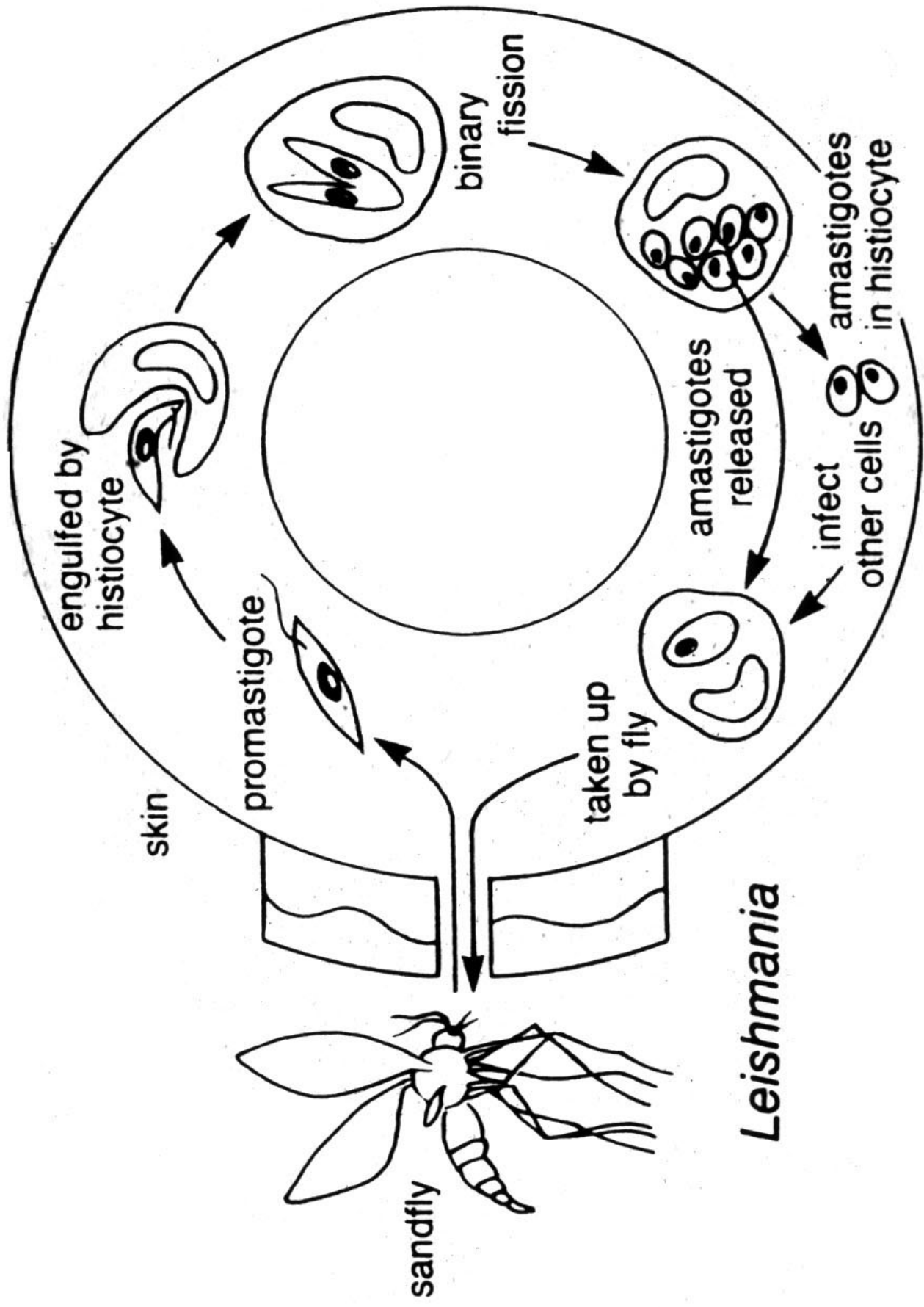


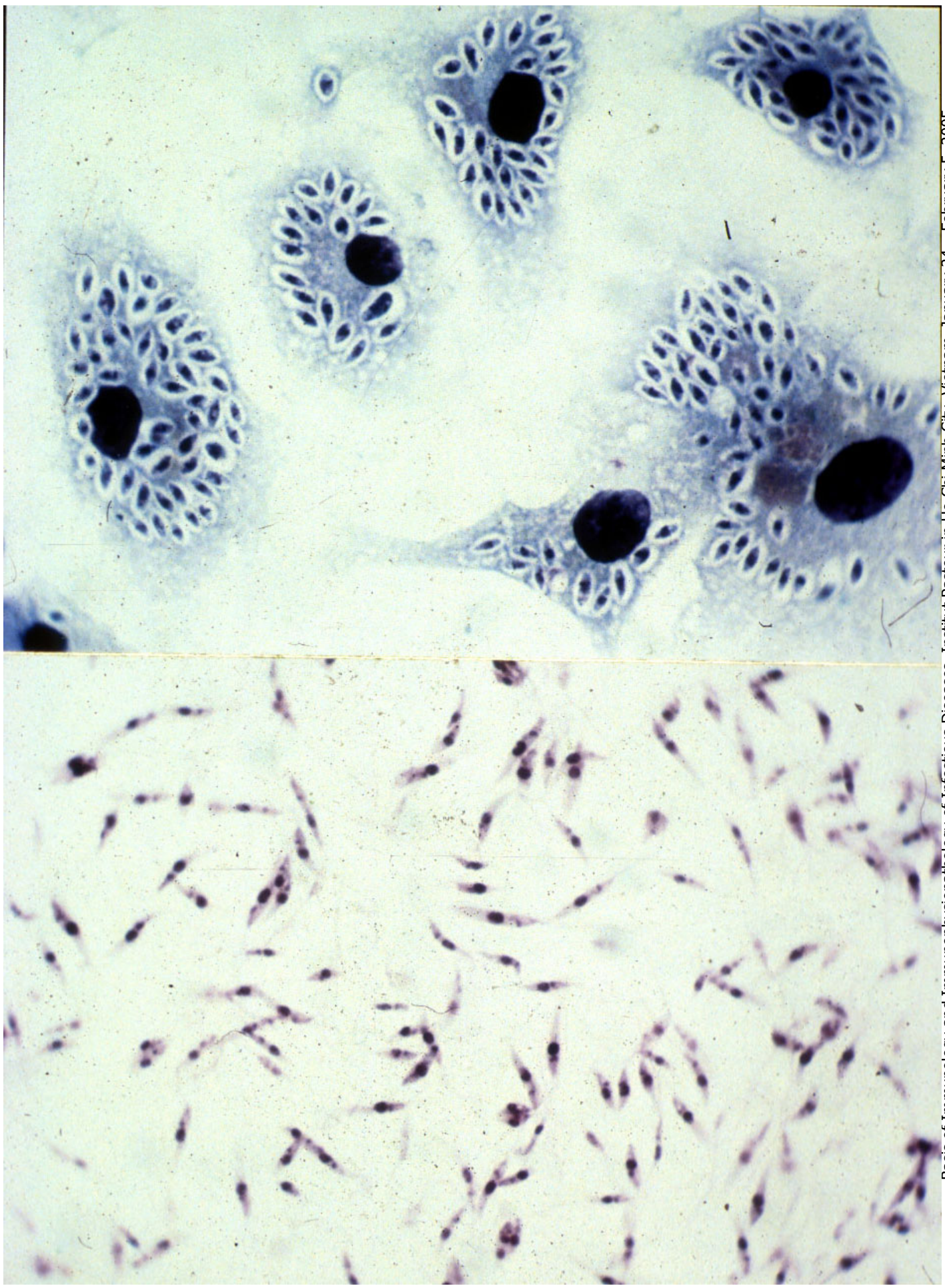




Toll and control of adaptive $CD4^+$ T cell responses

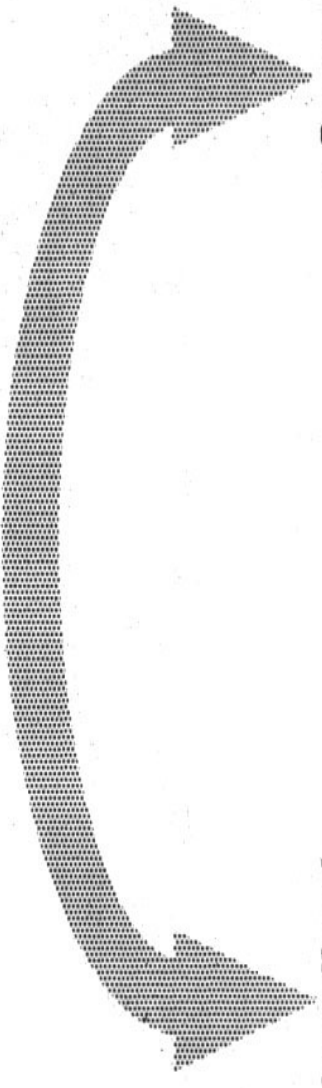
- * Th1 responses are Toll-dependent responses
- * Th2 responses are Toll-independent responses





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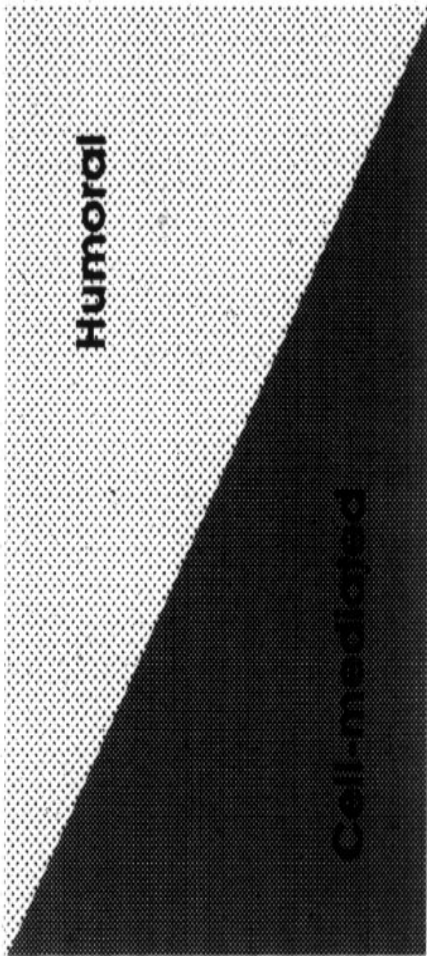
Spectrum of clinical manifestations of infection with *Leishmania*

