

# HEPATITIS C VIRUS AND HUMAN IMMUNODEFICIENCY VIRUS: PATHOGENESIS, IMMUNITY AND TREATMENT

Unit 3

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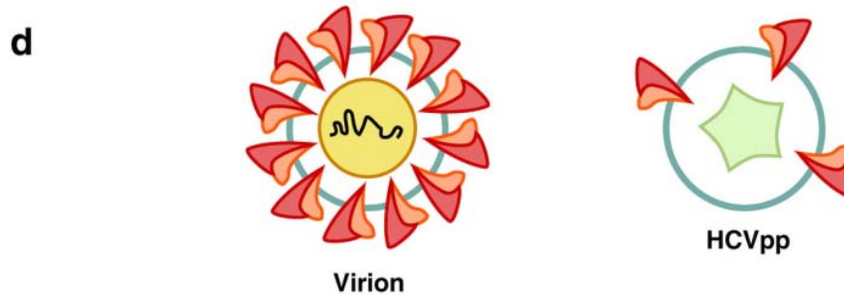
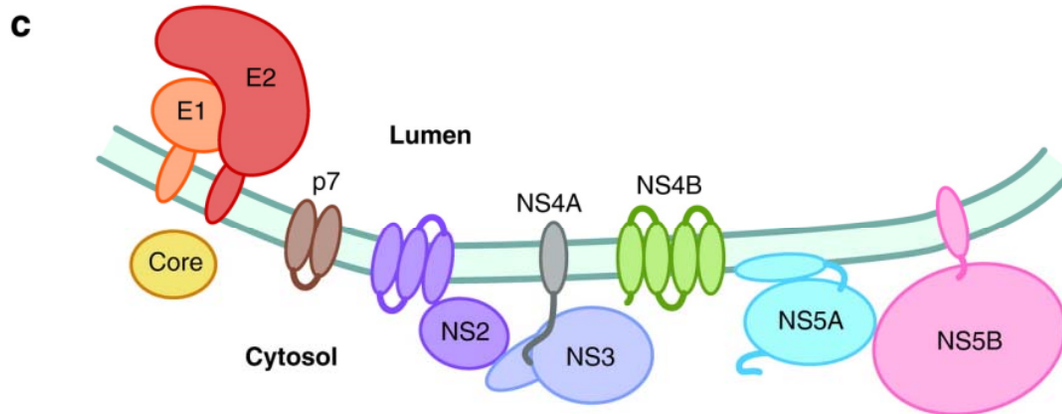
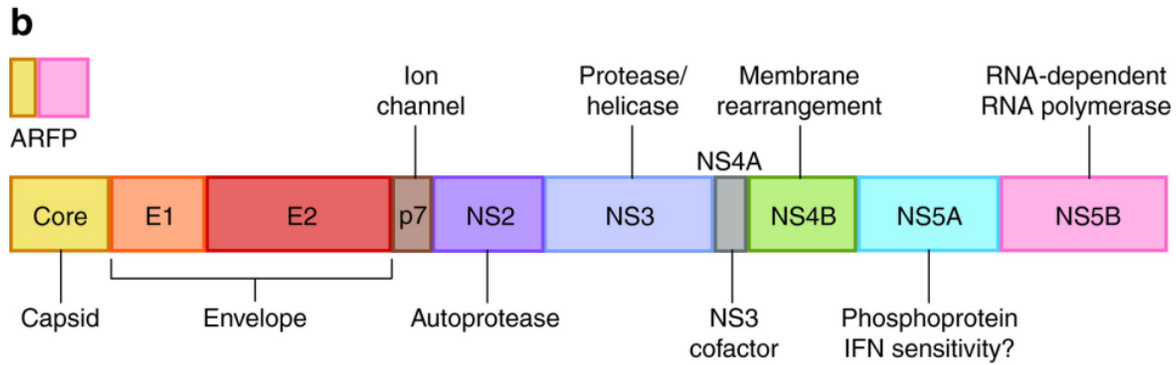
St. Jude Children's Research Hospital

## HEPATITIS C VIRUS

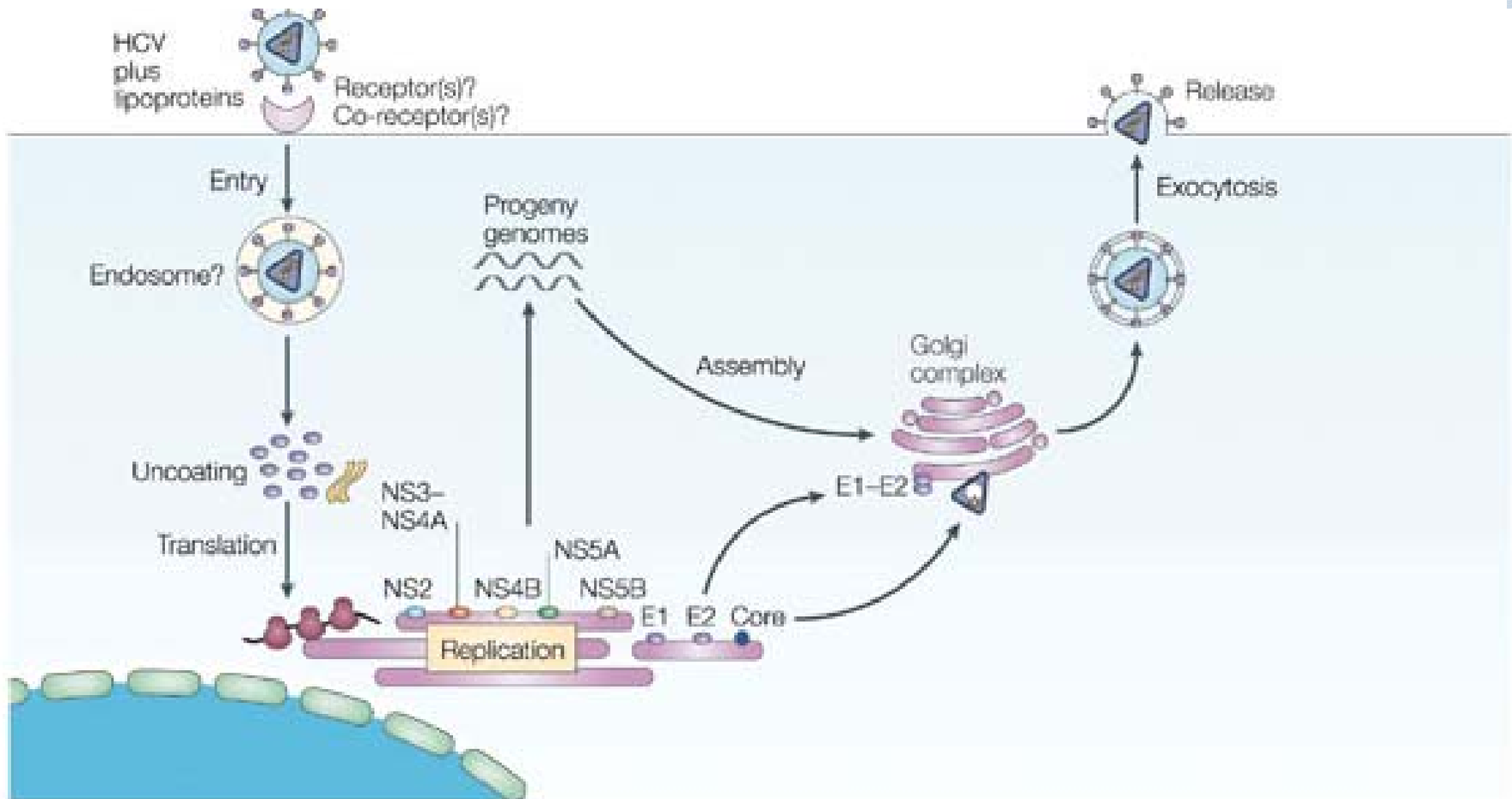
- Enveloped, positive strand RNA virus, *Flaviviridae*
- Isolated in 1989, treatments first emerged in early 1990s
- ~120 million-200 million infections worldwide, number one indication for liver transplant in the U.S.
- $10^{12}$  viral particles produced/day,  $\frac{1}{2}$  life 3 hours in circulation
- Six major genotypes, 3 dominate in the U.S. (1, 2, 3)
  - 30-50% genetic variation among genotypes
  - 1-5% variation among viruses within a single patient
- Replicates via negative-stranded RNA in membranous web in cytoplasm