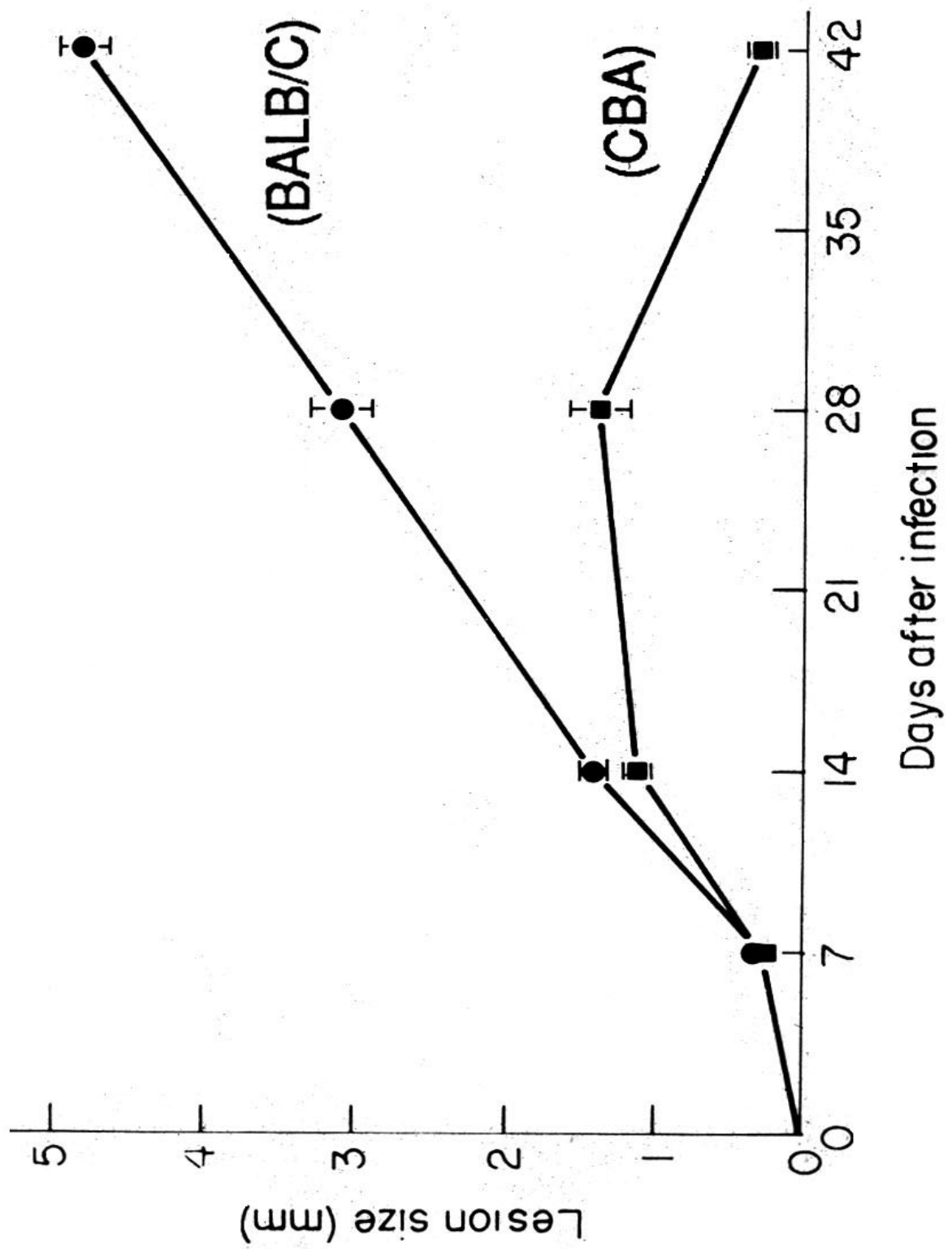


✓ Localized
cutaneous
lesion

✓ Generalized
systemic
disease

Type of immune response which predominates



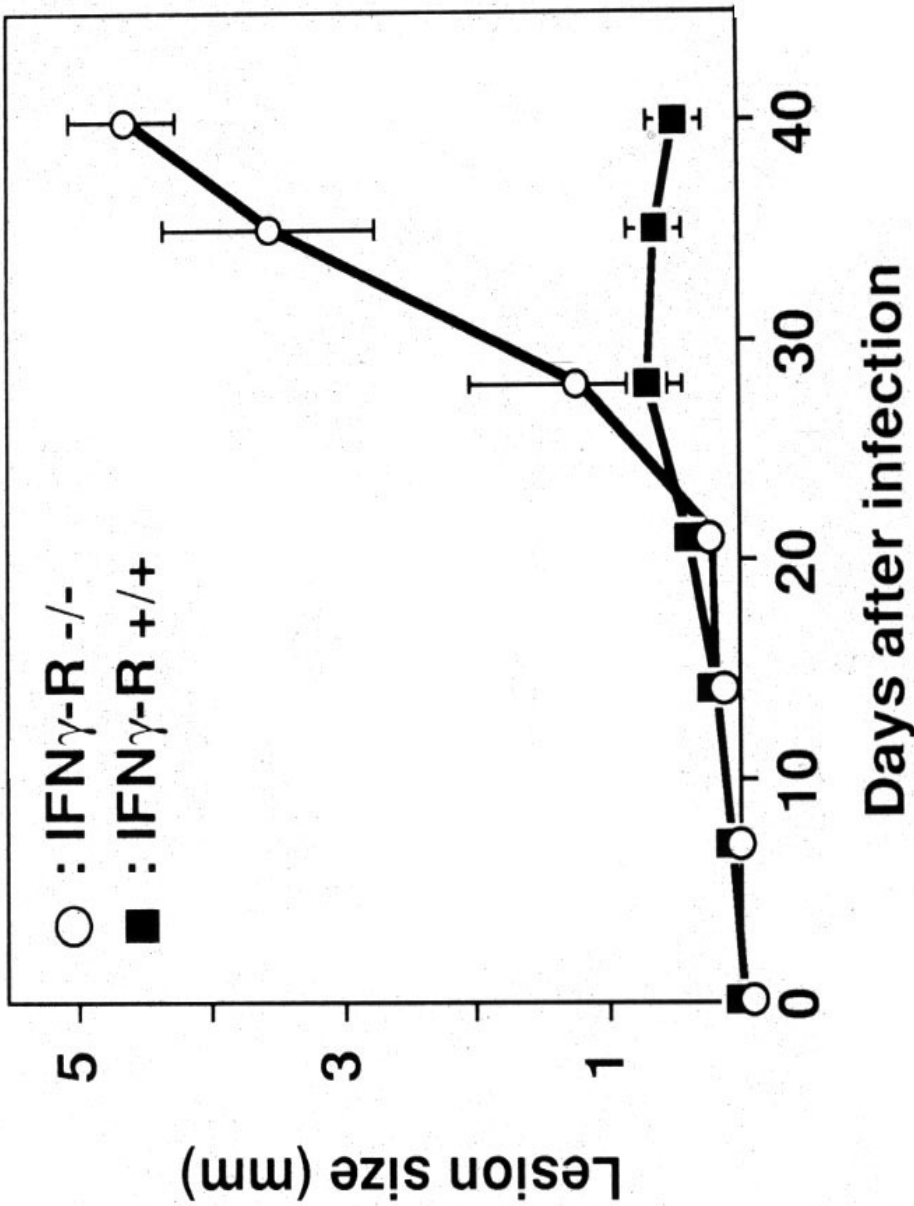


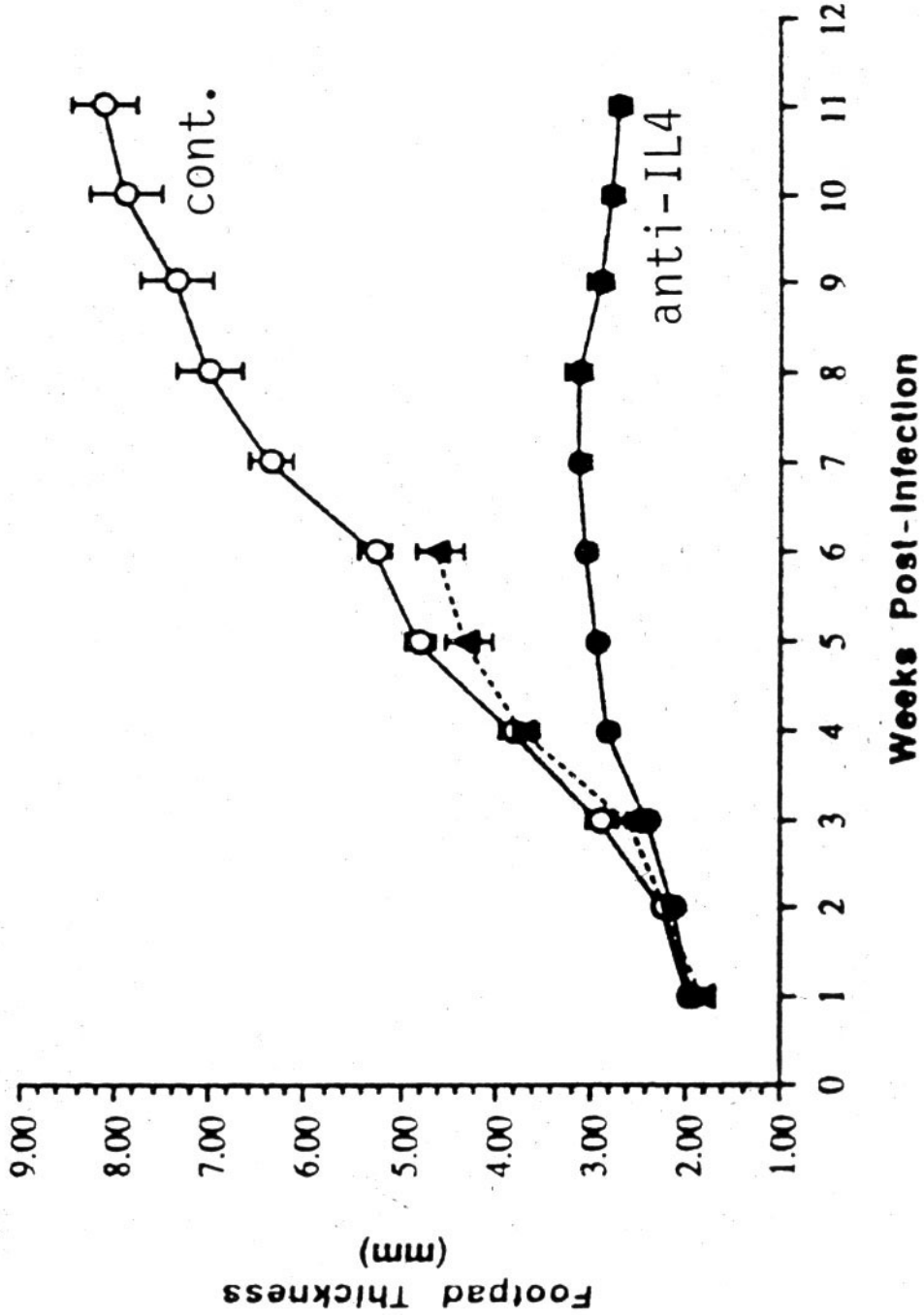
Characteristics of the T Cell Response in Mice Infected with *Leishmania major*.

Resistant Phenotype: Preferential expansion of TH1 cells.

Susceptible Phenotype: Preferential expansion of TH2 cells.

Development of Lesion induced by *L.maior* in Mice From a Genetically Resistant Strain Lacking IFN γ receptor





From Sadick et al.: J.Ex.Med., 1990, 171, 115

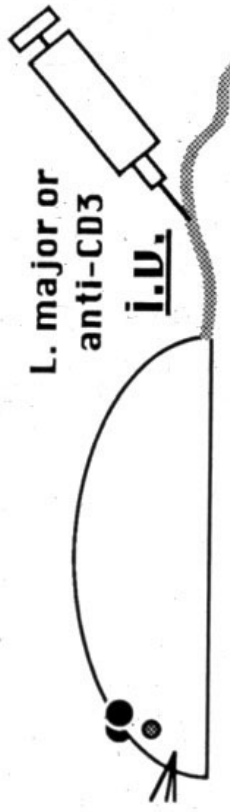
Factors Influencing the Functional Differentiation of CD4 T Cells

- ✓ Antigen structure
- ✓ Route of administration of antigen
- ✓ Genetic background
- ✓ Type of antigen presenting cells
- ✓ Costimulatory signals
- ✓ **Cytokines environment**

Importance of cytokines in the differentiation of CD4⁺ T cell precursors towards the Th1 or Th2 functional phenotype

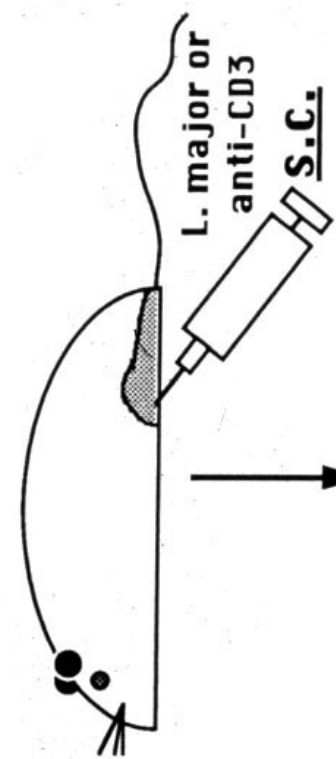
- IL-4:** Critical for priming CD4⁺ T cells to become IL-4 producers (Th2)
- IL-12:** Enhances priming for IFN γ -producing CD4⁺ T cells (Th1)
- IFN γ :** Plays a role in Th1 cell development, but is not sufficient. Differences between strains of mice in the IFN γ requirement for the development of Th1 responses?

Protocol 1



spleen removed
after 1.5, 4, 24, ... hrs

Protocol 2

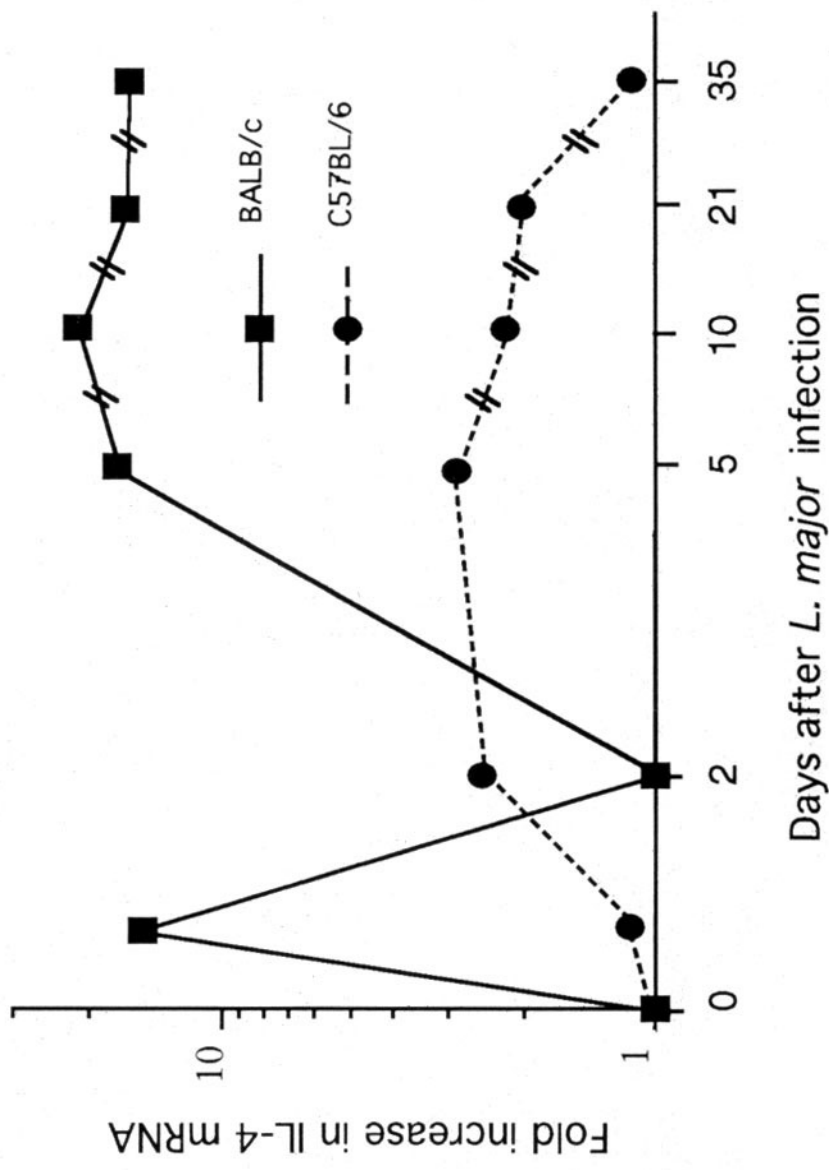


draining LN removed
after 6, 16, 24, ... hrs

mRNA extracted

**Quantitative RT-PCR
for IL-4 message**

**Kinetics of IL-4 mRNA expression in lymph nodes
of susceptible and resistant mice following
infection with *L. major***



Possible cellular origin of cytokines involved in the differentiation of CD4⁺ T cell precursors

IL-12:

Macrophages

IFN γ :

NK cells

CD4⁺ T cells

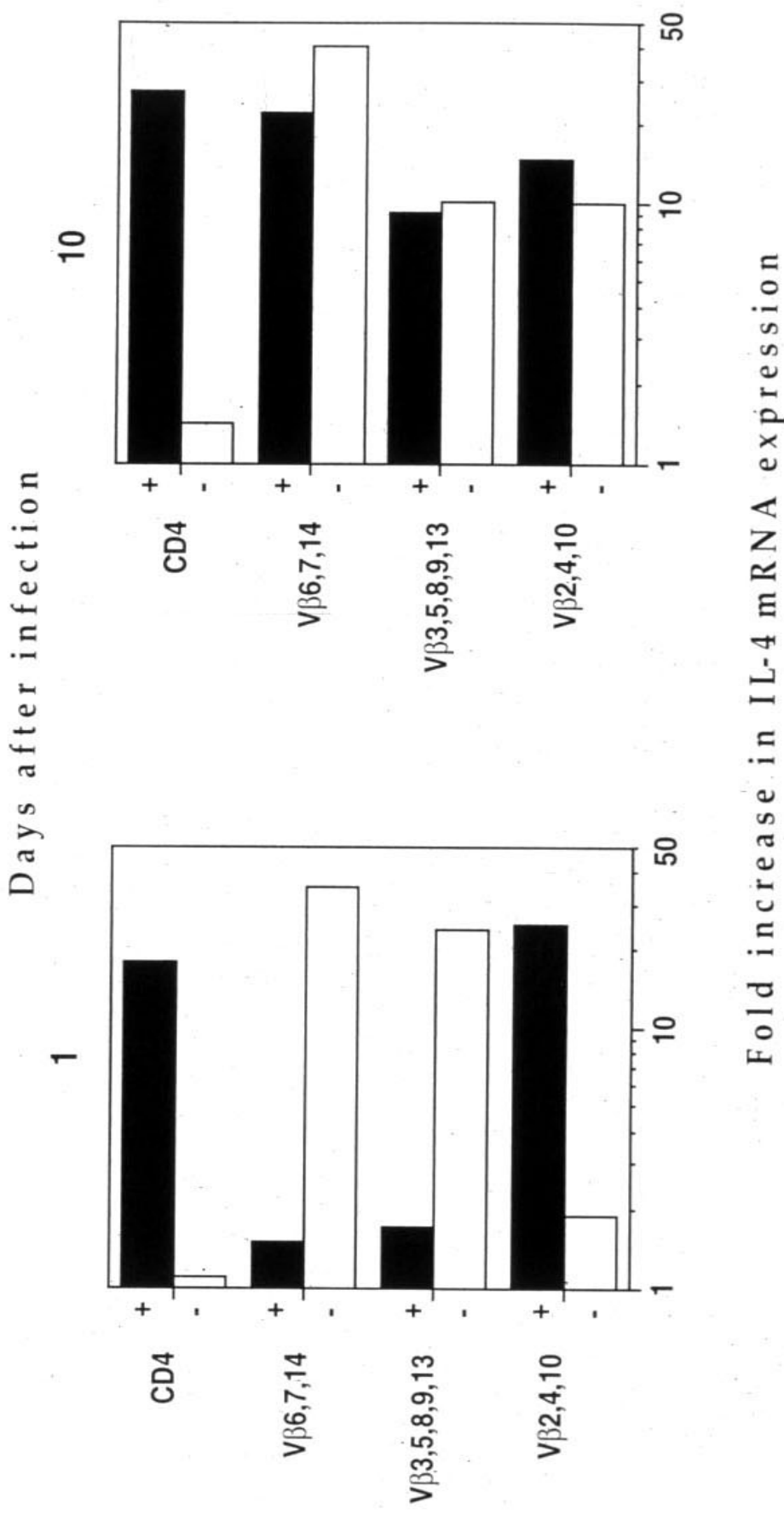
IL-4:

- Mast cells, basophils (stimulated to produce IL-4 by cross-linkage of Fc ϵ RI or Fc γ RII/III)