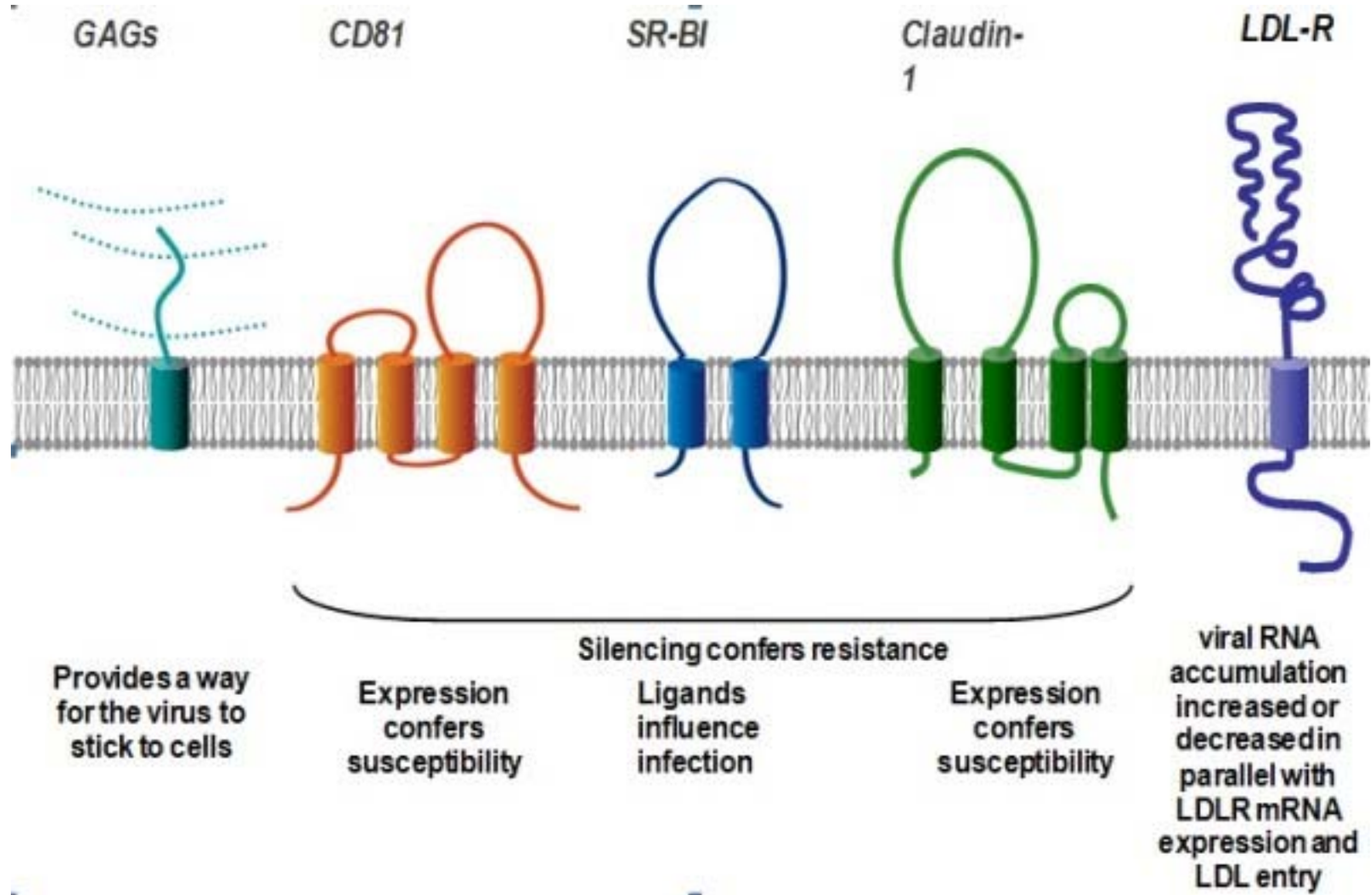
# HCV STRUCTURE



# HCV LIFE CYCLE



# RECEPTORS FOR VIRAL ENTRY

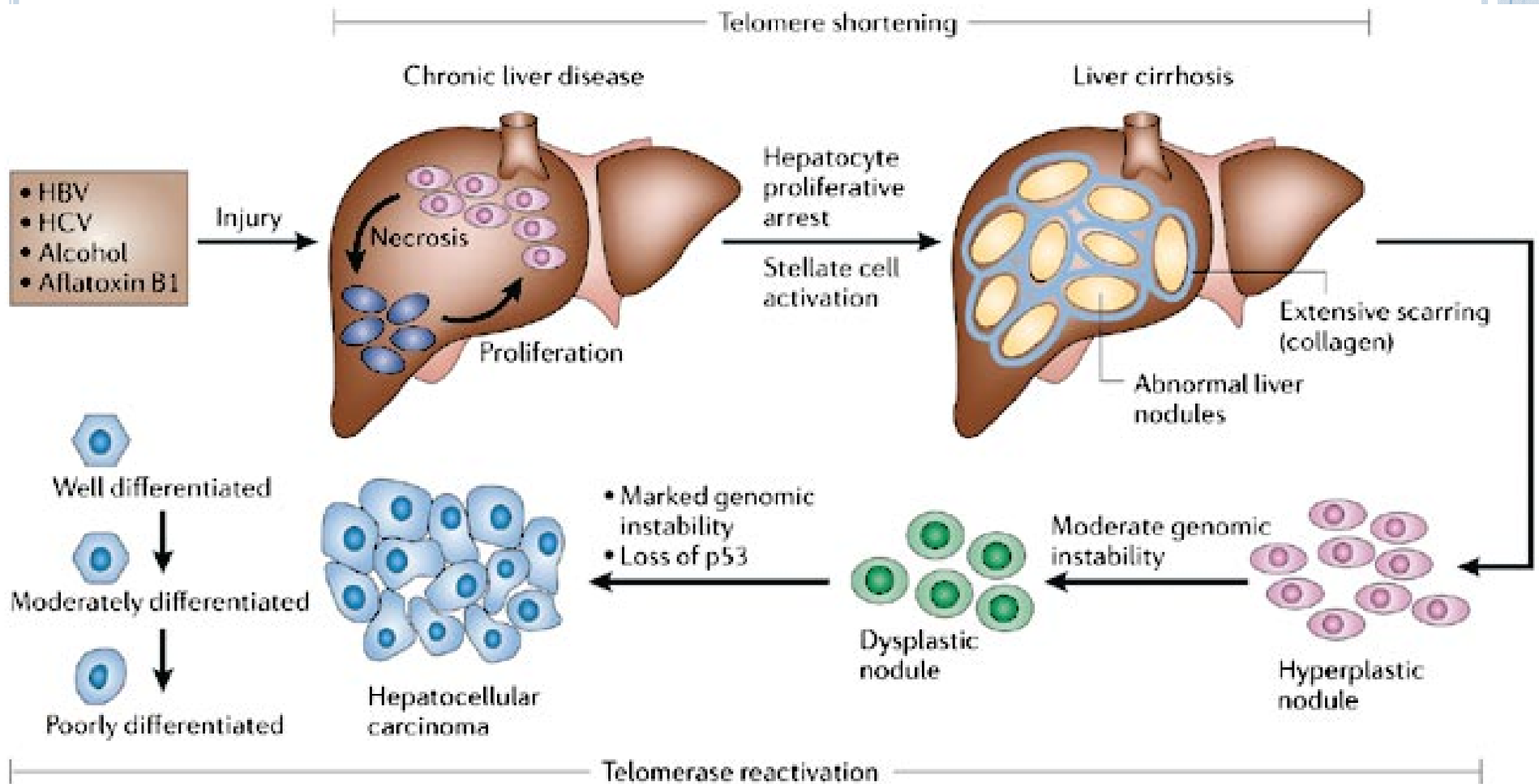


**HCV receptors for cell entry.**

Ashfaq *et al.* *Virology Journal* 2011 8:161 doi:10.1186/1743-422X-8-161

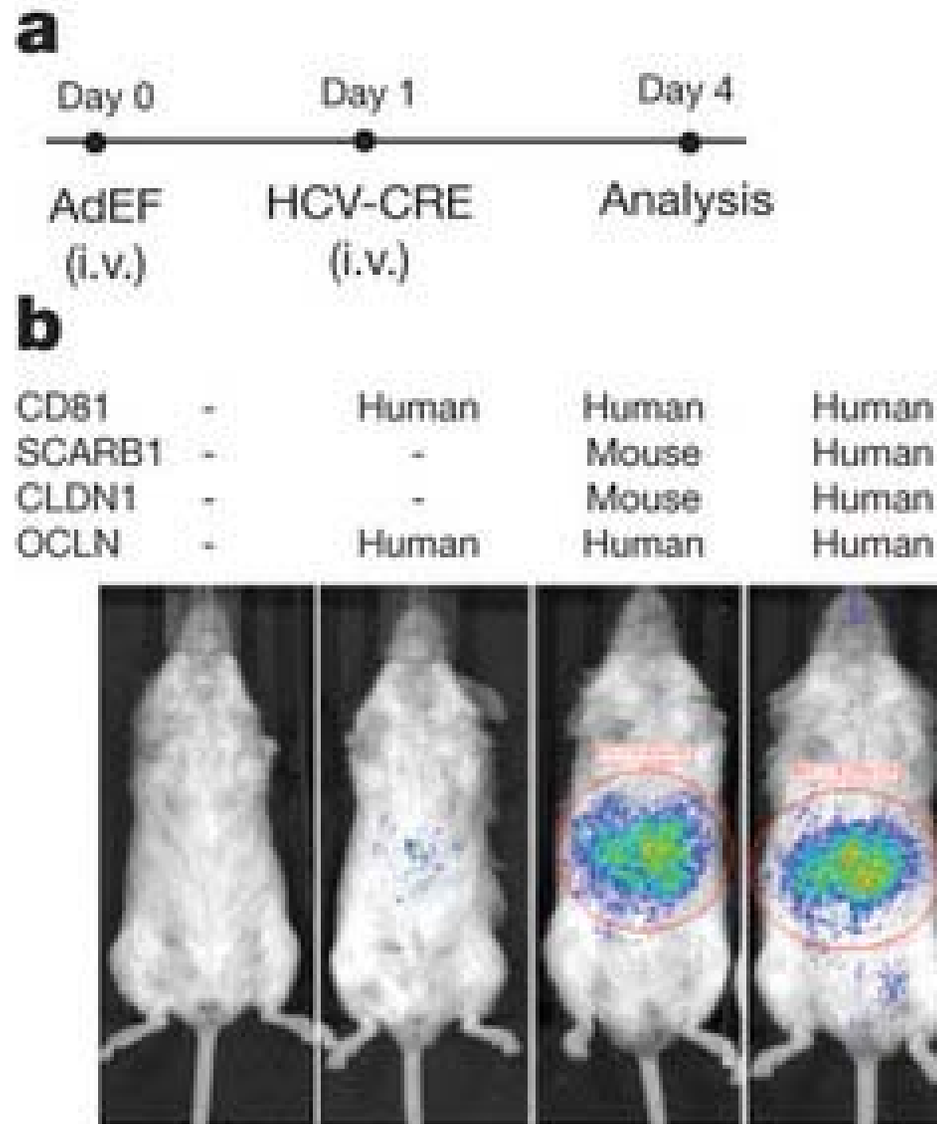
# HCV LIFE CYCLE 2

- HCV-associated disease results from viral persistence leading to long term inflammation and cell turnover



# MOUSE MODEL OF HCV REPLICATION

- Previous models relied human liver transplant into immunodeficient mice—limited usefulness
- Transgenic approach using four known entry factors—Occludin, CD81, SCARB and claudin 1



**c**

A genetically humanized mouse model for hepatitis C virus infection  
 Nature 474, 208–211 (09 June 2011)

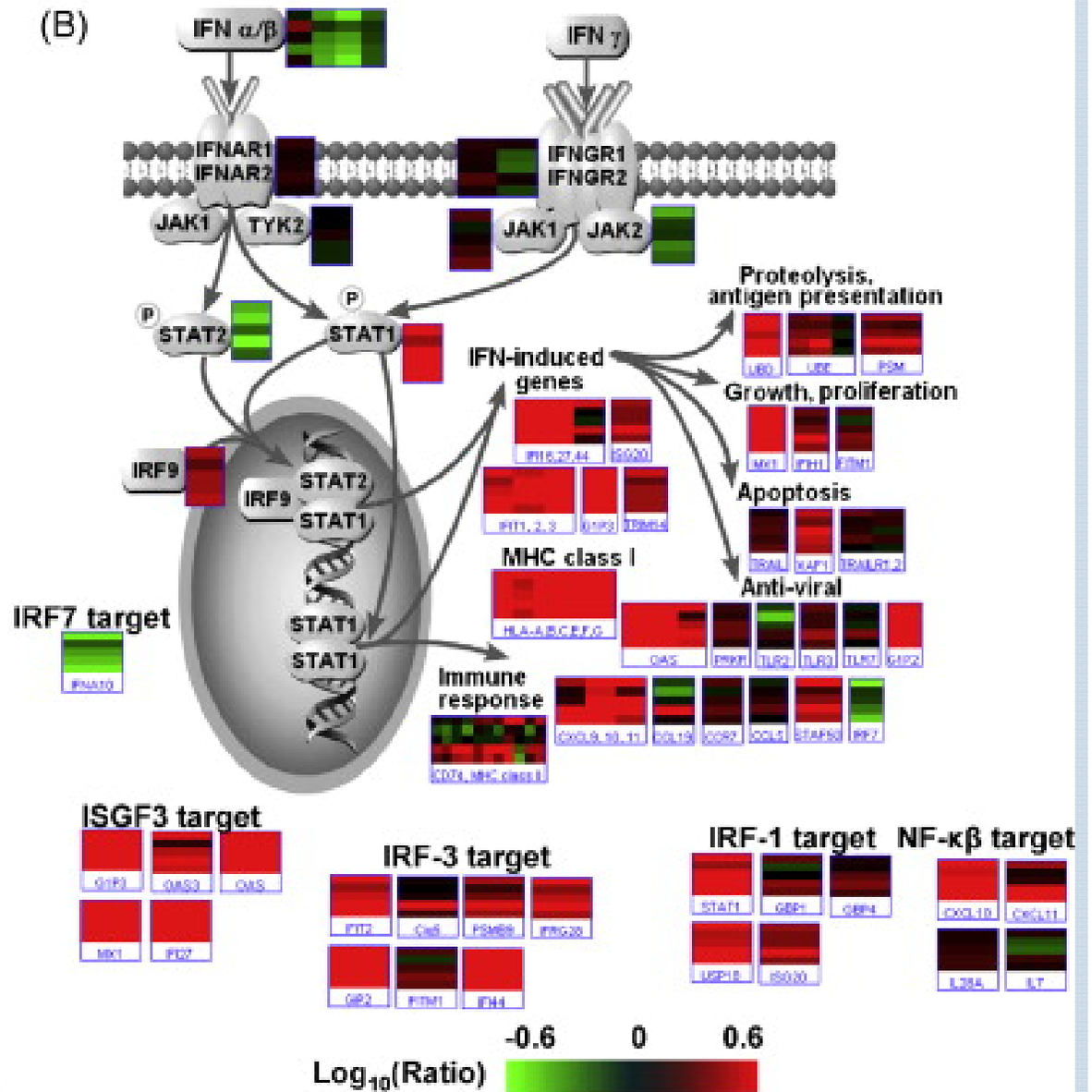
# WHAT ARMS OF THE IMMUNE RESPONSE ARE USEFUL AGAINST HCV?

- Innate immunity
  - Antiviral effectors such as IFN that act on host cells, regulating key components of cell biology to limit viral growth and spread
- Antibody-mediated clearance
  - In principle, antibodies should be able to remove virus as it spreads from cell to cell
  - In practice, the correlation of antibody with HCV clearance and outcome is controversial or lacking
  - Patients with high levels of *neutralizing* antibodies nevertheless maintain chronic infection, indicating that neutralizing antibodies are not *sterilizing*
- Cell-mediated clearance
  - Infected cells can be killed before releasing progeny virions
  - Thought to be the primary means of long term control in HCV infection



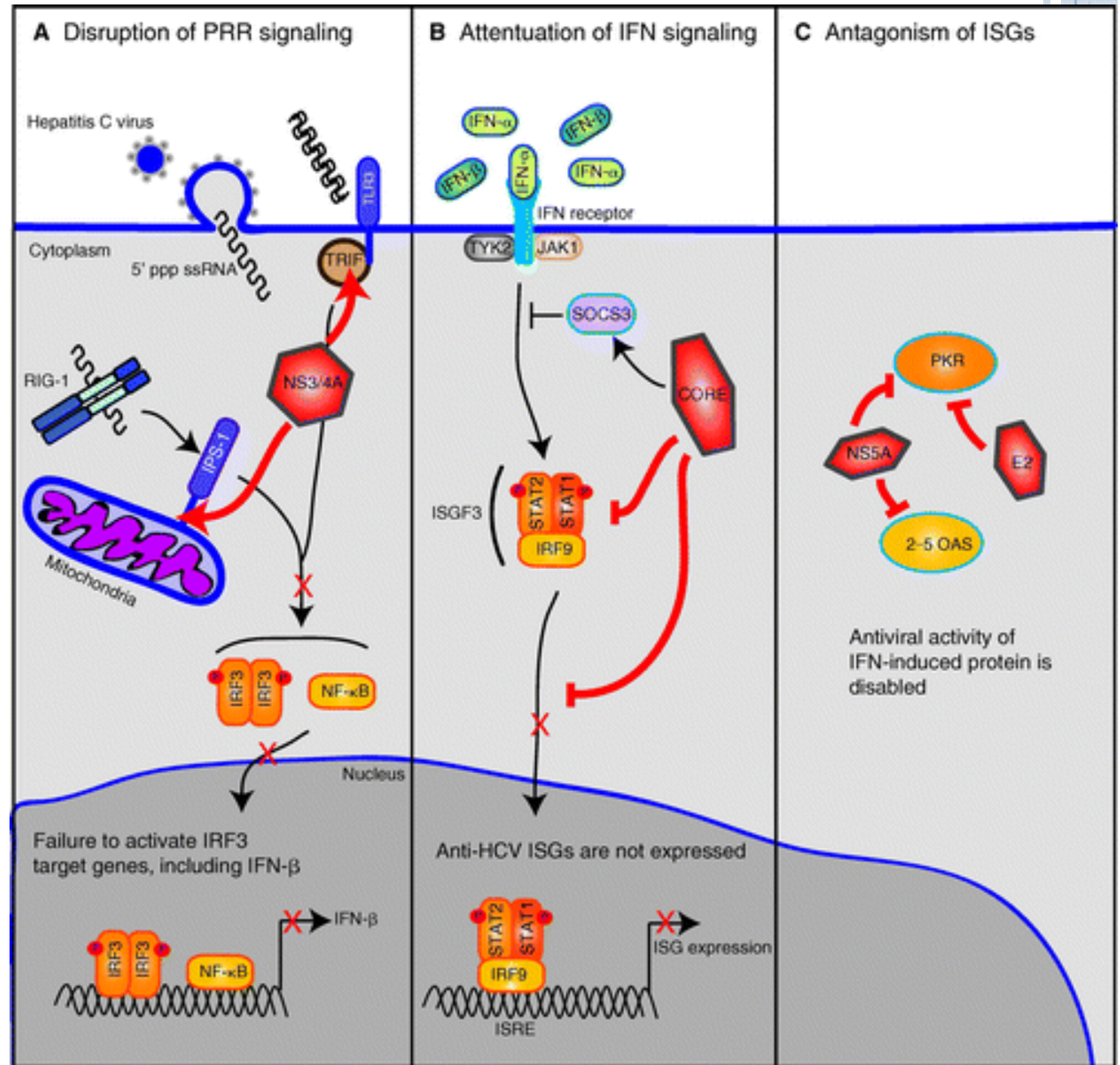
# INDUCTION OF INNATE IMMUNITY IN PATIENTS

- IFN-induced genes interfere with viral replication directly:
  - Reducing protein synthesis by inhibiting initiation factors (PKR, ISG56)
  - Targeting of viral RNA (OAS, RNaseL)
- Innate responses can enhance or initiate adaptive responses
  - MHC I expression
  - Chemokine secretion and recruitment of responder cells



# INNATE RECOGNITION OF HCV

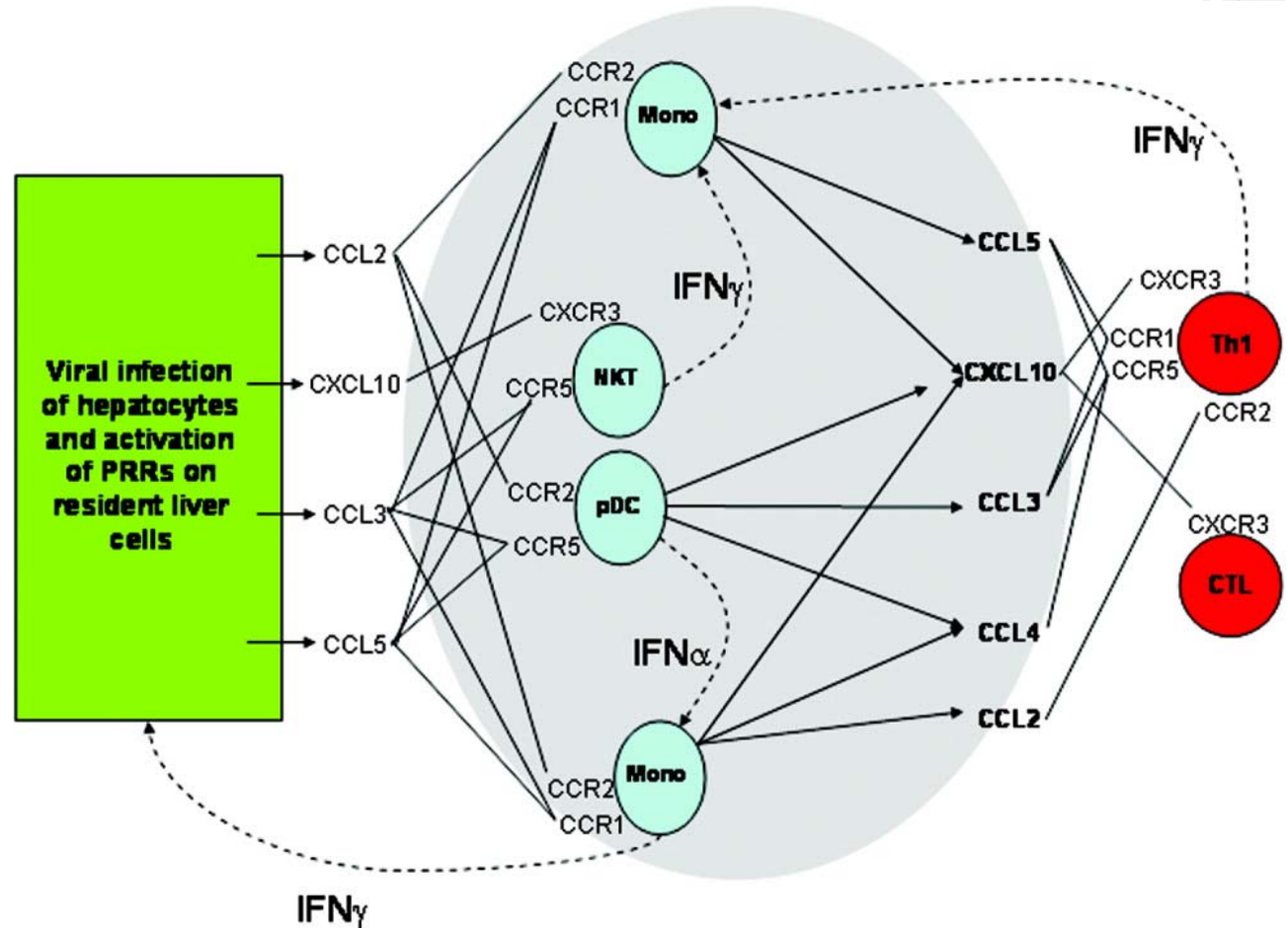
- The generation of dsRNA structures in HCV replication leads to recognition by multiple innate pathways
- HCV subverts these pathways by sequestering or cleaving key components of innate recognition
- The effects are both qualitative and quantitative on the ensuing innate response