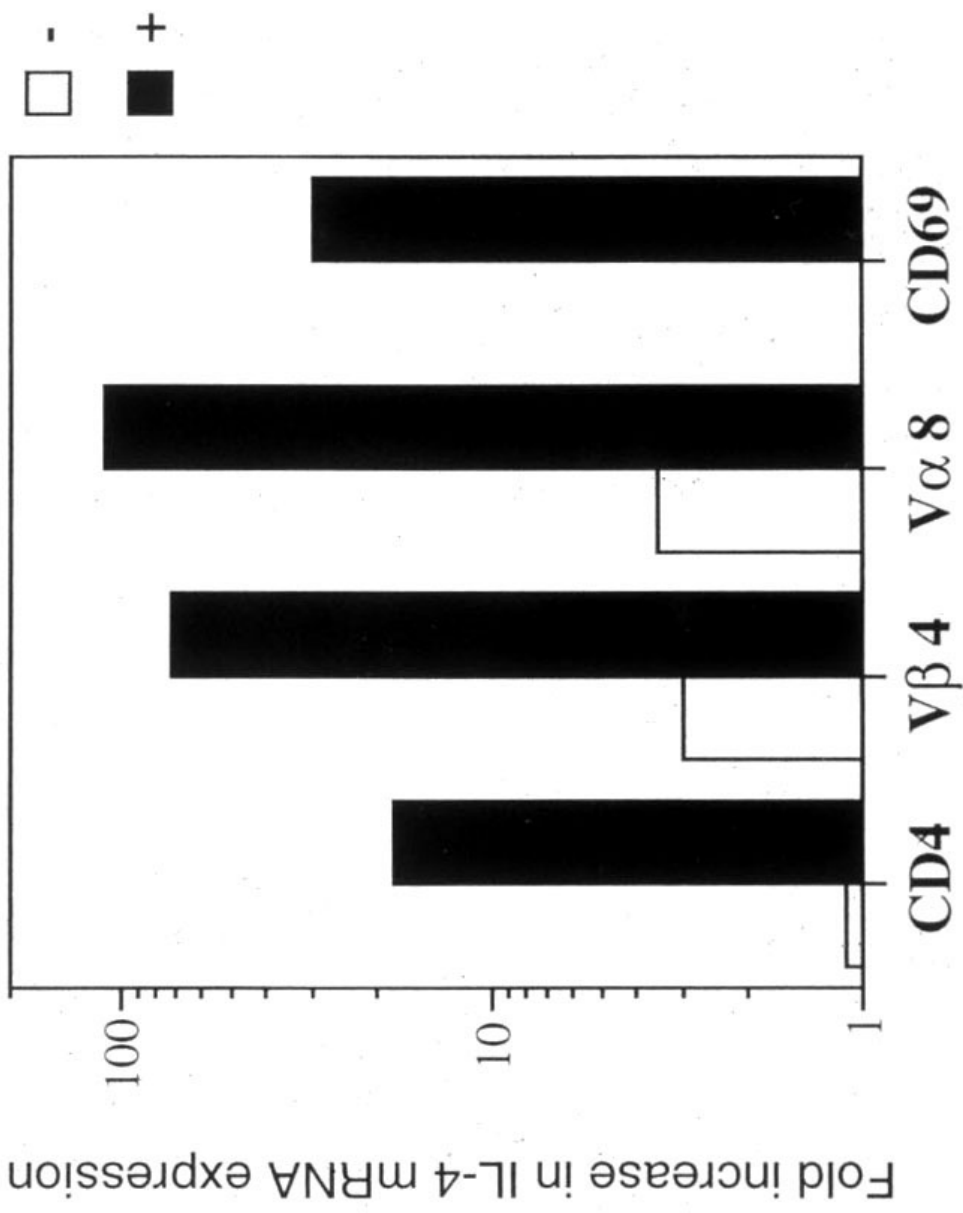
- Activated (memory) CD4⁺ T cells

- NK1.1 pos CD4⁺ cells

Characteristics of the T cells producing IL-4 in BALB/c mice infected with *L. major*



**Phenotype of CD4+ cells producing IL-4 in lymph nodes
16 hrs after infection with *L. major***

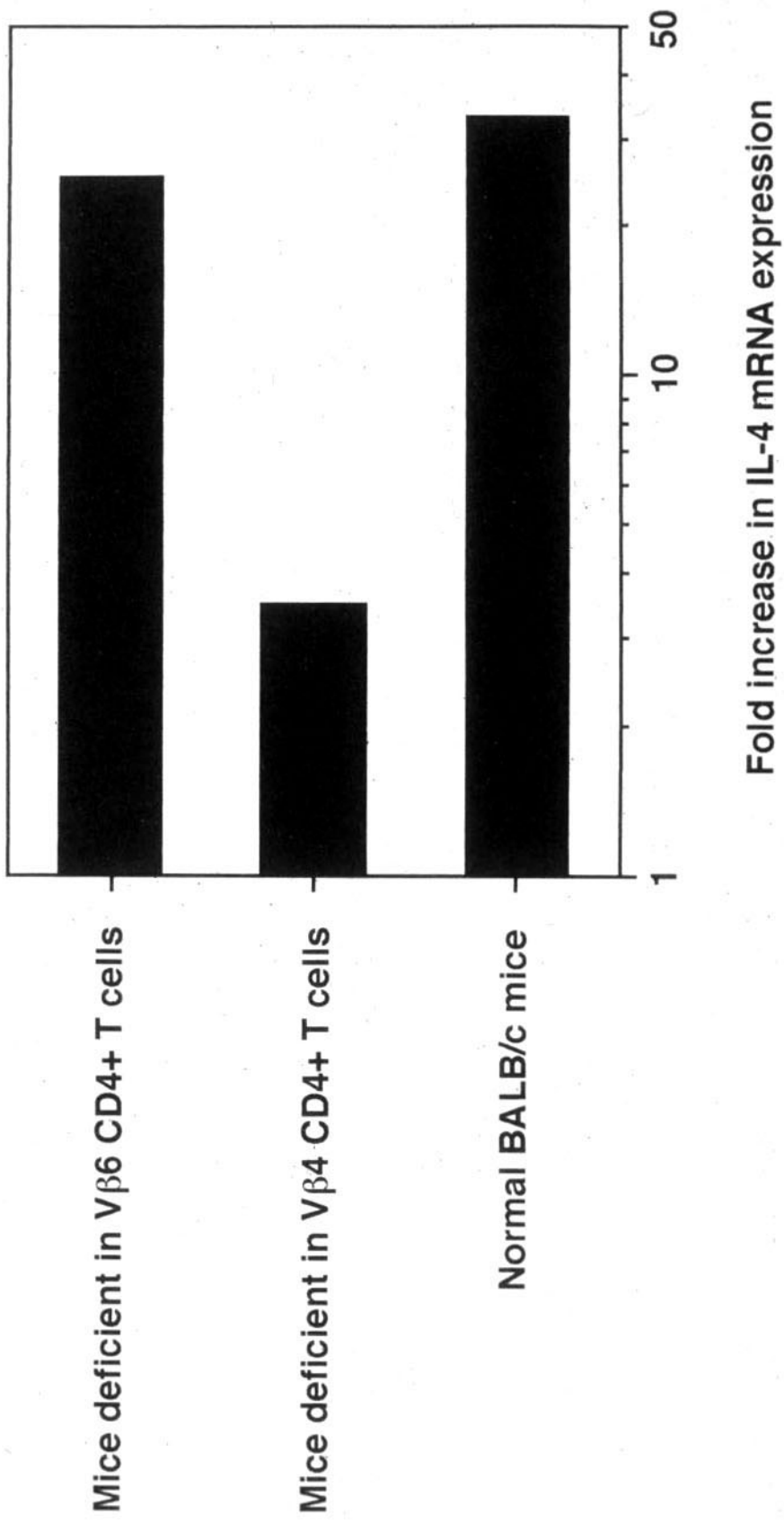


BALB/c mice deficient in T cells expressing the V β 4 TCR chain

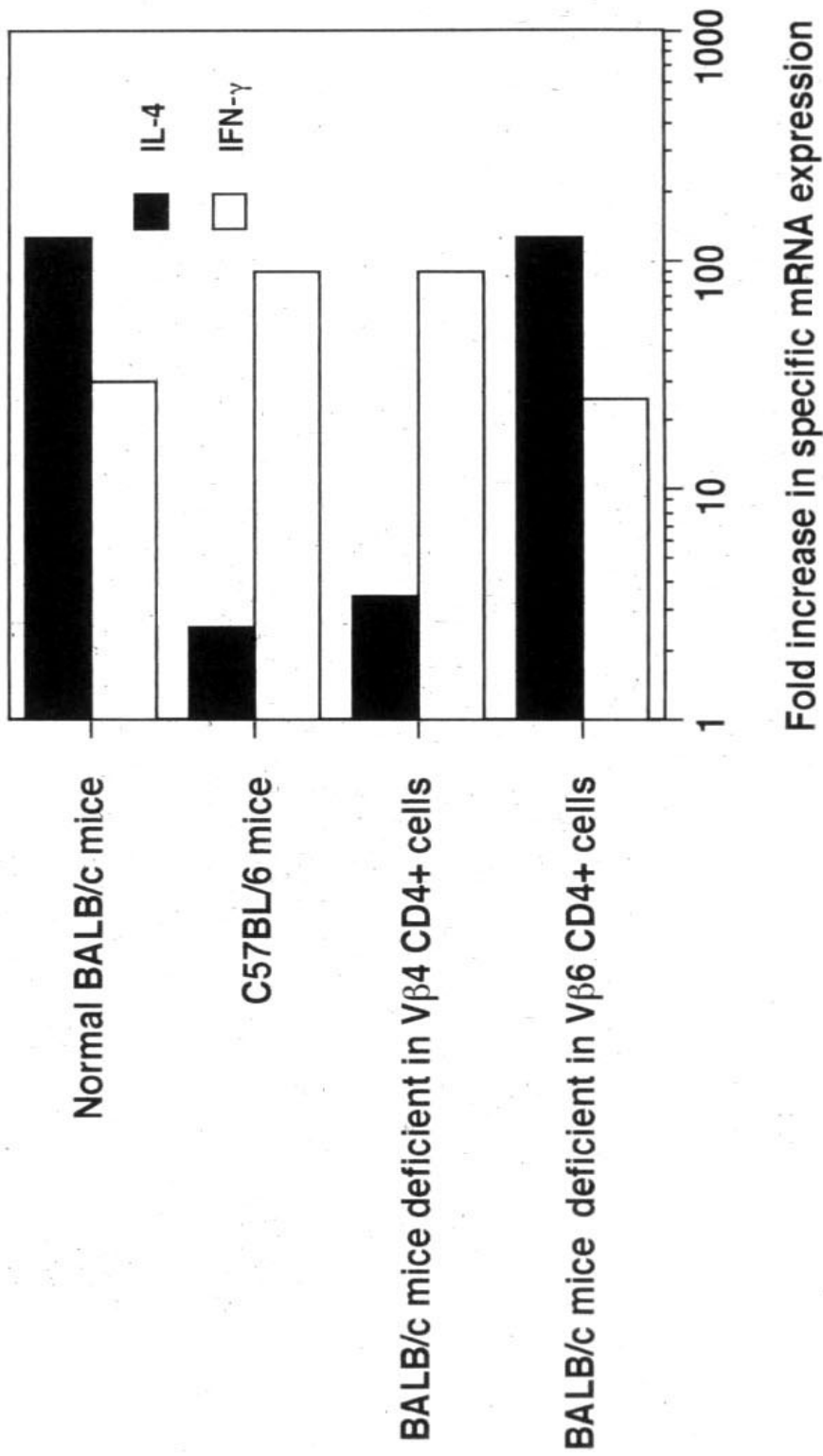
Mouse mammary tumor viruses (MMTV) encode a superantigen that ultimately leads to systemic deletion of CD4+ T cells expressing the V β TCR chain reacting with this superantigen