Stacy M. Horner, Michael Gale. Journal of Interferon & Cytokine Research. September 2009, 29(9): 489-498

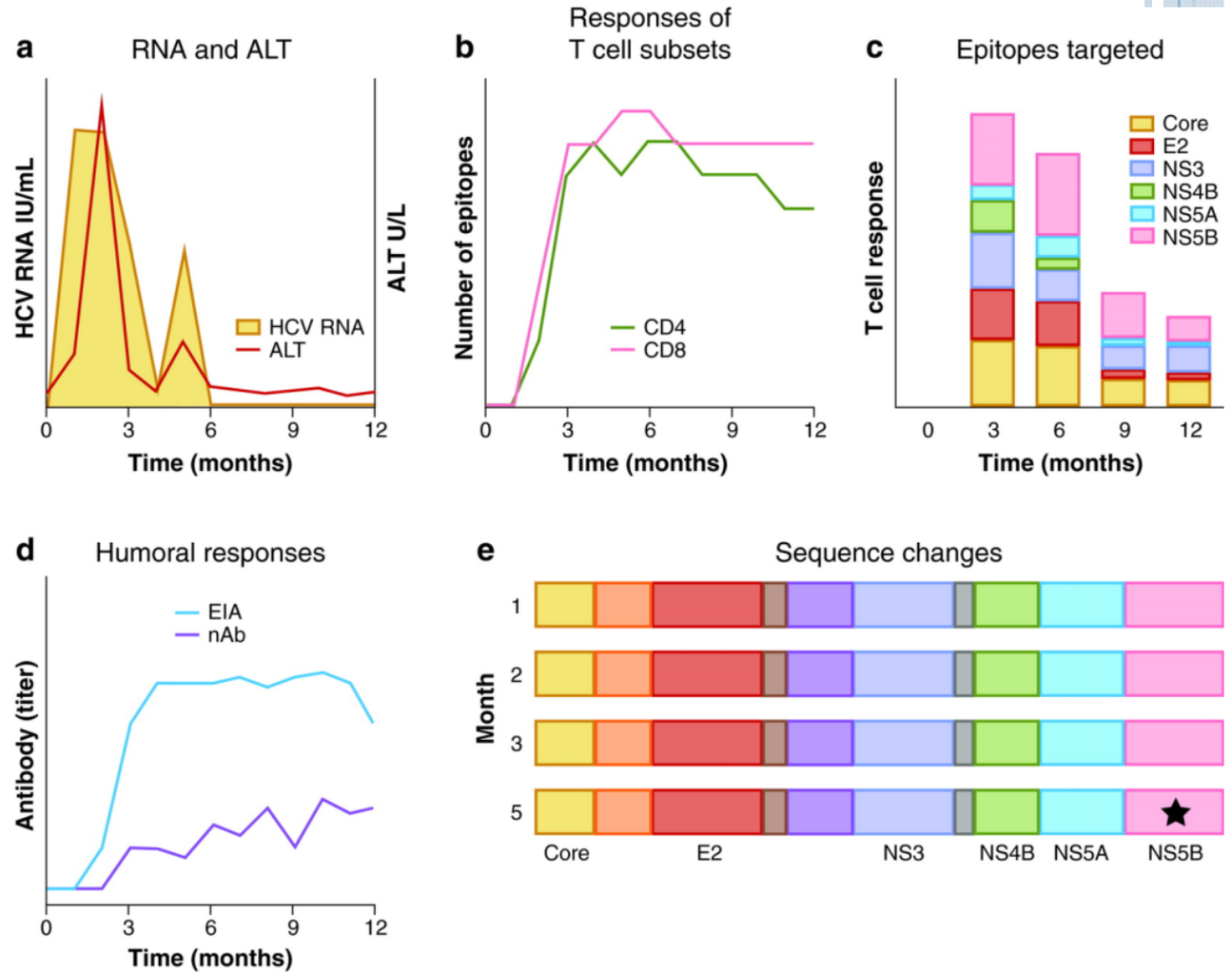
# INNATE ACTIVATION OF ADAPTIVE RESPONSES

- The innate response results in the recruitment and “biasing” of key innate and adaptive cell types, including NK cells, NKT cells, antigen-presenting cells (monocytes/macrophages) and ultimately CD4 T cells that will orchestrate the adaptive response



# SUCCESSFUL HCV CONTROL (SUSTAINED VIROLOGICAL RESPONSE) IS MEDIATED BY ROBUST ADAPTIVE IMMUNITY

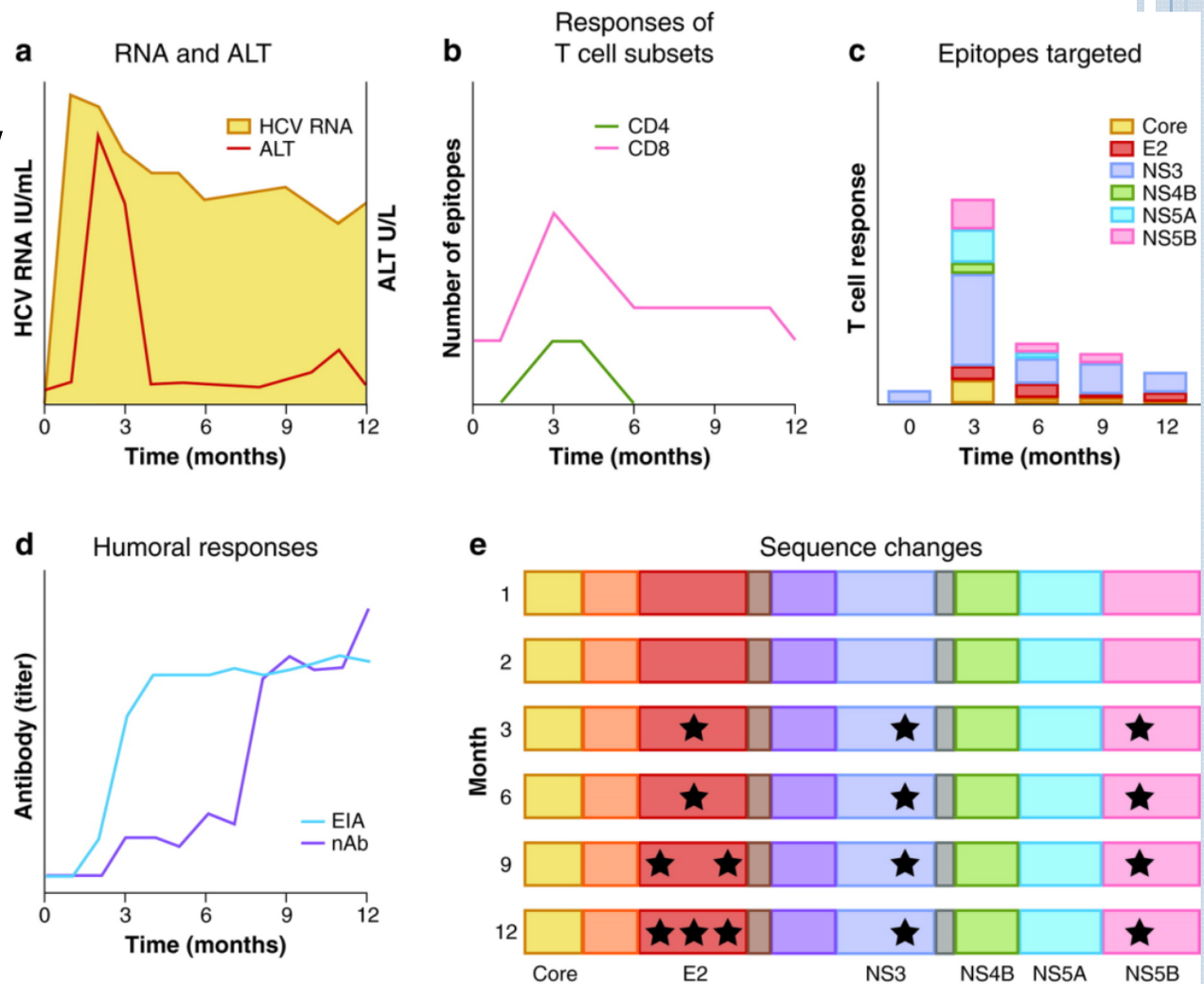
- Broad-based immunological repertoires (targeting multiple epitopes with diverse populations) control acute and prevent the development of chronic infections—particularly CD4 and CD8 cells (the role of antibody is controversial)



**AR** Dustin LB, Rice CM. 2007. Annu. Rev. Immunol. 25:71–99

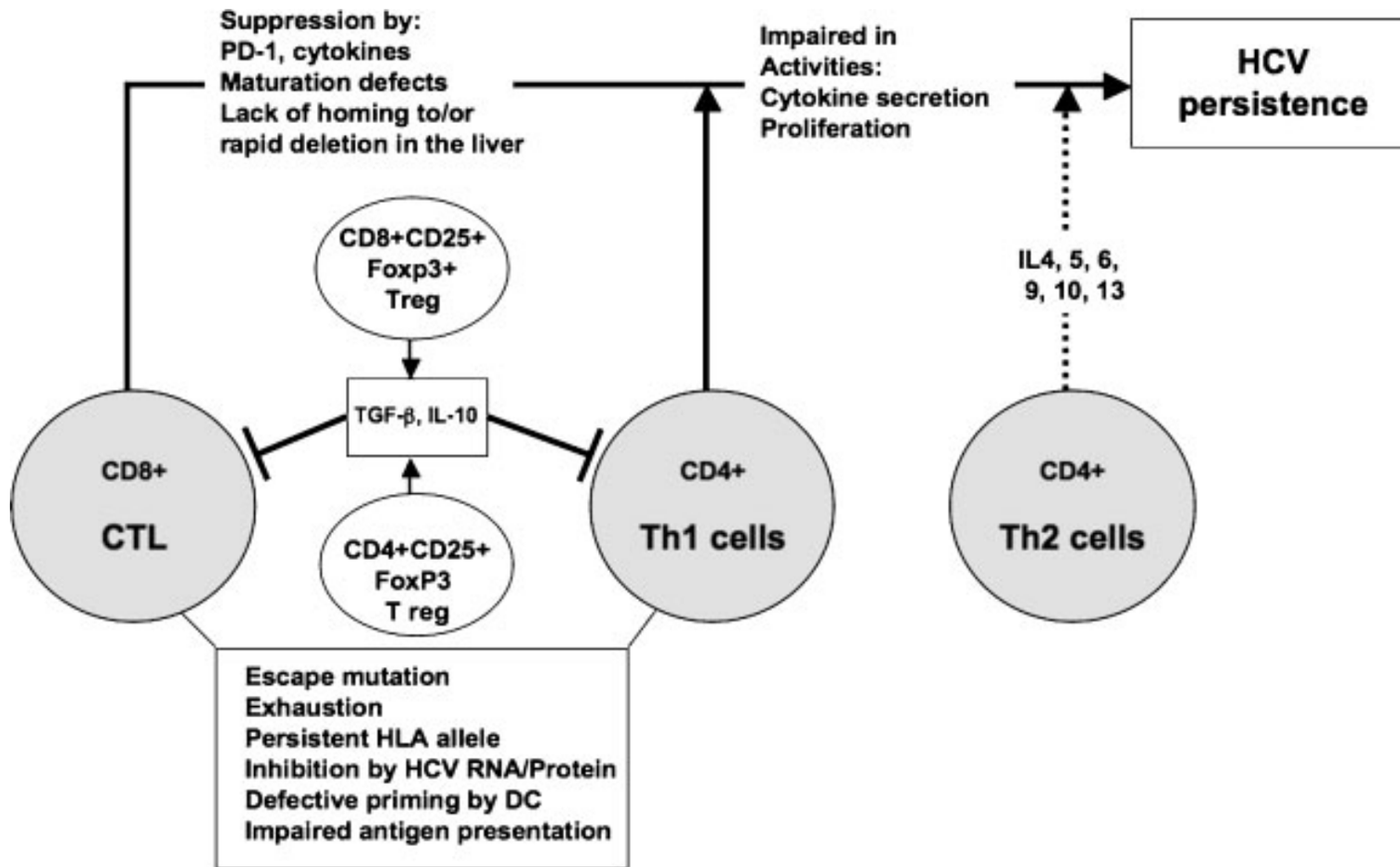
# CHRONIC HCV INFECTIONS RESULT FROM POOR T CELL CONTROL, EPITOPE ESCAPE AND LIMITED REPERTOIRES

- Limited TCR diversity, restricted epitope targets and dysfunctional T cell regulation result in weak T cell responses that are unable to avoid immunological escape



**AR** Dustin LB, Rice CM. 2007.  
Annu. Rev. Immunol. 25:71-99

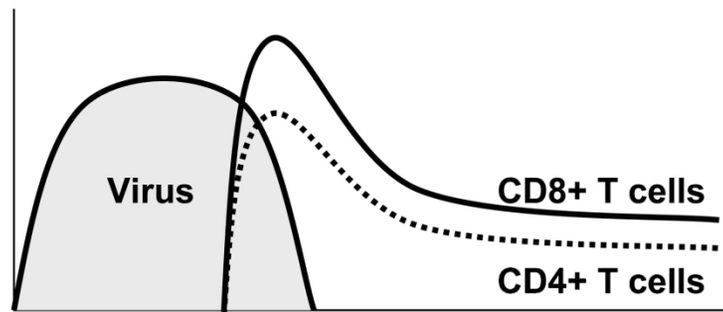
# CHRONIC INFECTIONS AND IMMUNOSUPPRESSION



- Th2 biasing or immune senescence result in the downregulation of aggressive immunological control by CTL, providing the opportunity for viral escape and establishment of chronic infection

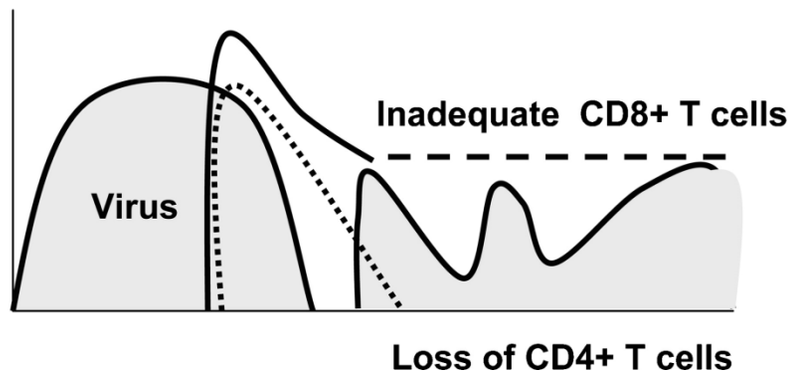
# SUSTAINING AN EFFECTIVE CELLULAR RESPONSE IS MORE IMPORTANT THAN PEAK RESPONSE NUMBERS