- MMTV-SIM encodes a superantigen leading to systemic deletion of **V β 4+** CD4+ T cells (Maillard, I. et al., Eur. J. Immunol. 1996, 26, 1000).
- MMTV-SW encodes a superantigen leading to systemic deletion of **V β 6+** CD4+ T cells (Held, W. et al., J. Exp. Med. 1992, 175, 1623).

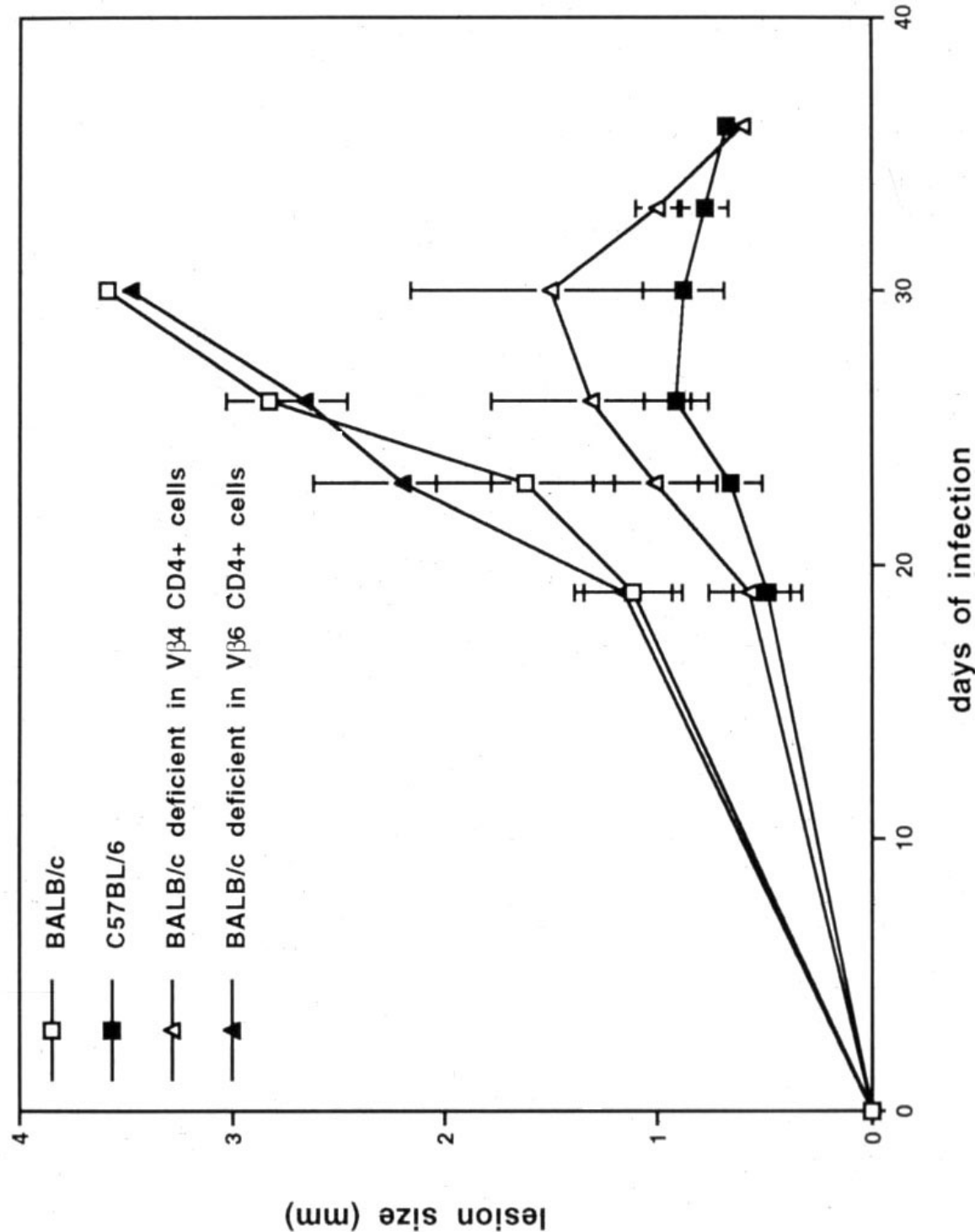
BALB/c mice deficient in V β 4+ CD4+ T cells do not exhibit early (16hrs) IL-4 mRNA expression following infection with *L. major*



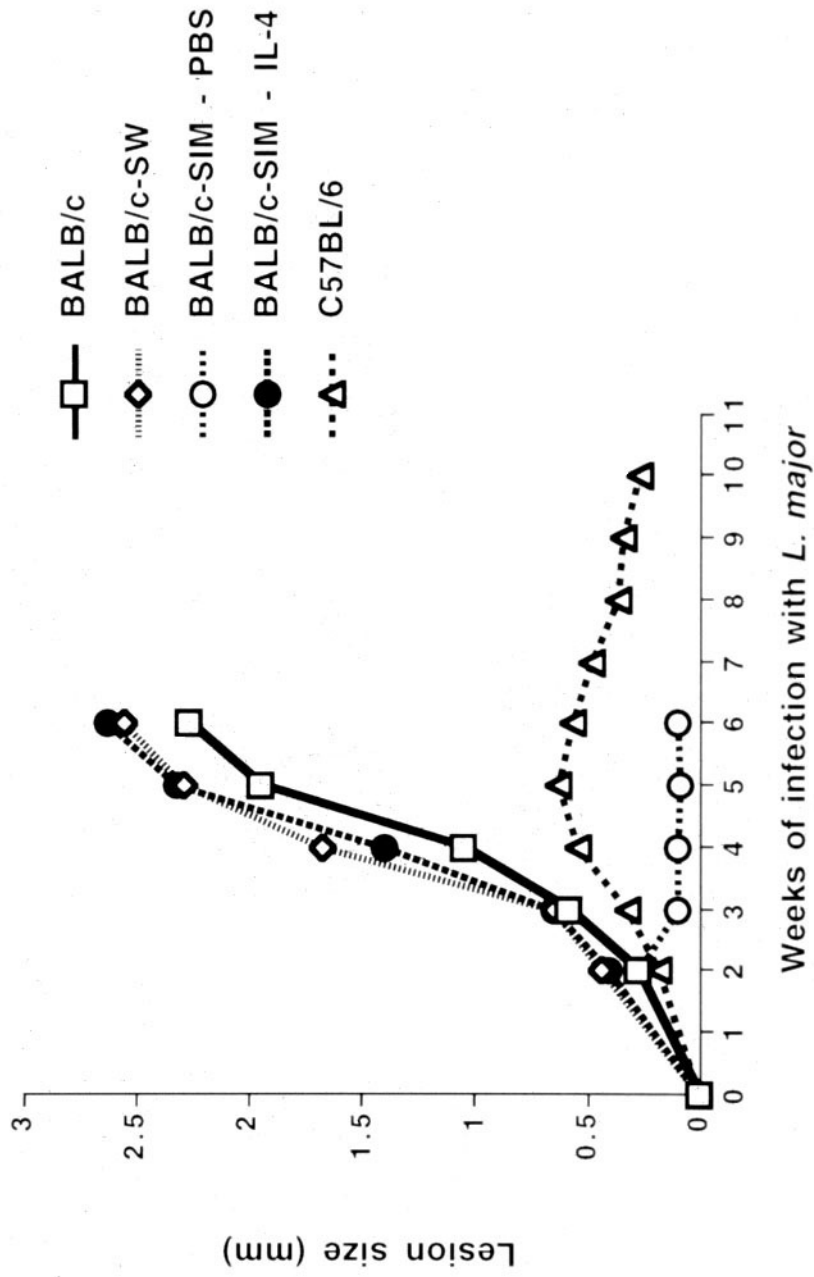
Th2 cell development does not occur in BALB/c mice deficient in V β 4+ CD4+ T cells following infection with *L. major*



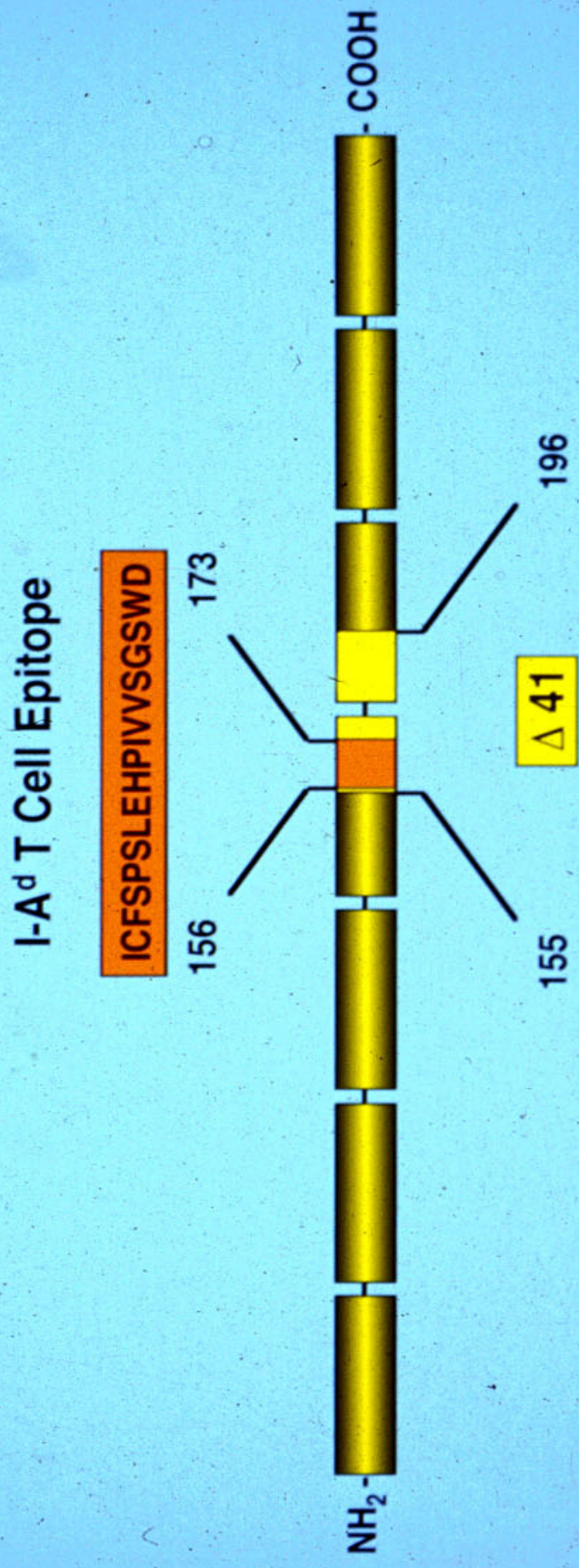
BALB/c mice deficient in V β 4+ CD4+ T cells are
resistant to infection with *L. major*