## A. Successful immune response



Long-lived memory

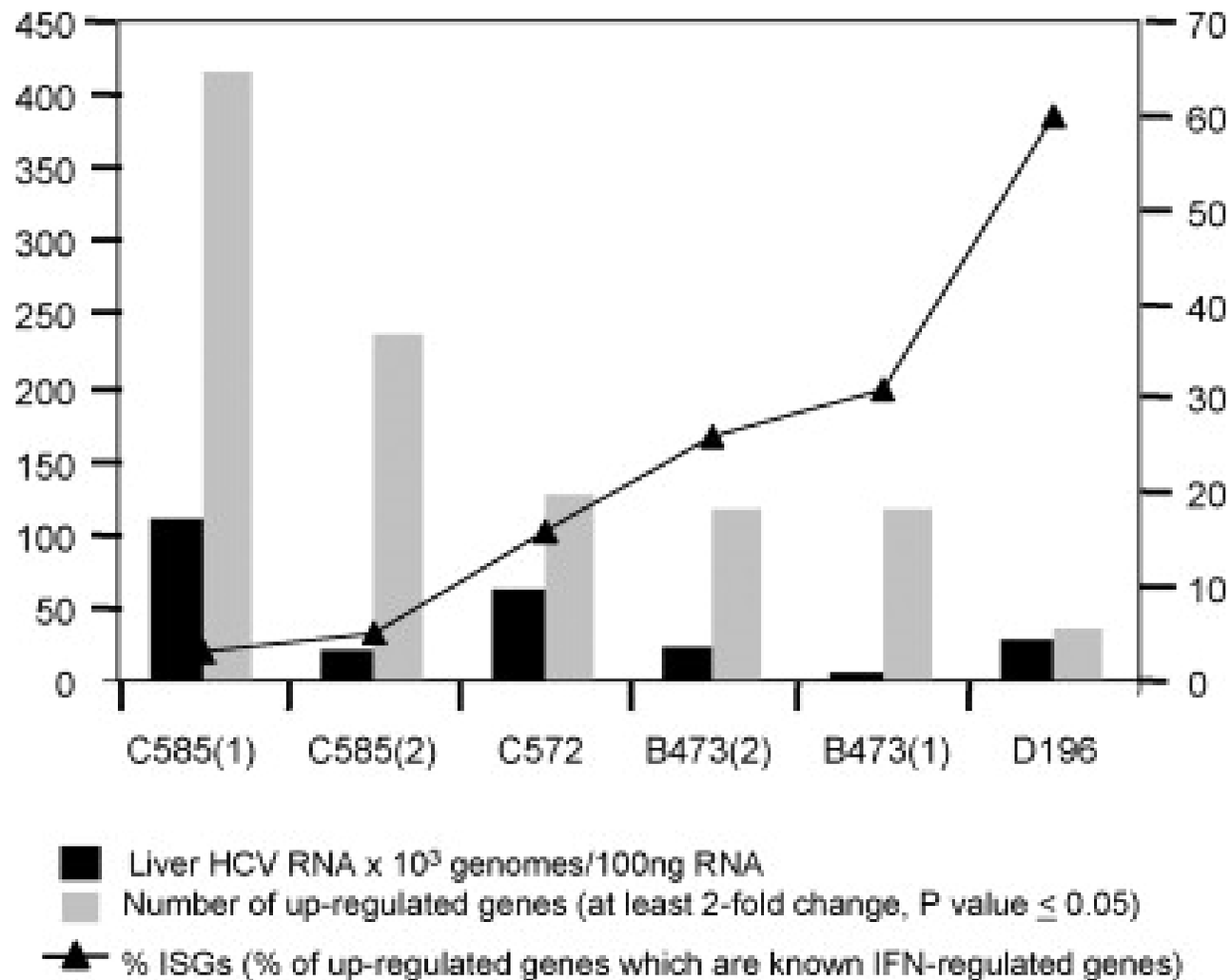
## B. Unsuccessful immune response



Persistent viremia

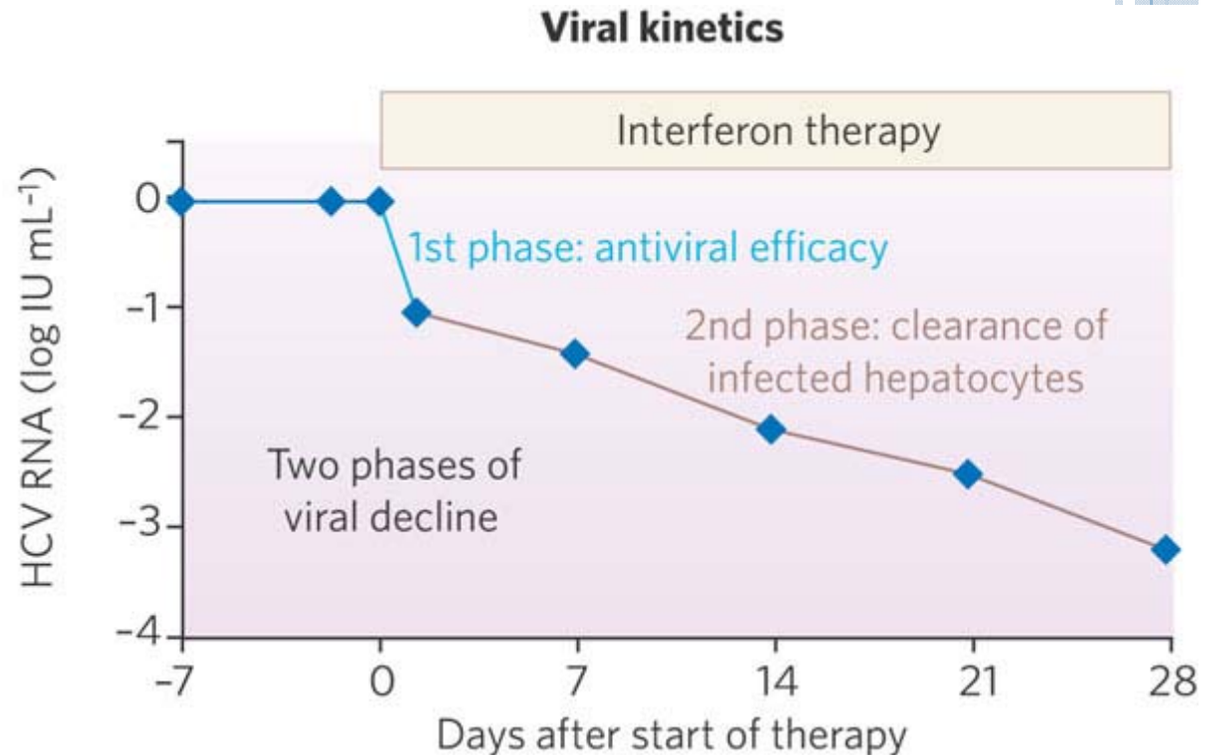


# CONTROL OF ACUTE INFECTION CORRELATES WITH INTERFERON-INDUCED GENES



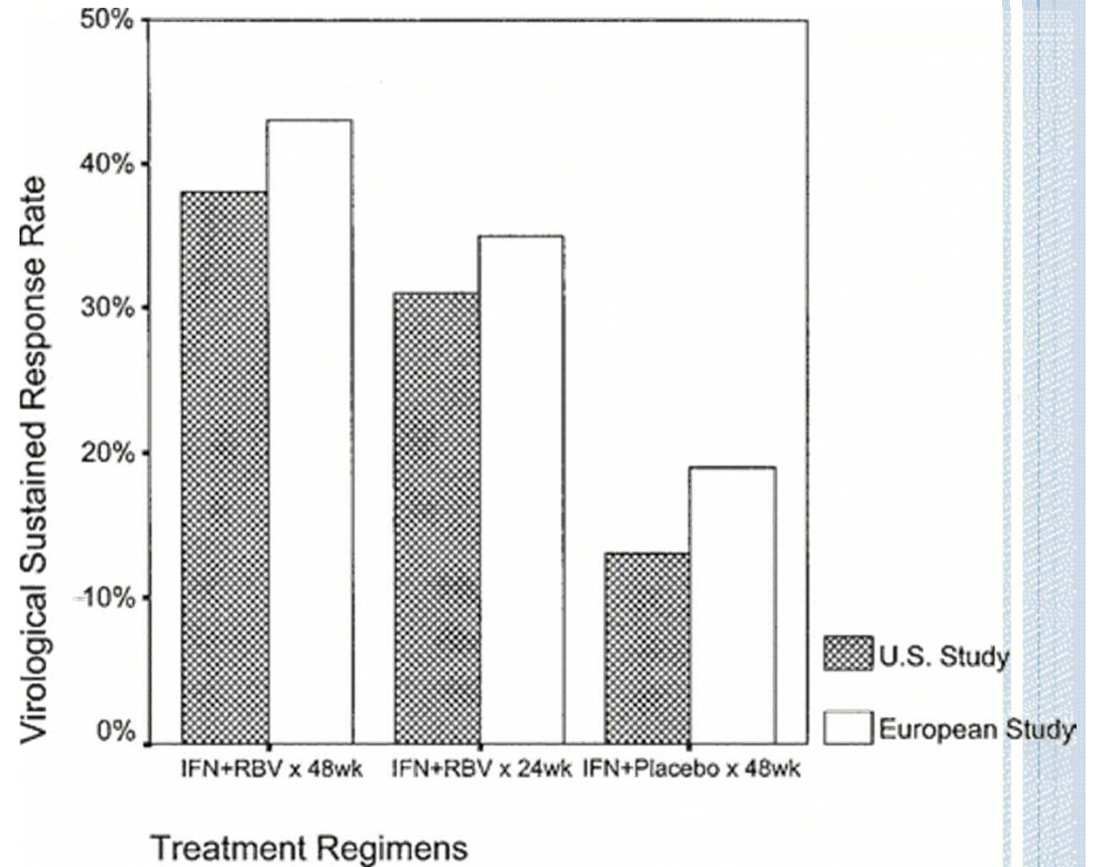
# TREATMENT: TYPE I INTERFERON

- First therapy introduced for HCV
- Full mechanism of action unclear—presumably enhances the “normal” interferon response pathways
- Genotype of virus, low baseline levels of HCV RNA and stage of infection are the strongest correlates of efficacy
- Suggestions that immunomodulation may play a role and that high dose-interferon may overcome some of the “regulatory” negative feedback loops active in the infected host
- Overall, the specific mechanism has not been clearly demonstrated biologically



# COMBINATION THERAPY IS SIGNIFICANTLY MORE EFFECTIVE

- Inteferon alone only yields a 20-25% response rate following a 12-18 month course
- Combination therapy with the “broad based” antiviral ribavirin results in 40% of individuals with SVR (30% genotype 1, 65% genotype 2 or 3)

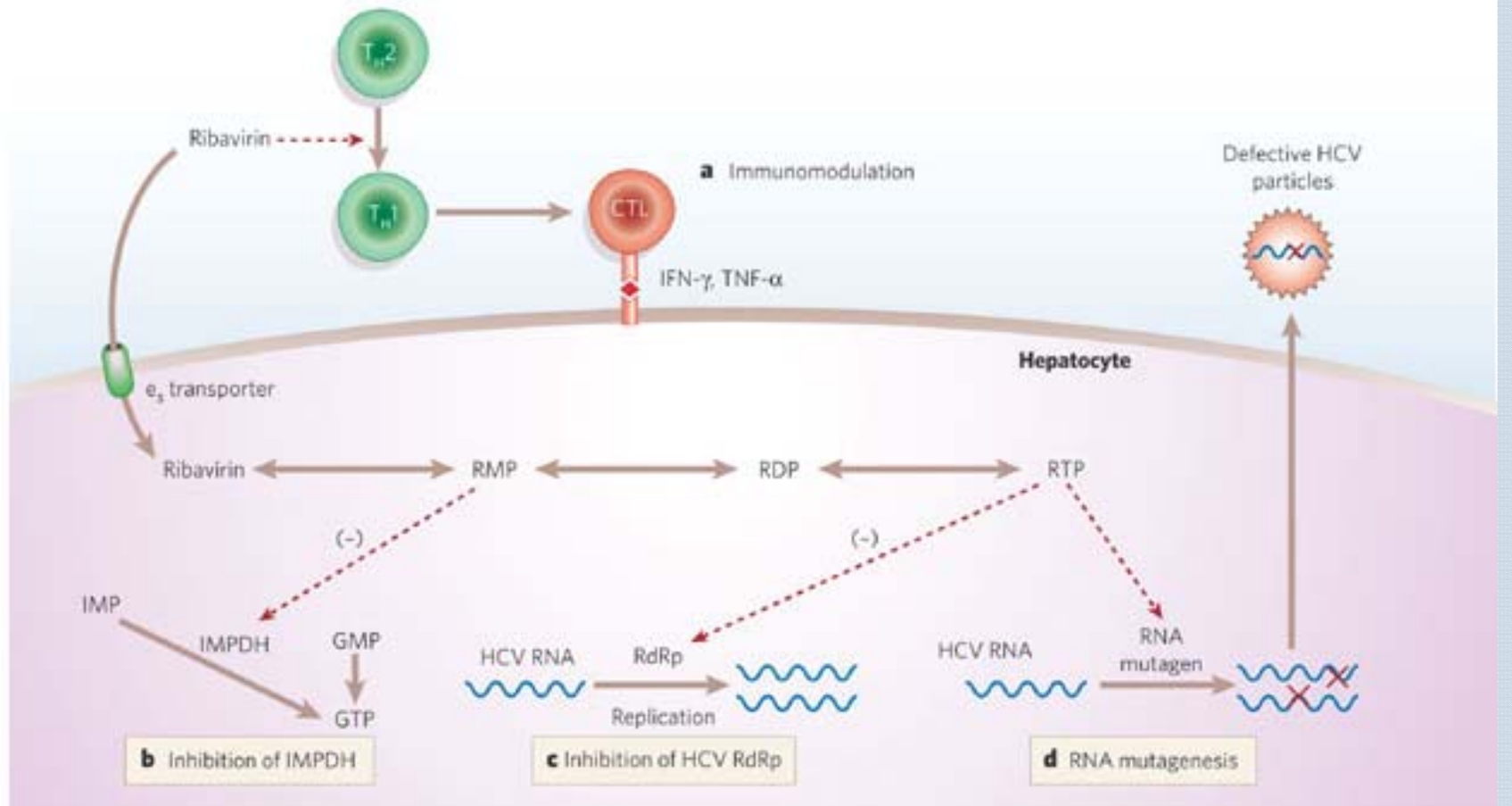


## HOW DOES RIBAVIRIN WORK AGAINST HCV?

- Ribavirin was initially designed as a nucleoside analog and developed as an anti-influenza drug, but failed to receive FDA approval or show significant efficacy in humans
- It has been used to treat hemorrhagic fevers, RSV and is again under consideration as combination therapy for influenza
- Proposed Mechanisms:
  - 1) Immunomodulatory properties
  - 2) Inhibition of the inosine monophosphate dehydrogenase (IMPDH)
  - 3) Direct inhibition of the HCV-encoded NS5B RNA polymerase
  - 4) Induction of lethal mutagenesis
  - 5) Modulation of interferon-stimulated gene (ISG) expression