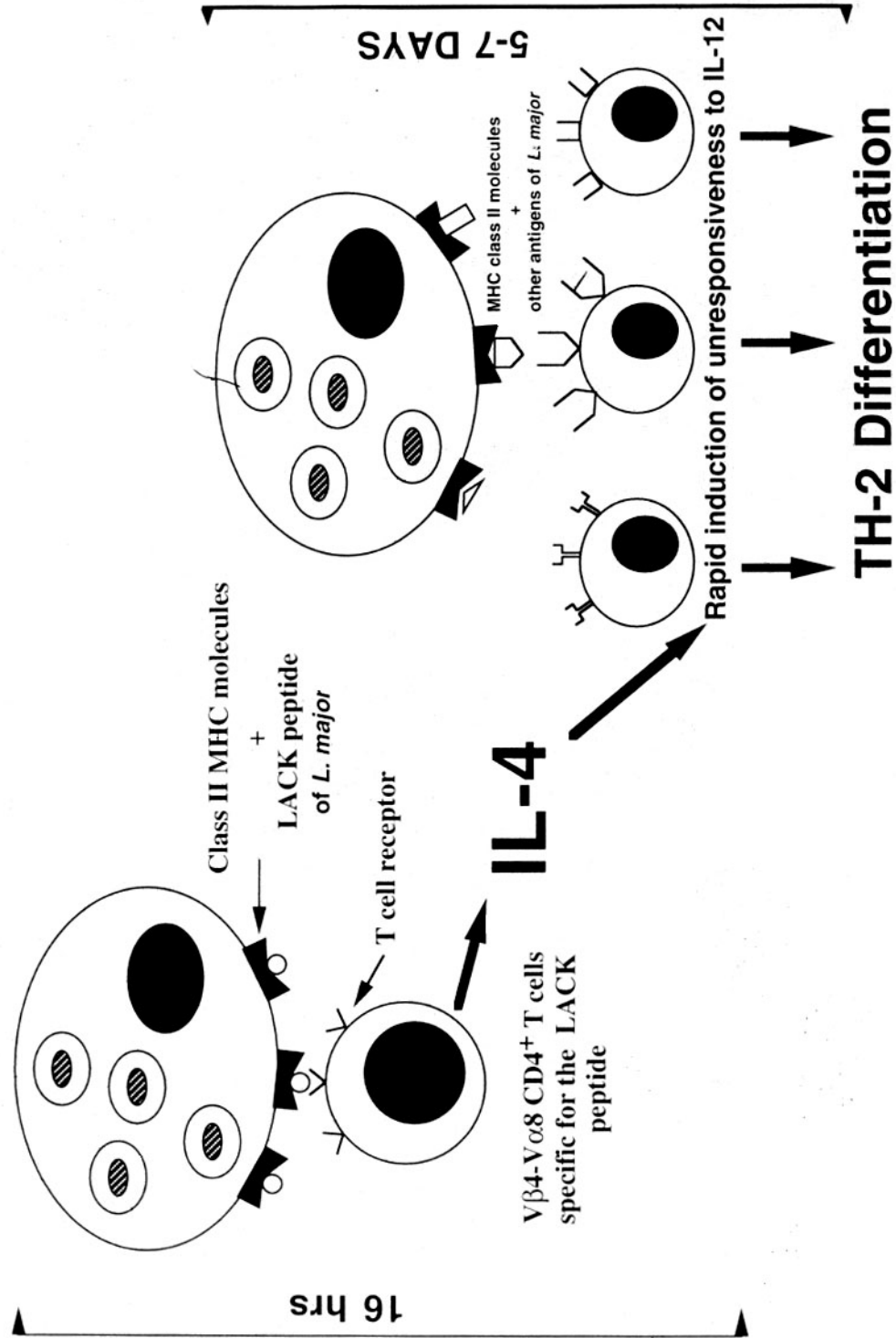
IL-4 during the early stage of infection is necessary and sufficient to instruct Th2 development and susceptibility to *Leishmania major* in BALB/c mice



Protein Structure of the Immunodominant Antigen LACK from *L. major*



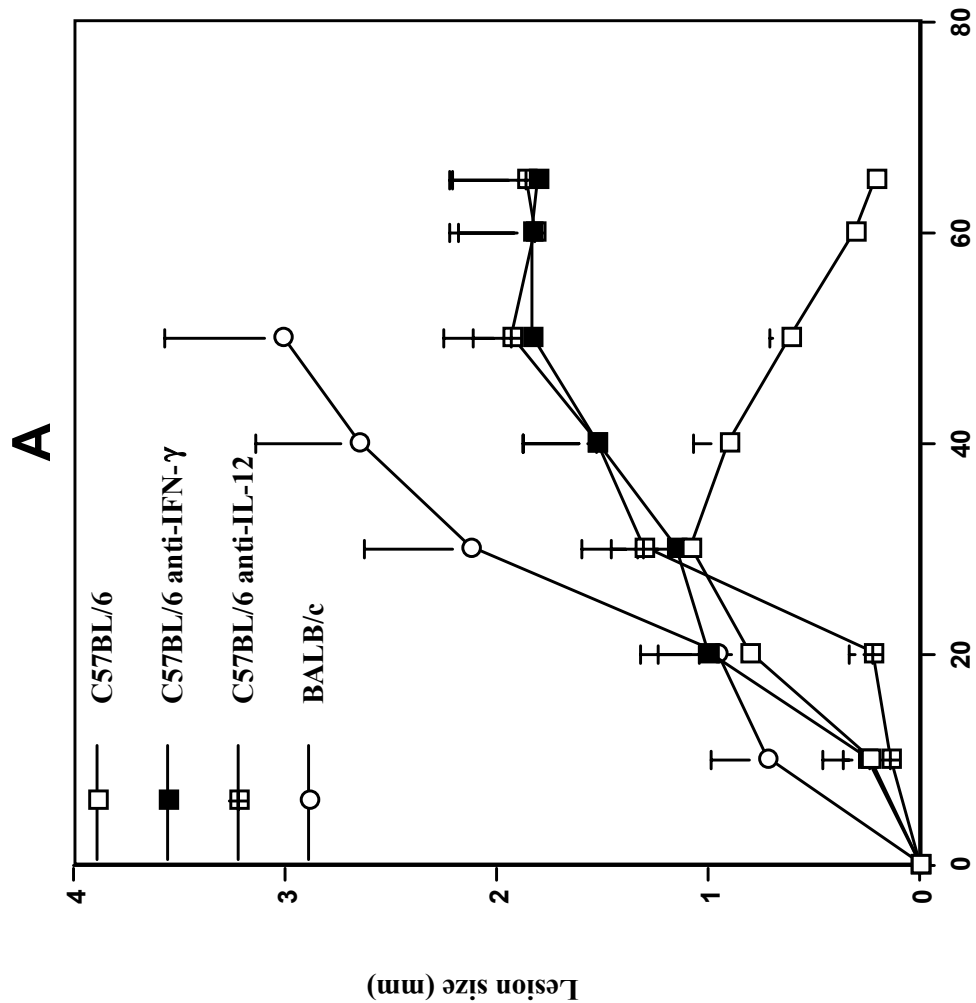
IL-4 rapidly produced by V β 4V α 8 CD4 $^+$ T cells instructs Th2 cell development and the susceptibility to L. major in BALB/c mice