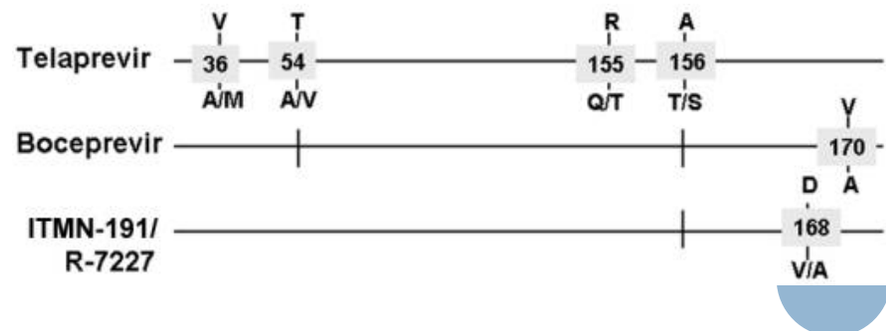
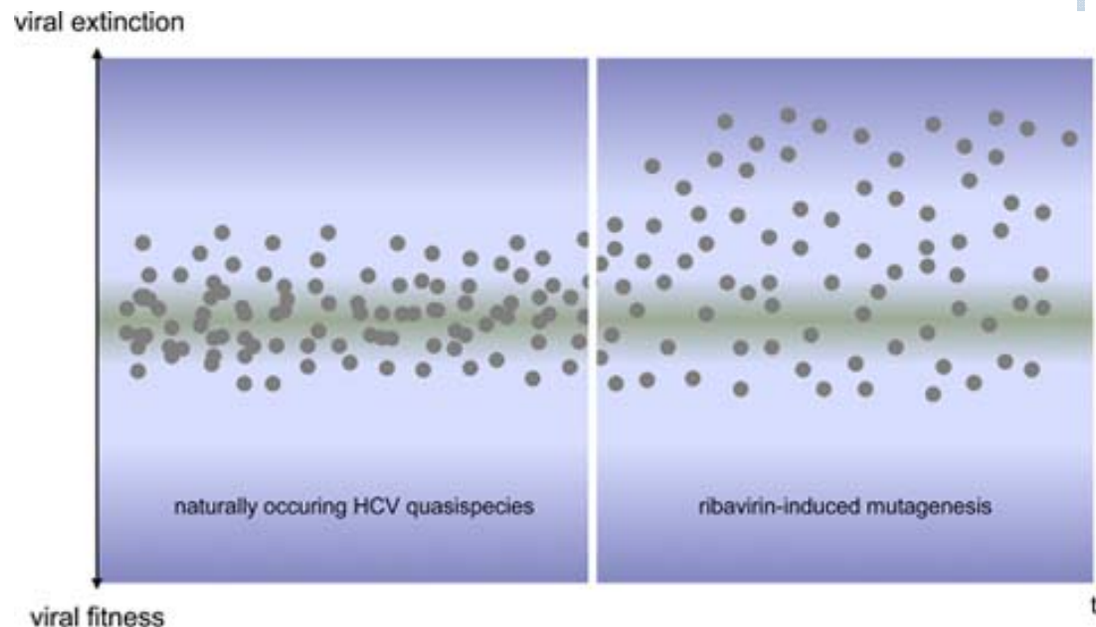
# POSSIBLE MECHANISMS FOR RIBAVIRIN MODE OF ACTION



# WHAT DATA WOULD HELP RESOLVE RIBAVIRIN'S MECHANISM?

Interferon reduces viral production-- given the proposed mechanisms, how should ribavirin work?

- 1) Immunomodulatory properties—**Should act independently of interferon**
- 2) Inhibition of the inosine monophosphate dehydrogenase (IMPDH)—**Should reduce viral production, be guanosine dependent**
- 3) Direct inhibition of the HCV-encoded NS5B RNA polymerase—**Should reduce viral production, put pressure on polymerase to mutate**
- 4) Induction of lethal mutagenesis—**Viral production maintained, infected cell number maintained (clearance by decay), new cells infected at a lower rate**
- 5) Modulation of interferon-stimulated gene (ISG) expression—**Direct antiviral effects like interferon, should shift ISG expression from negative feedback pathways and be synergistic with poor interferon responders.**

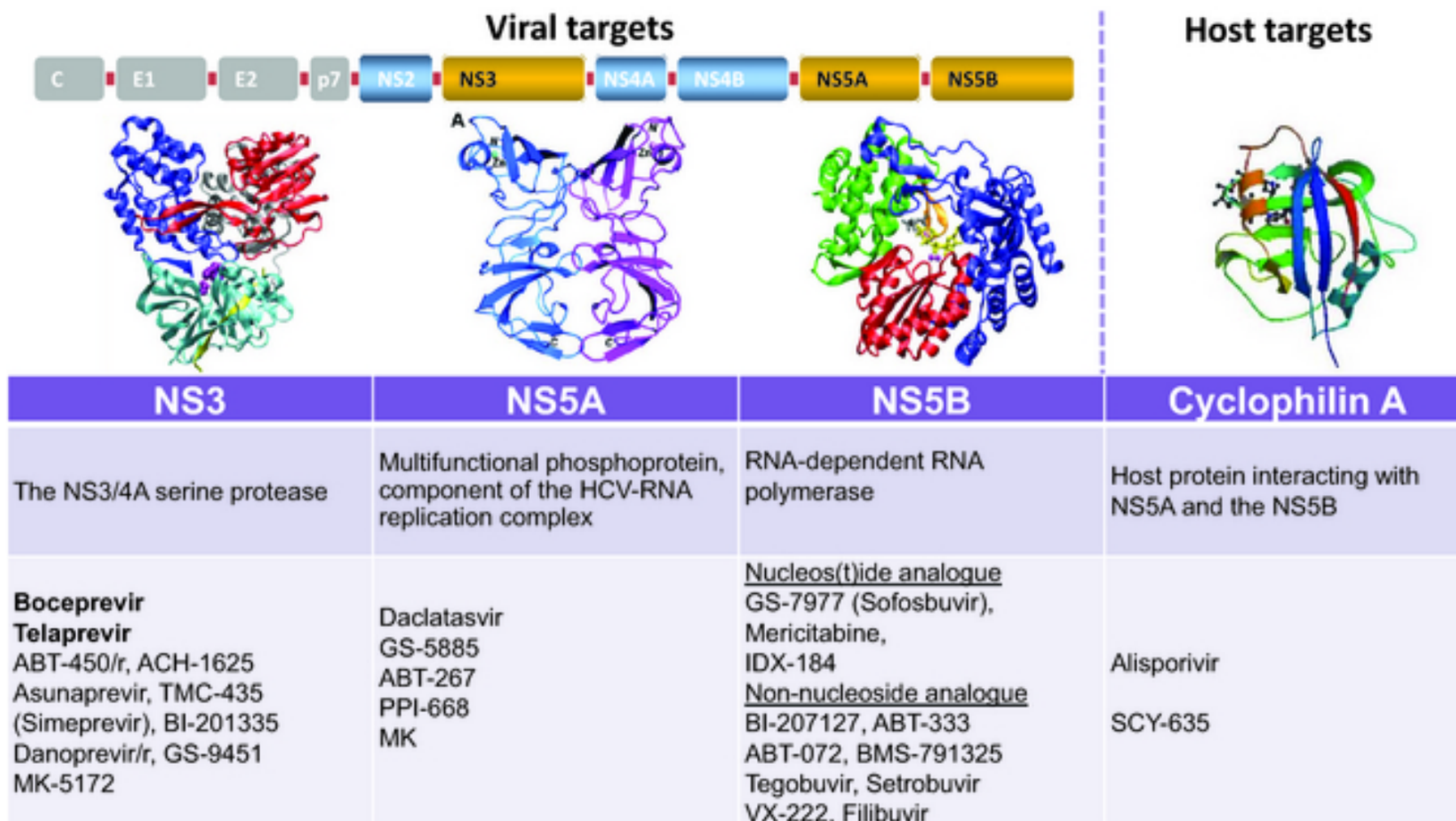


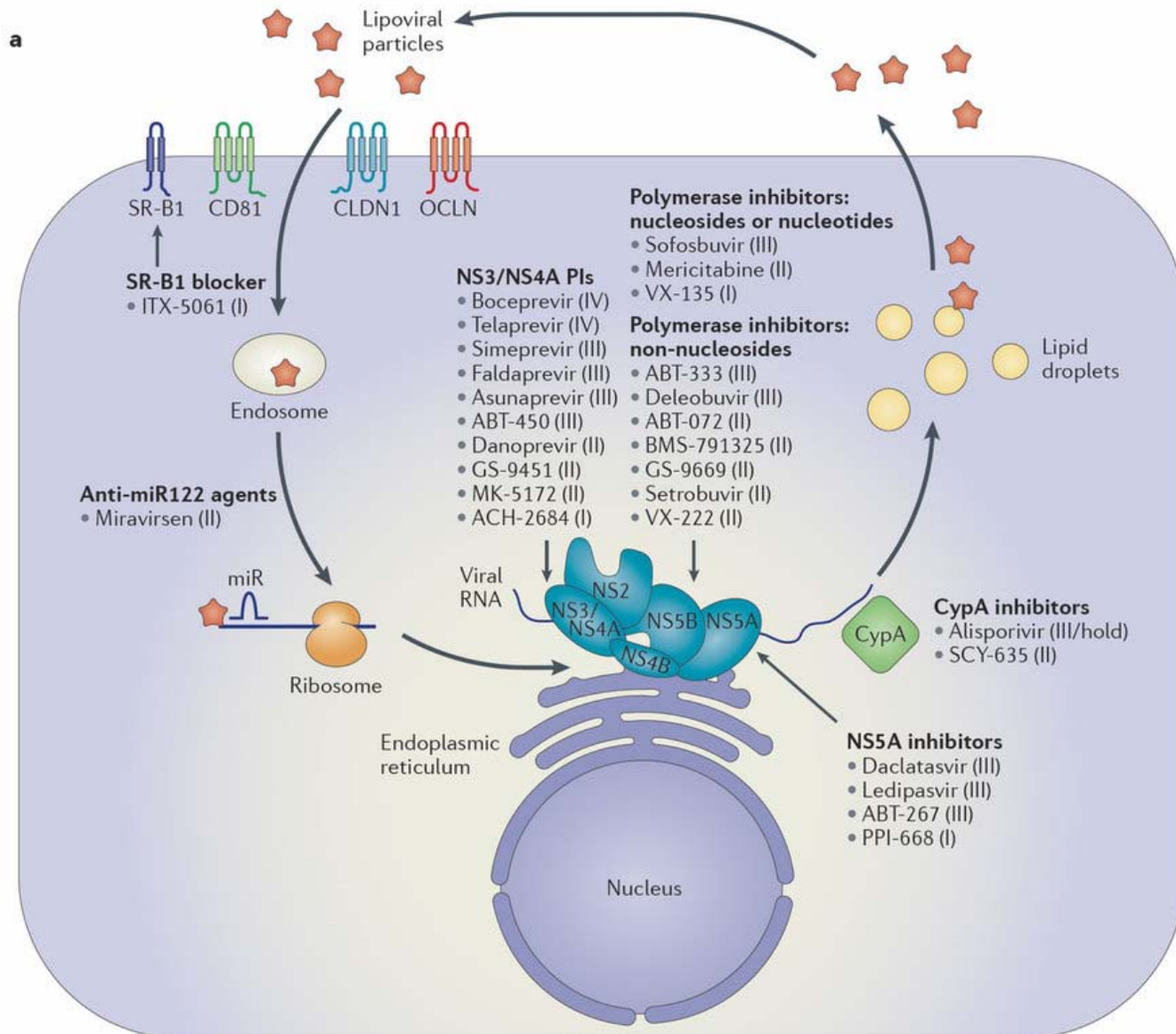
## DETERMINING AN ANTIVRAL TREATMENT'S MODE OF ACTION

- Biological *in vitro* experiments with HCV have been difficult to perform as a result of the limited nature of developed culture systems
- Alternative drugs that perform a single “ribavirin function” do not recapitulate ribavirin efficacy, suggesting that multiple pathways may be acting together
- Biological mechanisms can often seem plausible, but can be difficult to prove conclusively that they play an important role (particularly when the drug is “reverse engineered” to the pathogen)
- Mathematical modeling from real infection data provides a compelling argument for the viral life cycle stage(s) that might be affected



# NEW DRUG TREATMENTS FOR HCV





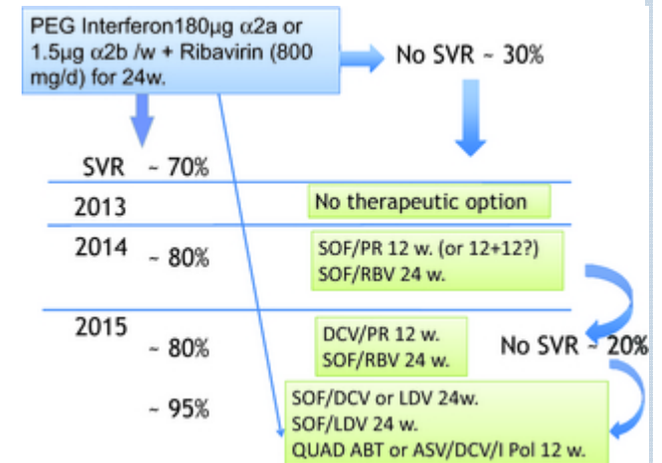
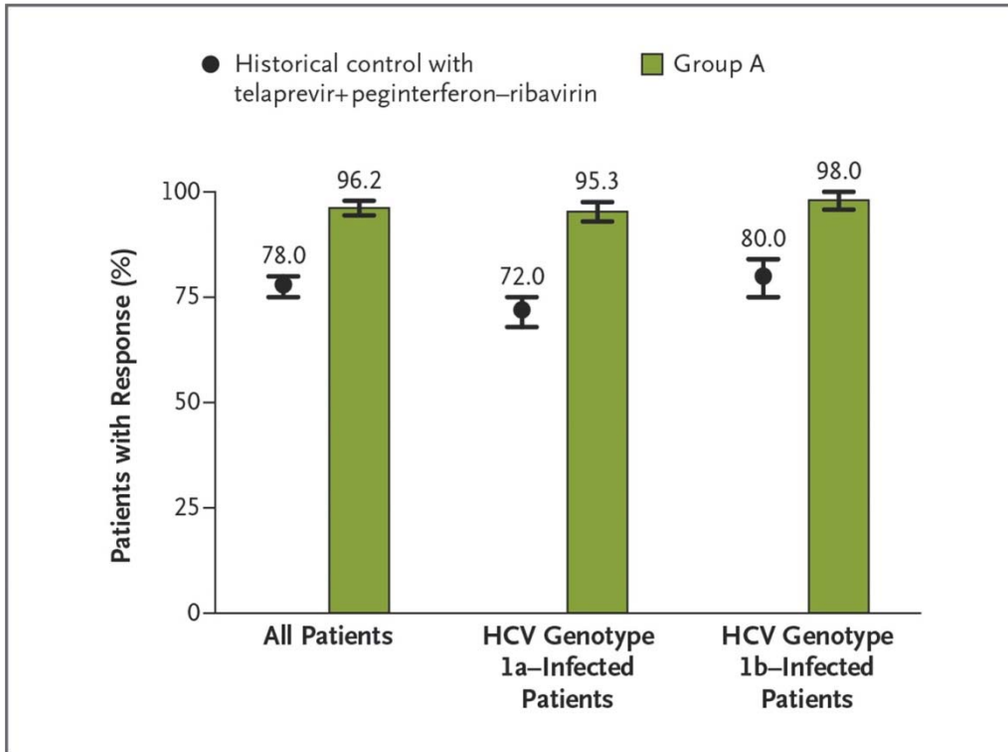
Novel therapies for hepatitis C — one pill fits all?

Michael P. Manns

& Thomas von Hahn

Nature Reviews Drug Discovery 12, 595–610 (2013) doi:10.1038/nrd4050

## Rates of Sustained Virologic Response among All Patients and According to HCV Genotype in the Historical Control Group and in Group A.

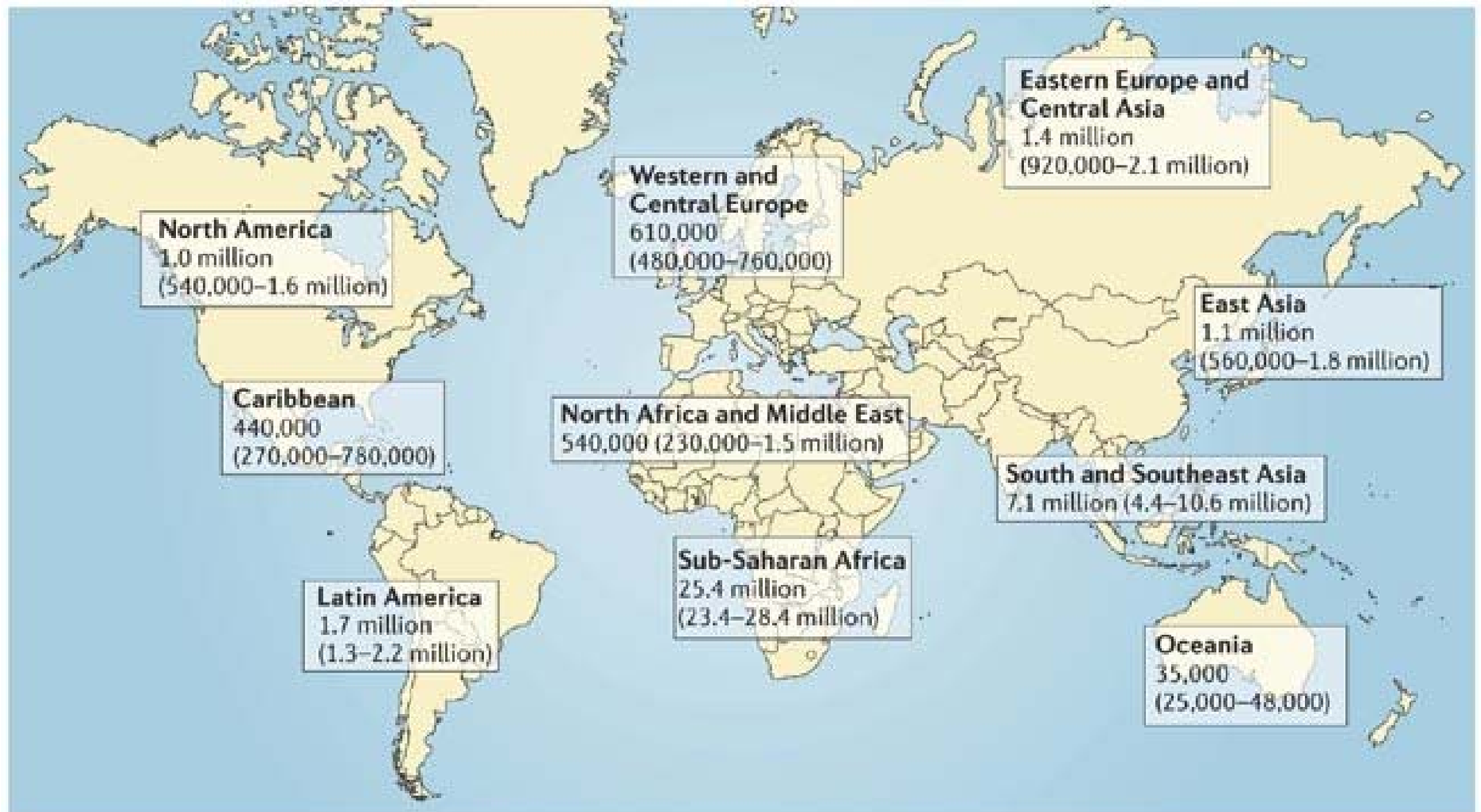


FELD JJ ET AL. N ENGL J MED 2014;370:1594-1603.

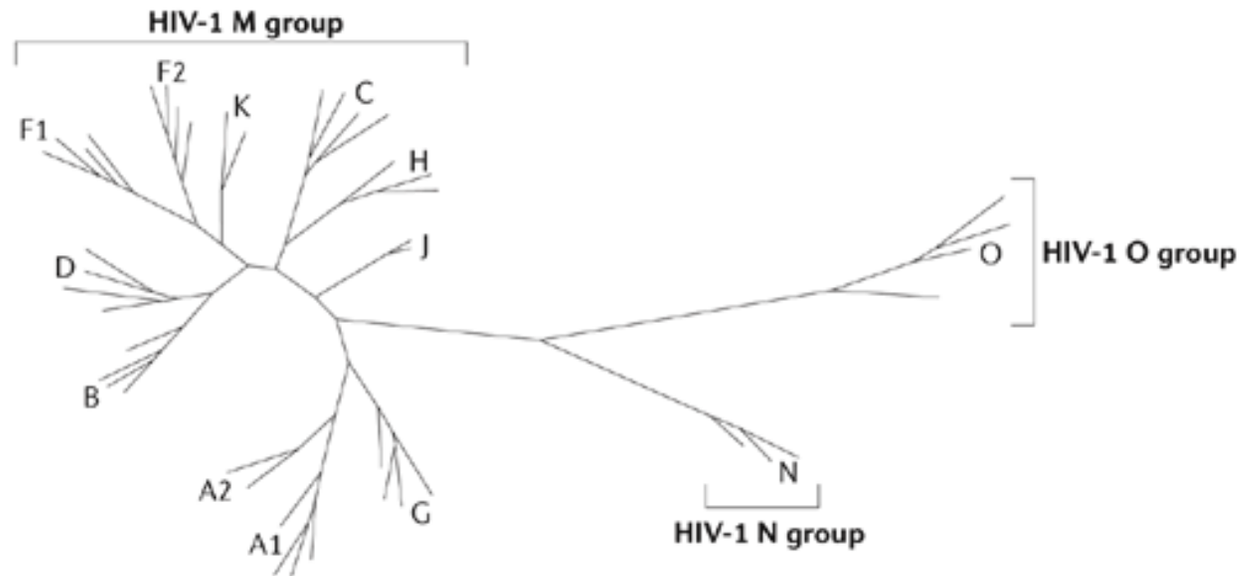


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# PREVALENCE OF HIV INFECTION



# GENETIC DIVERSITY OF HIV-1

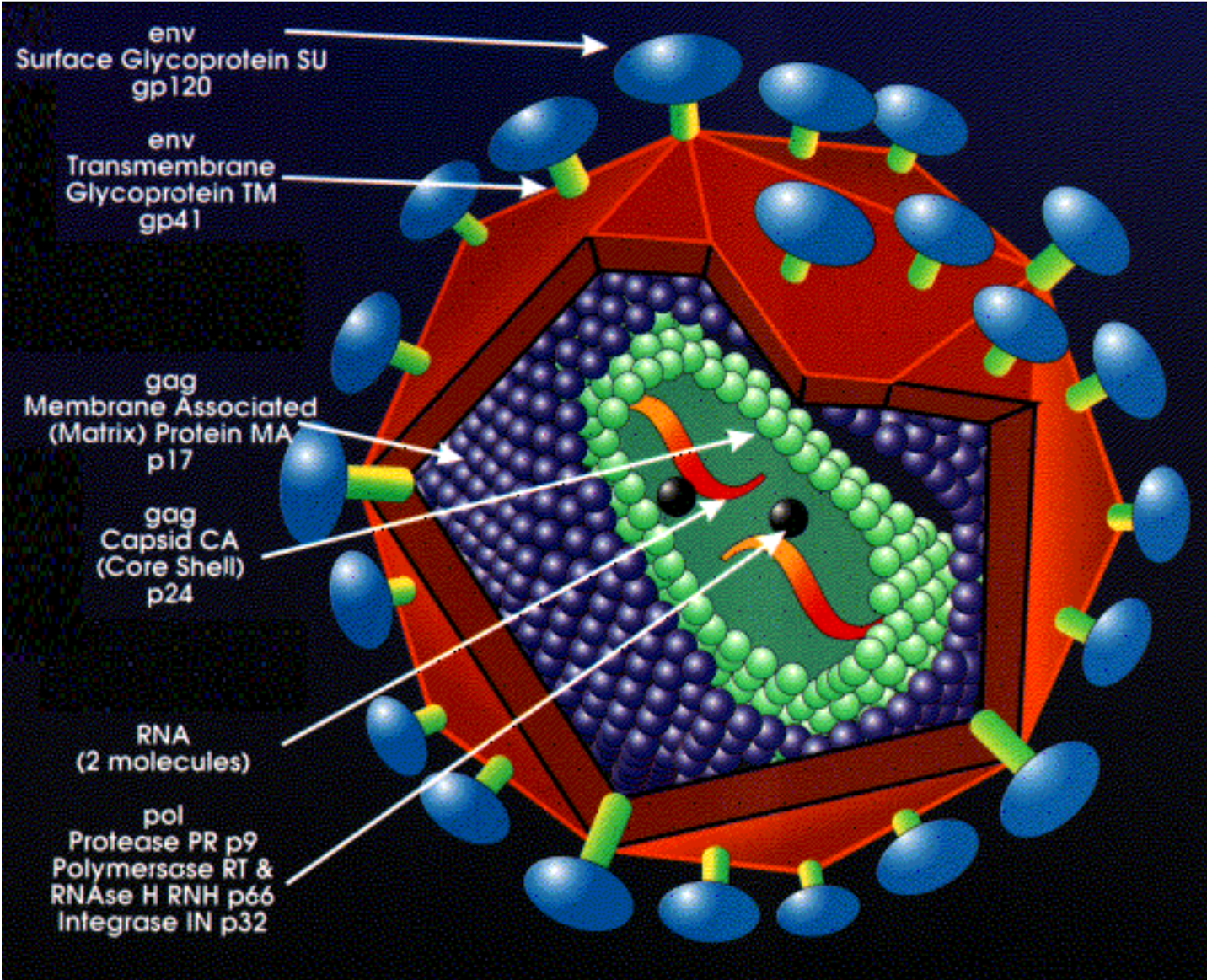


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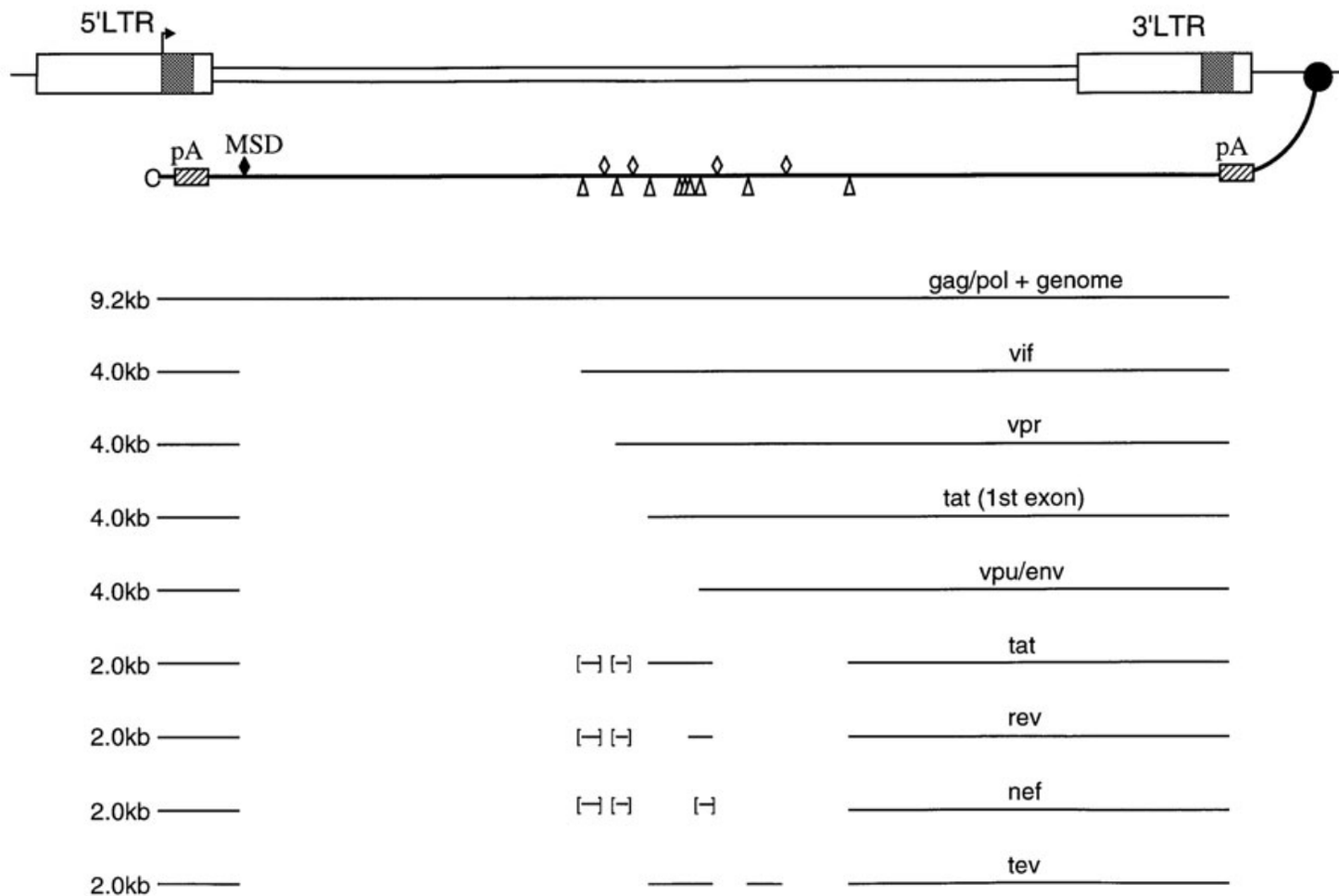
- Within HIV-1, a large sequence diversity exists with viral clades being geographically isolated
- Several studies have suggested that the clades have different biological characteristics, including disease pathogenicity and transmissibility