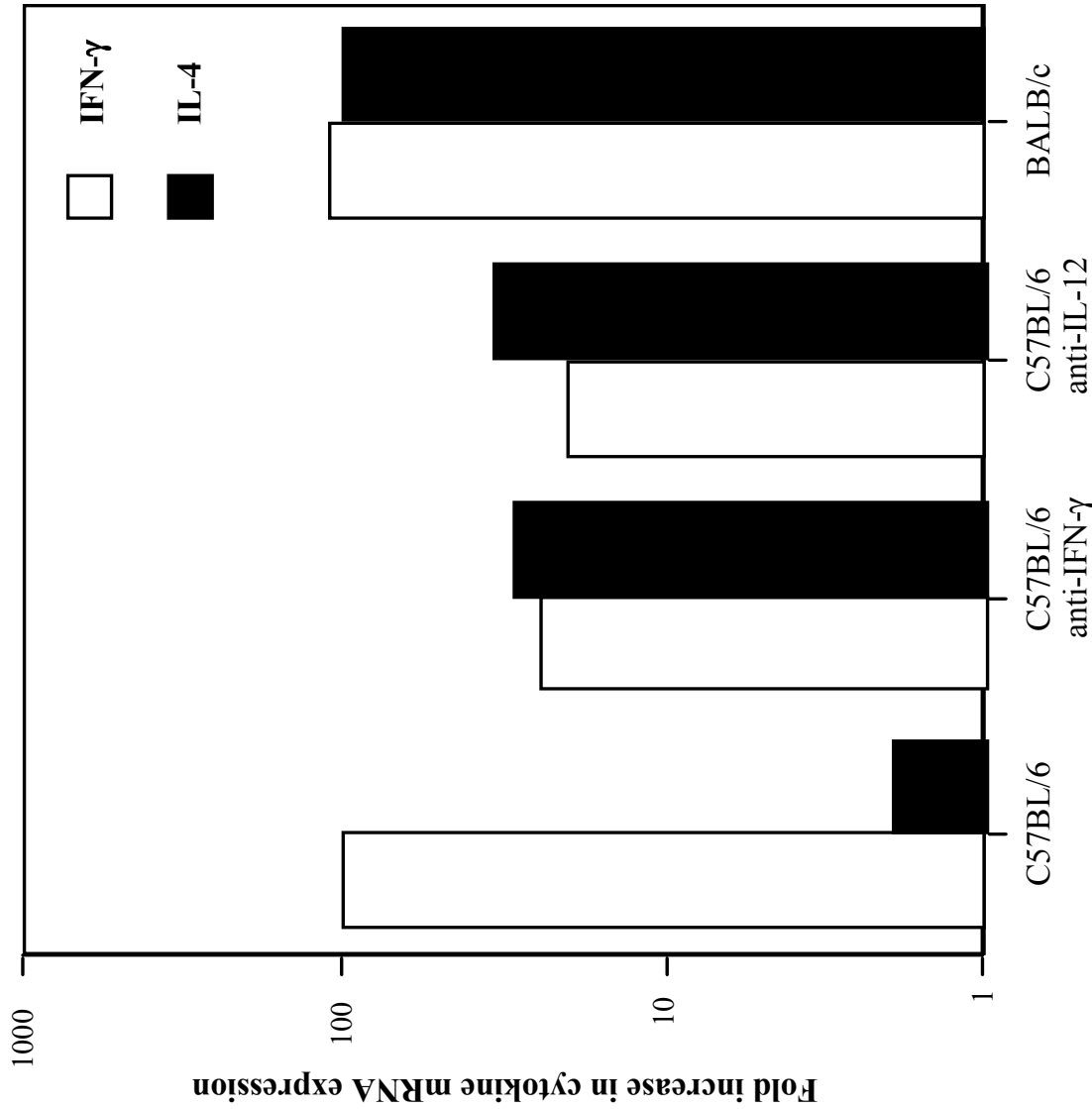
What might underlie susceptibility to infection with L. major in BALB strain mice?

- A greater LACK-specific T cell precursor frequency could account for the capacity of the initial IL-4 production in response to LACK to exceed the threshold required for Th2 lineage commitment.
- Susceptibility of BALB mice might be linked to an inability to down-regulate early IL-4 production by LACK-reactive cells.

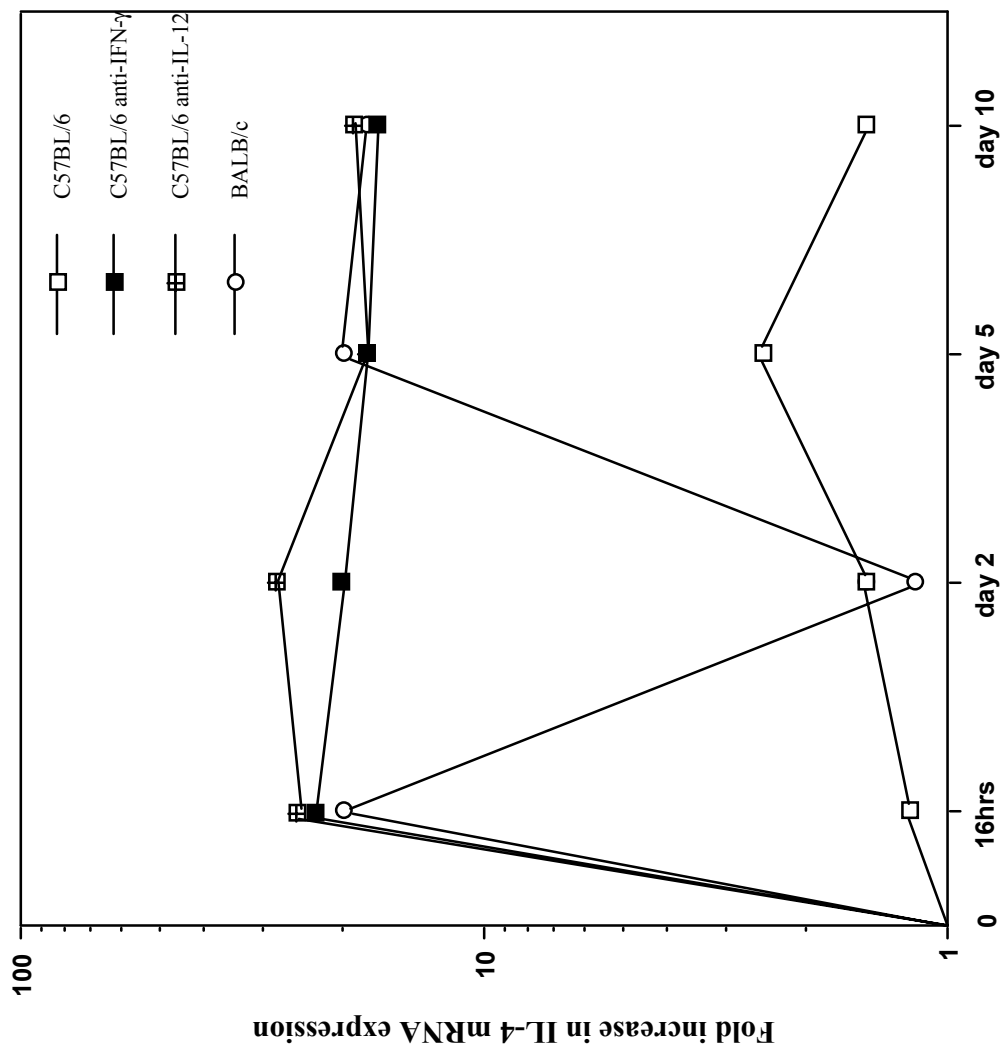
Lesions' development in C57BL/6 mice treated with anti-IL-12 or -IFN- γ at the onset of infection with *L. major*



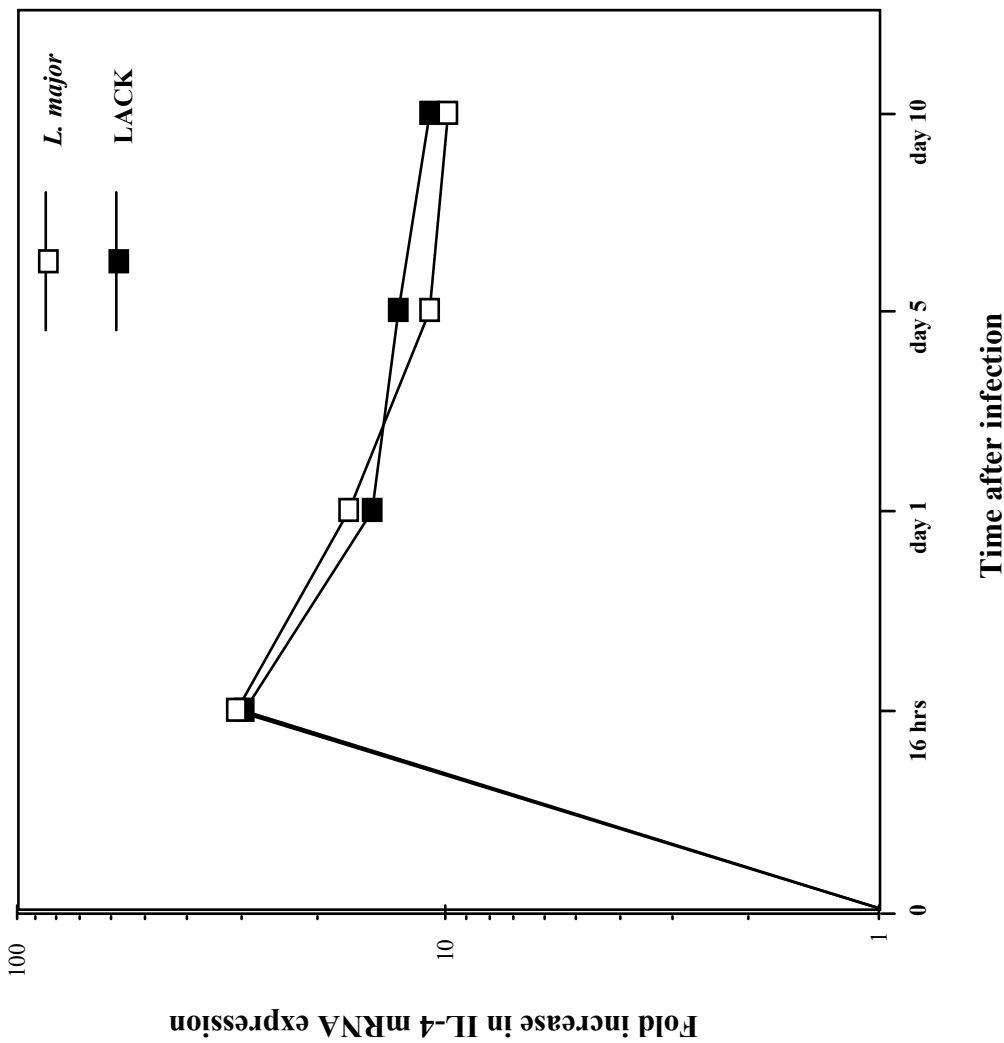
Cytokines transcripts in draining lymph nodes 45 days after infection with L. major in C57BL/6 mice treated with anti-IL-12 or IFN- γ



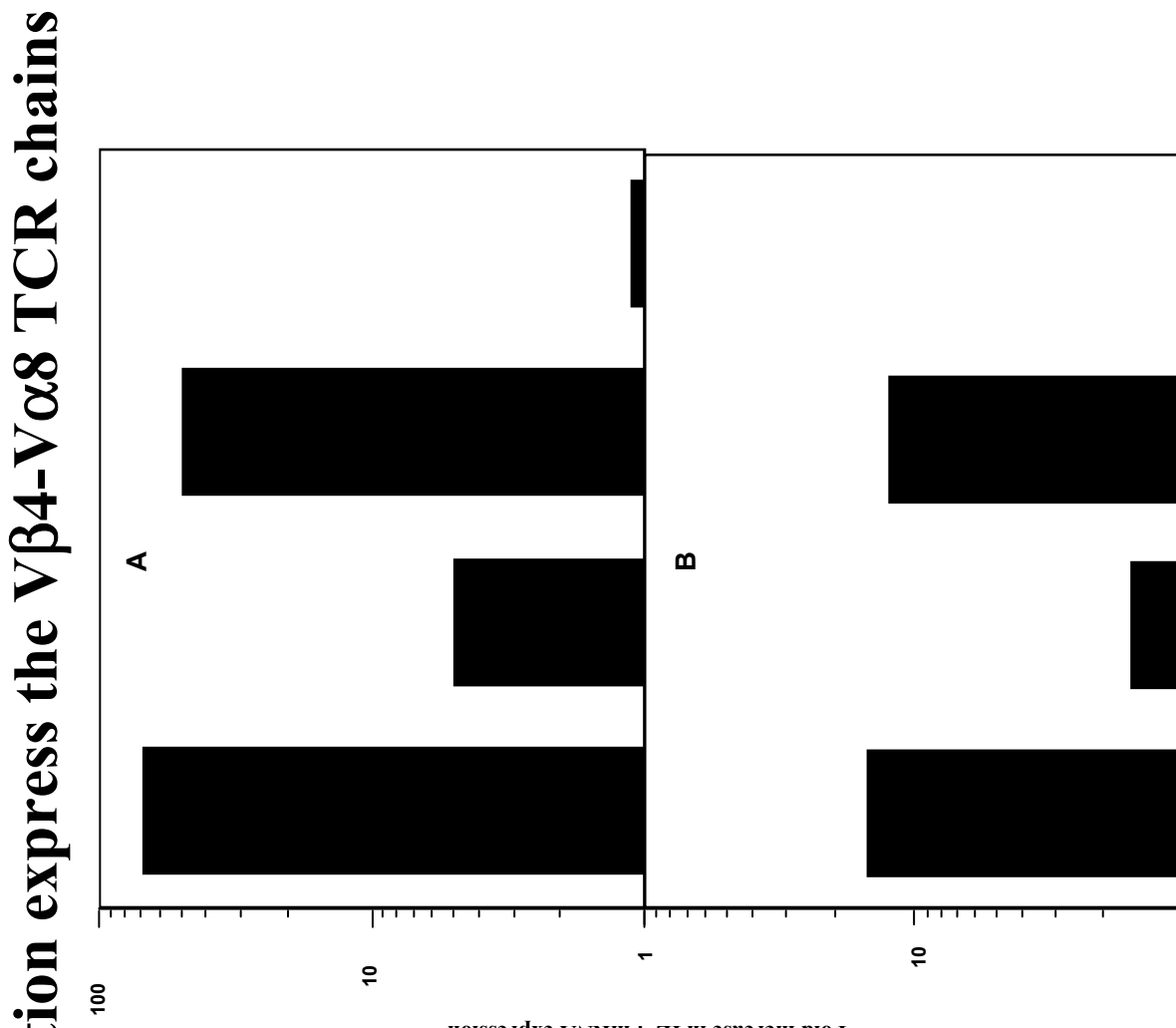
Kinetics of IL-4 mRNA expression in lymph nodes of C57BL/6 mice treated with anti-IFN- γ or-IL-12 at the onset of infection with Leishmania major



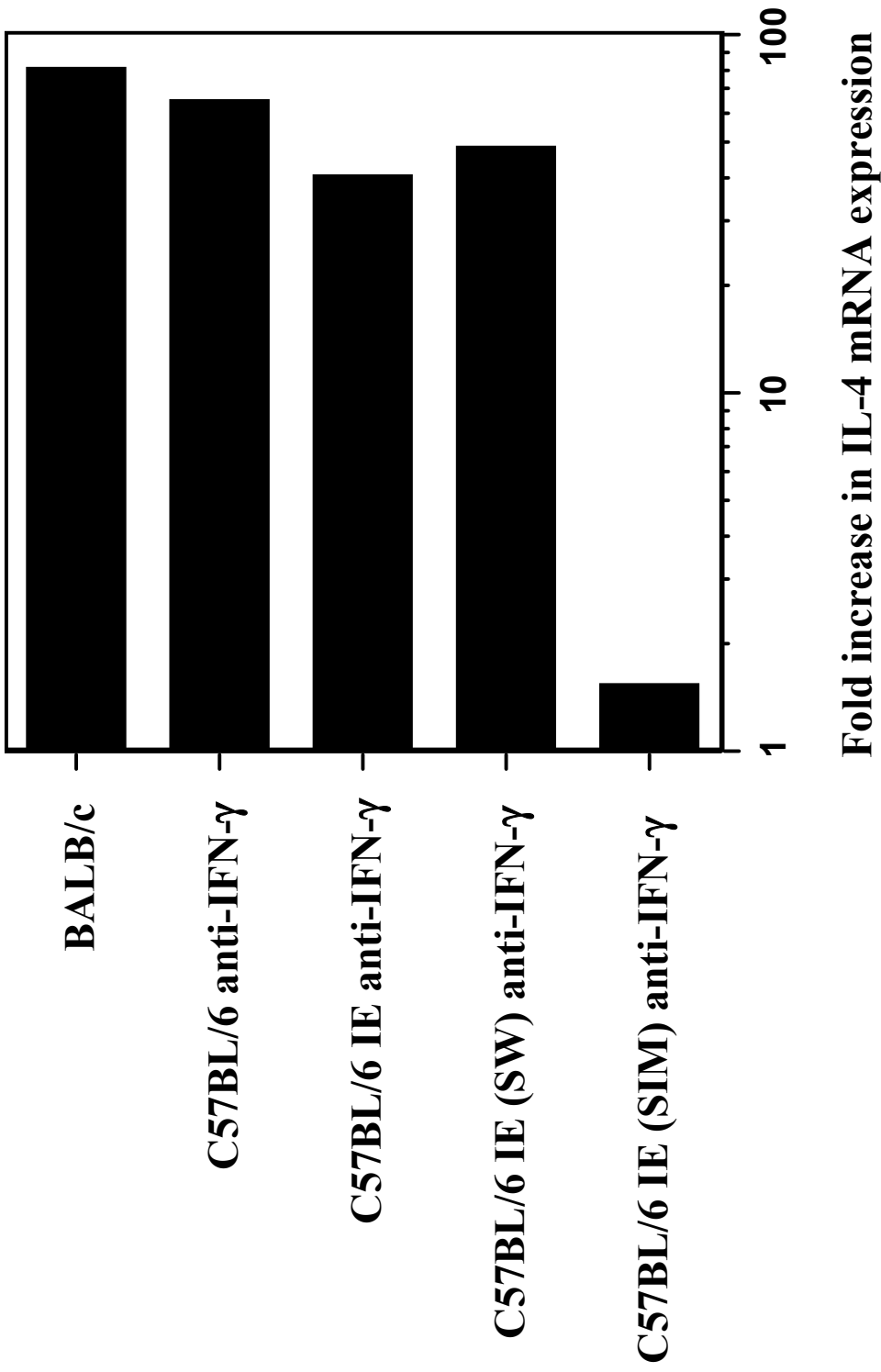
LACK induces a rapid IL-4 response in draining lymph node cells from C57BL/6 mice treated with anti-IL-12 or -IFN- γ



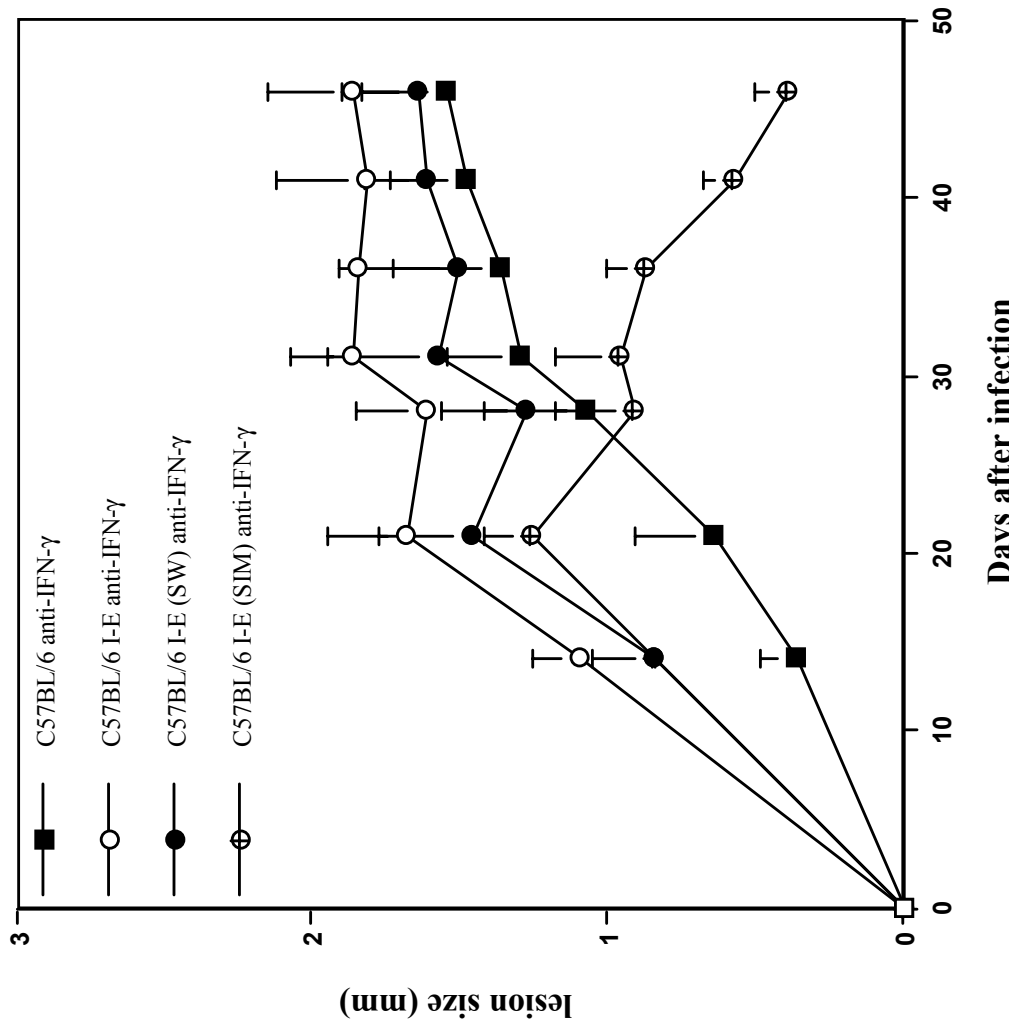
The CD4 cells producing IL-4 in response to L. major in C57BL/6 mice treated with anti-IL-12 at the onset of infection express the V β 4-V α 8 TCR chains



**Early IL-4 mRNA expression in response to L. major does
not occur in V β 4-deficient C57BL/6 mice treated with
anti-IFN- γ at the initiation of infection**



Lesions' development in anti-IFN- γ treated I-E transgenic C57BL/6 mice deficient in V β 4 CD4 T cells



Treatment of V β 4-deficient I-E transgenic C57BL/6 mice with anti-IFN- γ before infection with L. major does not interfere with Th1 cell development