# VIRION STRUCTURE

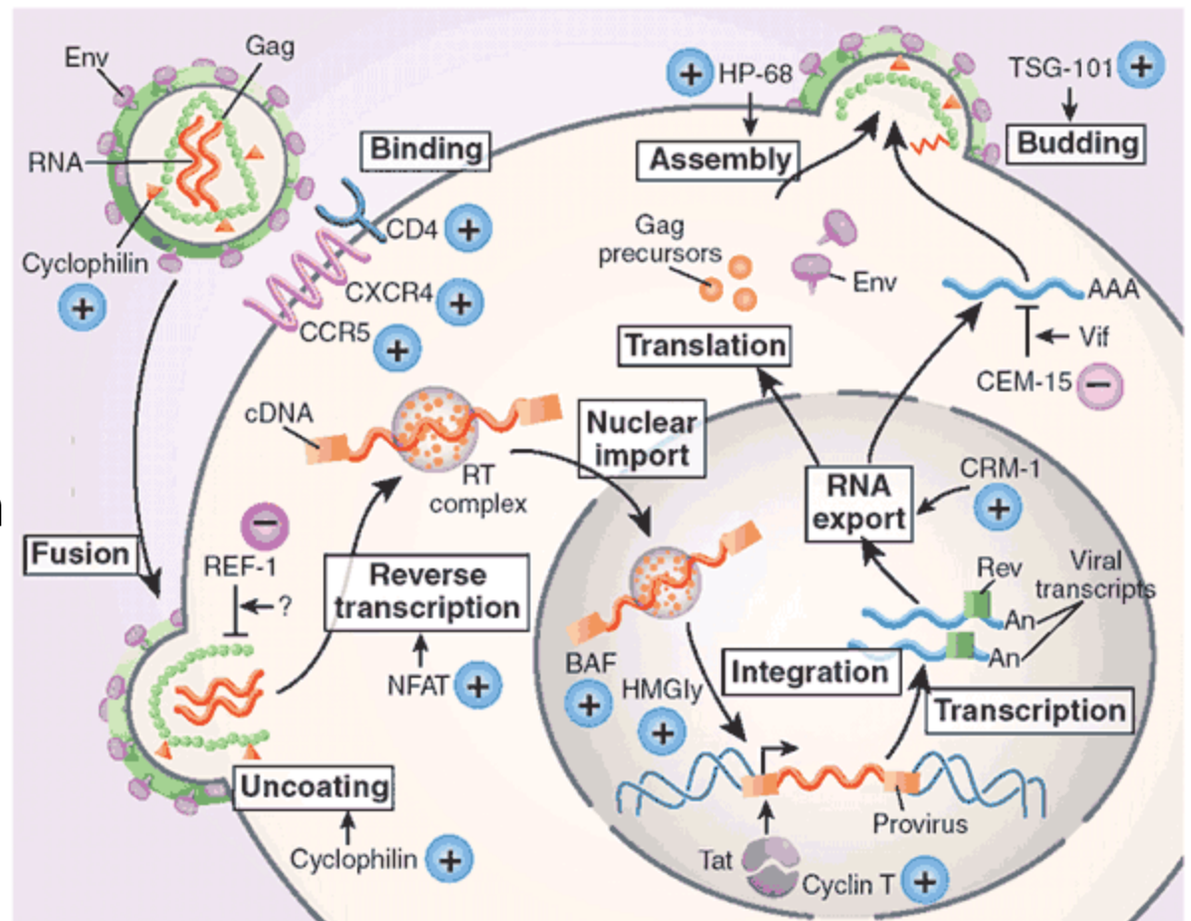


# SINGLE STRANDED GENOME, MULTIPLE MESSAGES FROM ALTERNATIVE SPLICING



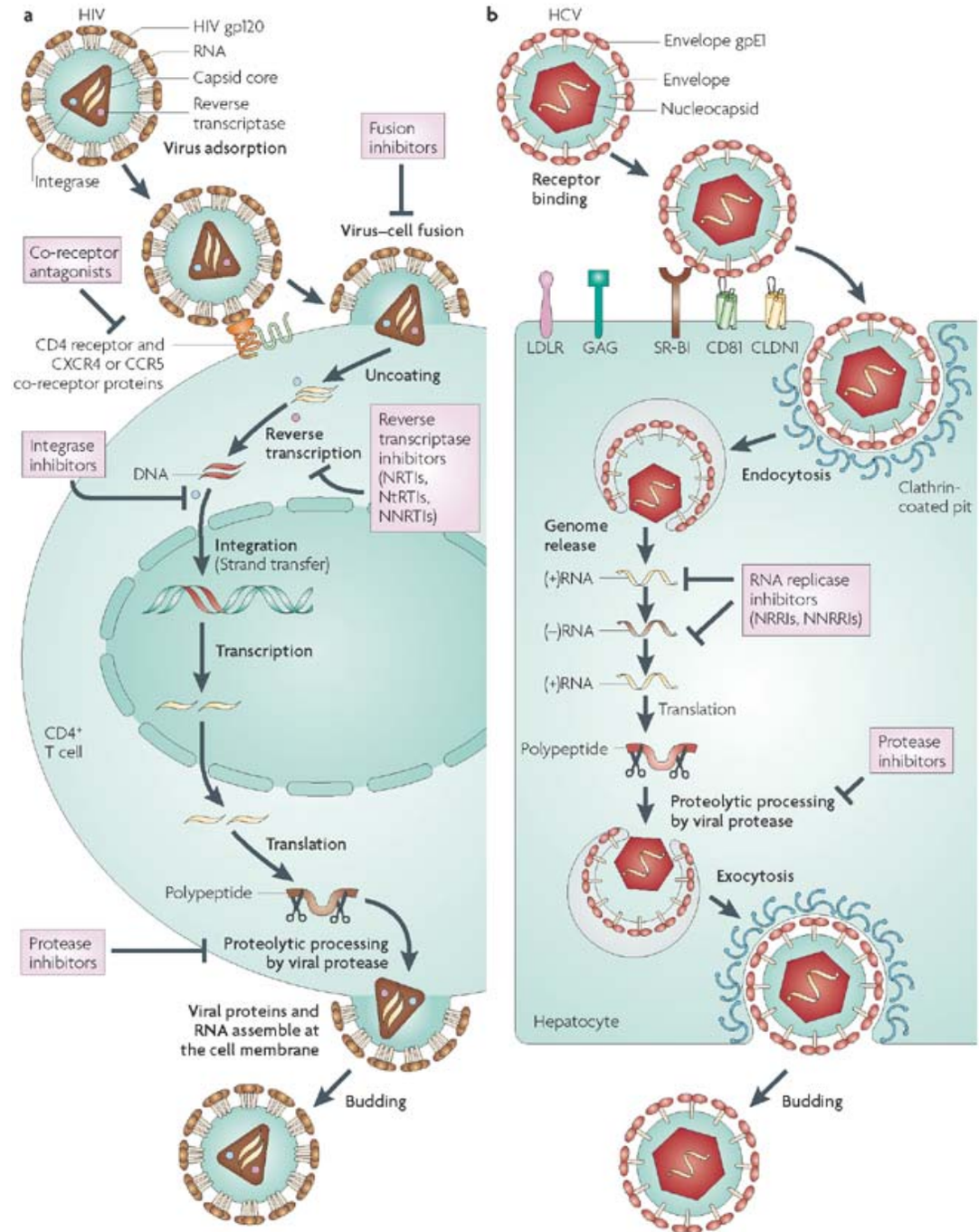
# VIRAL LIFE CYCLE

- As a retrovirus, HIV replicates by making a DNA copy of itself that is inserted into the host genome
- Thus, an infected cell can become a stable reservoir for the long term production of viral particles

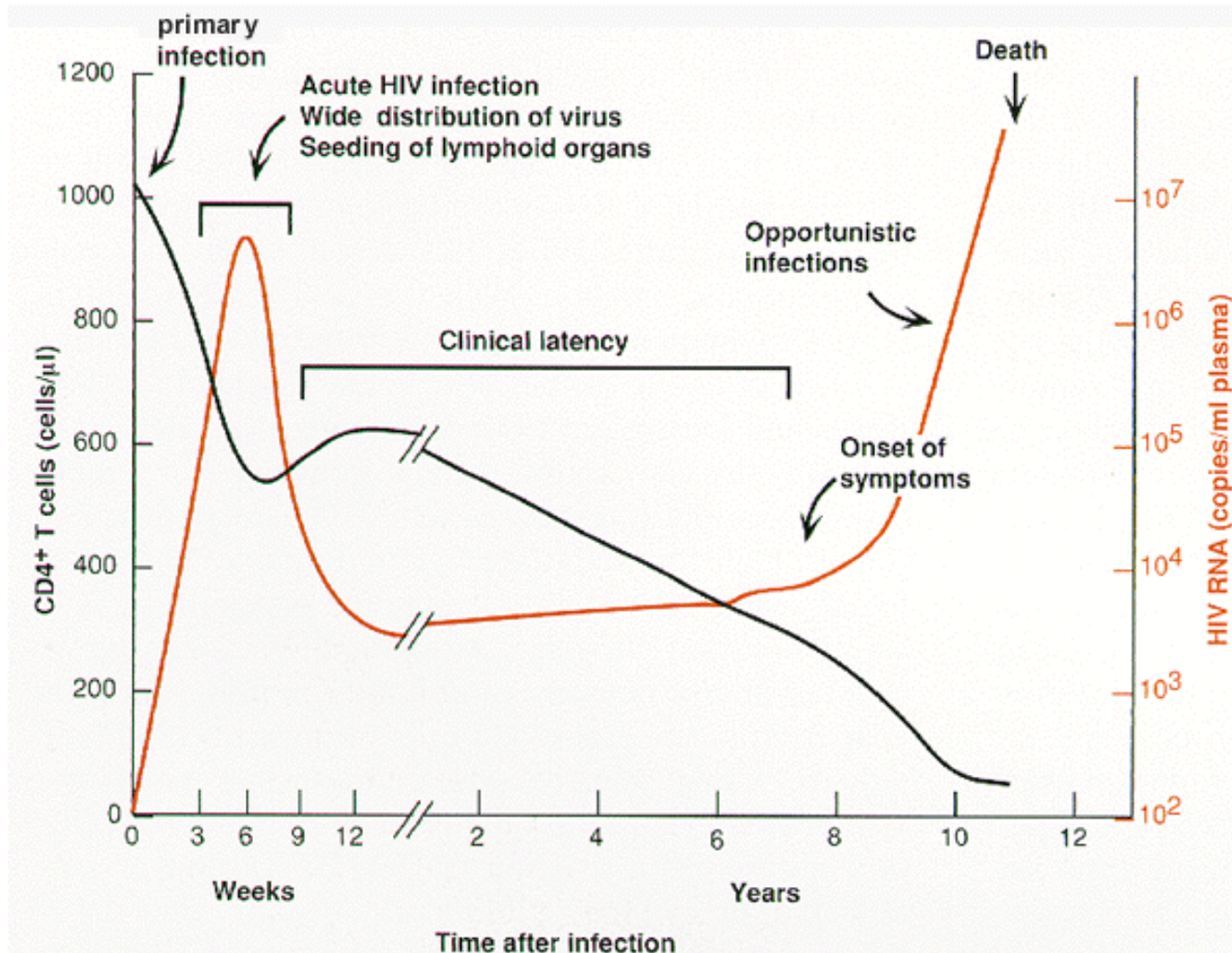


# COMPARISON OF HIV AND HCV

- HIV and HCV both produce chronic infections, but are biologically very different viruses
- HIV has a DNA intermediate that become heritably integrated
- HCV is a purely RNA virus

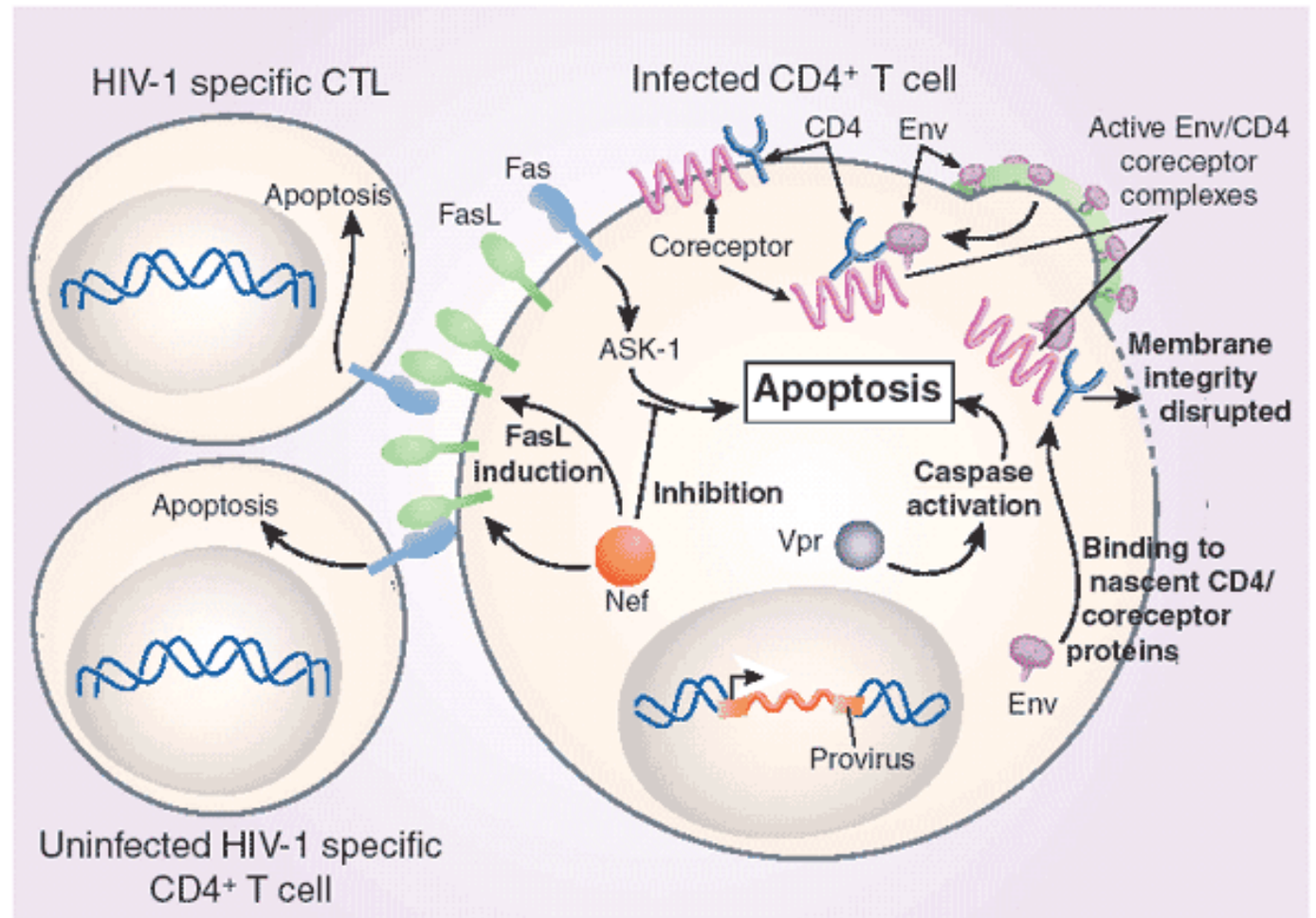


# CLINICAL COURSE OF INFECTION



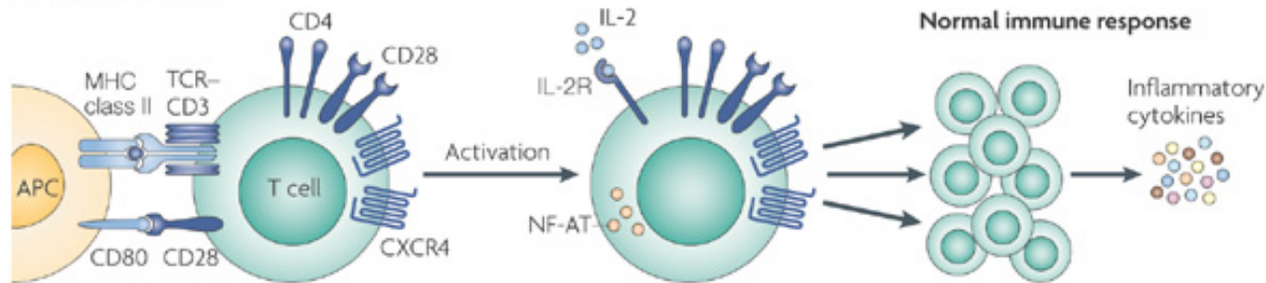
# MECHANISMS OF CYTOPATHOGENICITY

- Viral envelope fusogenicity (ER compromised)
- Vpr activates caspases
- Nef contributes indirectly to apoptosis via FasL



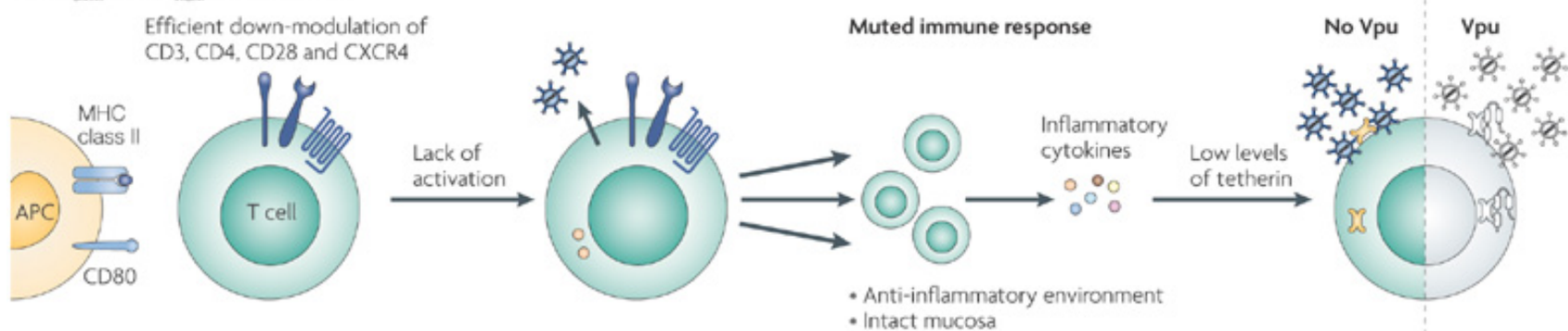
# MECHANISMS OF IMMUNE DYSREGULATION

## a Uninfected T cell



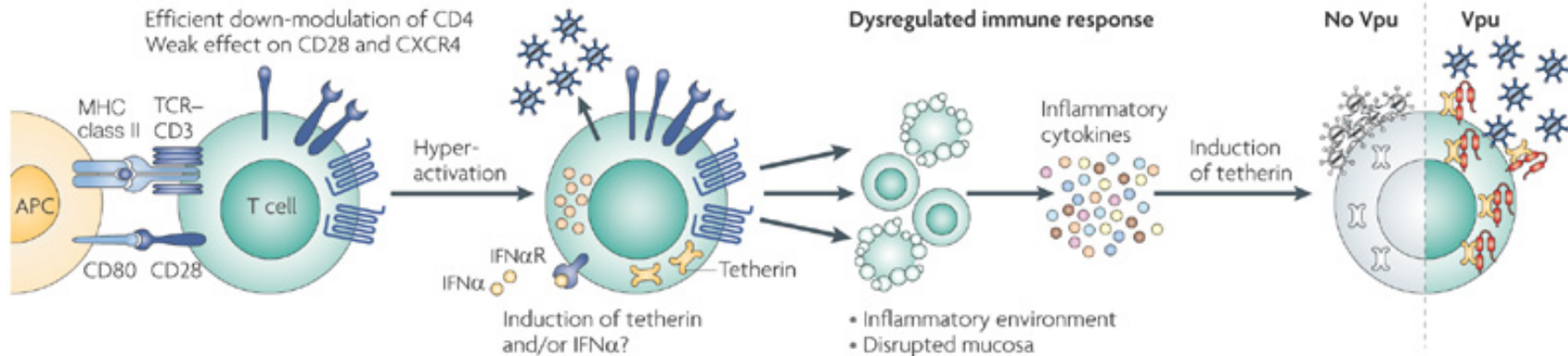
## b SIV<sub>SMM</sub><sup>-</sup> or SIV<sub>AGM</sub>-infected T cell

Efficient down-modulation of CD3, CD4, CD28 and CXCR4



## c HIV-1-infected T cell

Efficient down-modulation of CD4  
 Weak effect on CD28 and CXCR4



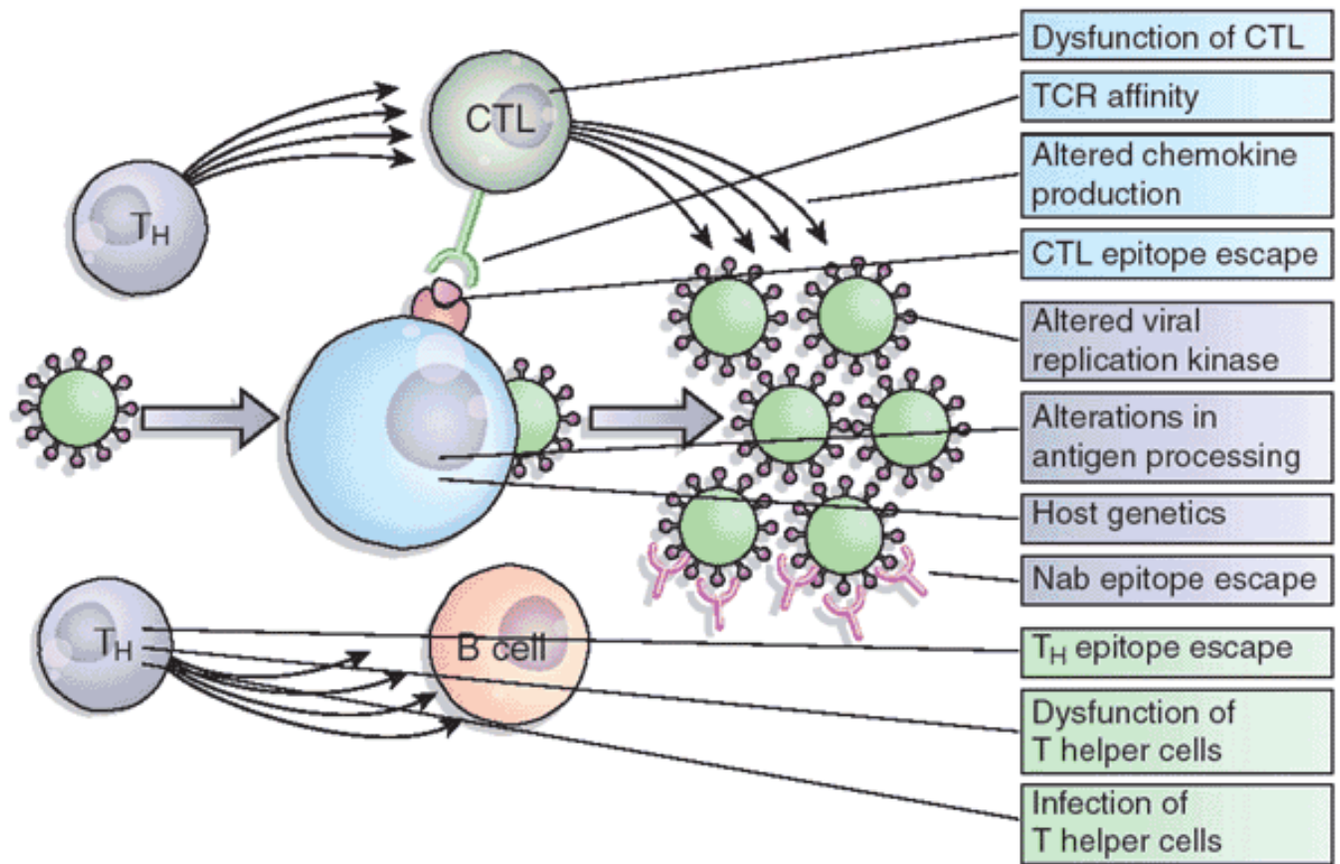
# WHAT MAKES HIV LETHAL?

Infections		Malignancies
Parasites	<i>Toxoplasma</i> spp. <i>Cryptosporidium</i> spp. <i>Leishmania</i> spp. <i>Microsporidium</i> spp.	Kaposi's sarcoma - HHV8 Non-Hodgkin's lymphoma, including EBV-positive Burkitt's lymphoma Primary lymphoma of the brain
Intracellular bacteria	<i>Mycobacterium tuberculosis</i> <i>Mycobacterium avium intracellulare</i> <i>Salmonella</i> spp.	
Fungi	<i>Pneumocystis carinii</i> <i>Cryptococcus neoformans</i> <i>Candida</i> spp. <i>Histoplasma capsulatum</i> <i>Coccidioides immitis</i>	
Viruses	Herpes simplex Cytomegalovirus Varicella zoster	

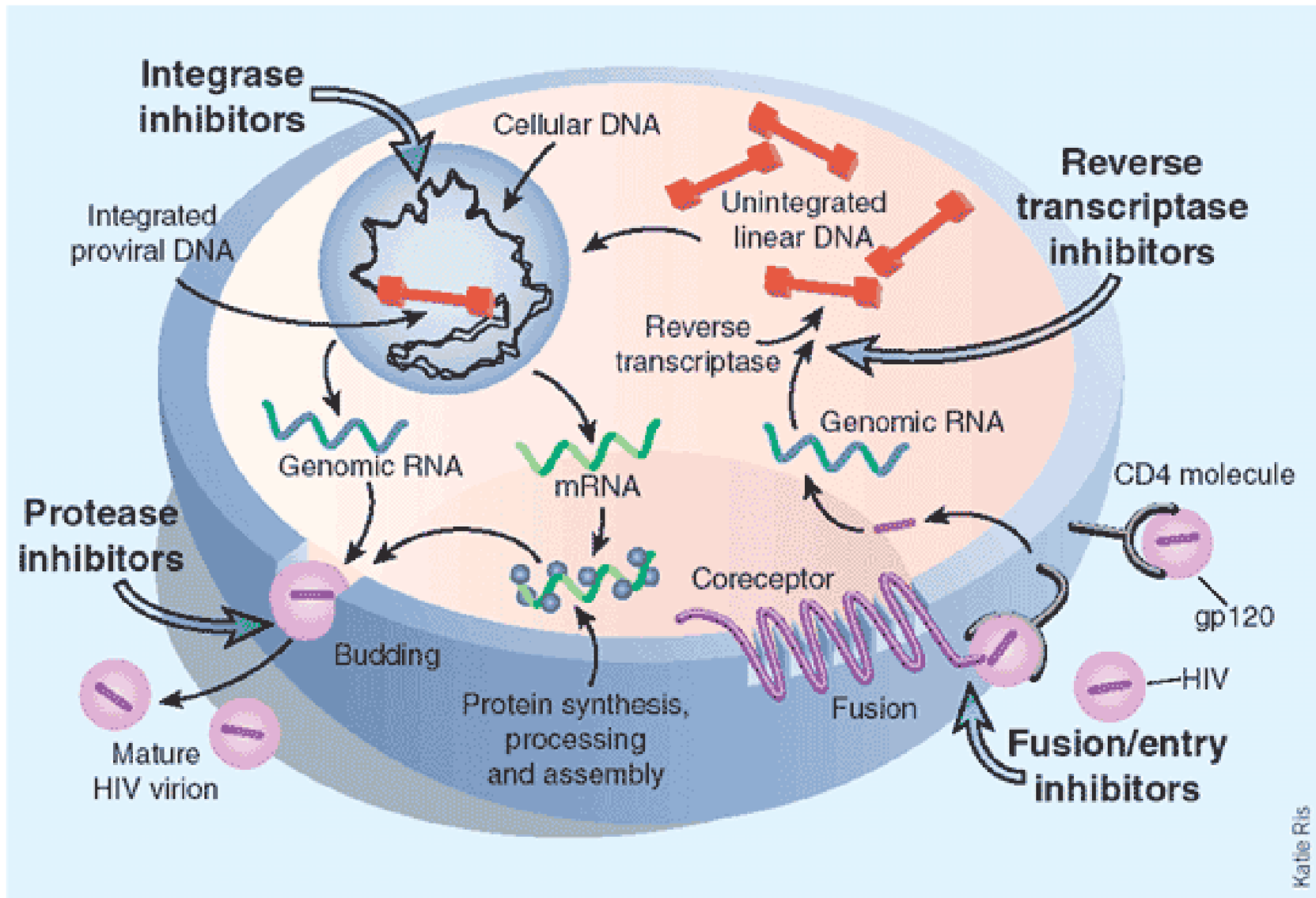
Figure 11-30 Immunobiology, 6/e. (© Garland Science 2005)

# WHY IS HIV UNLIKE ANY OTHER CHRONIC INFECTION?

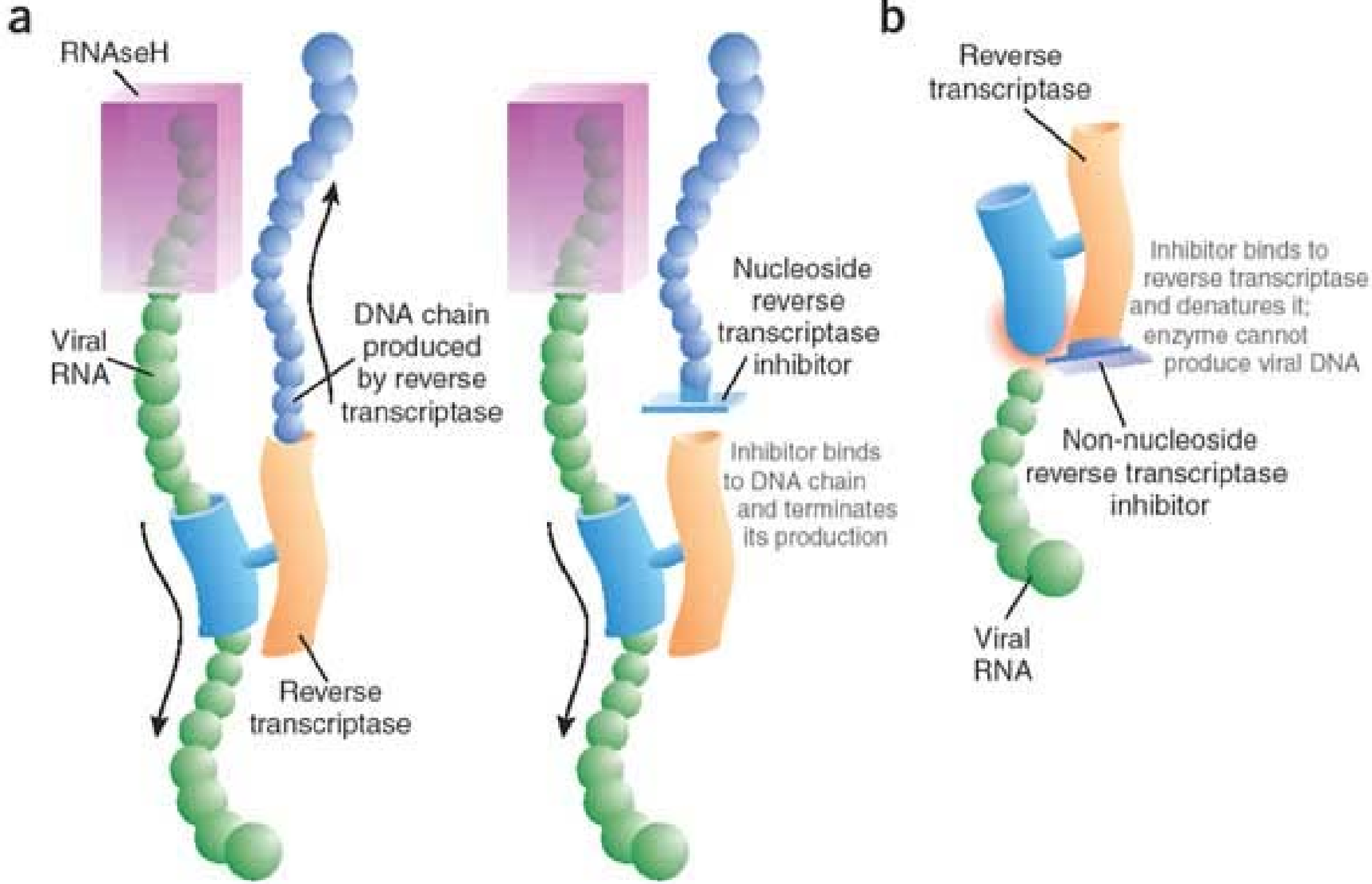
- A combination of “traditional” immune evasion mechanisms (CTL escape, antigen masking) and non-traditional (attacking immune function and cell compartments directly)



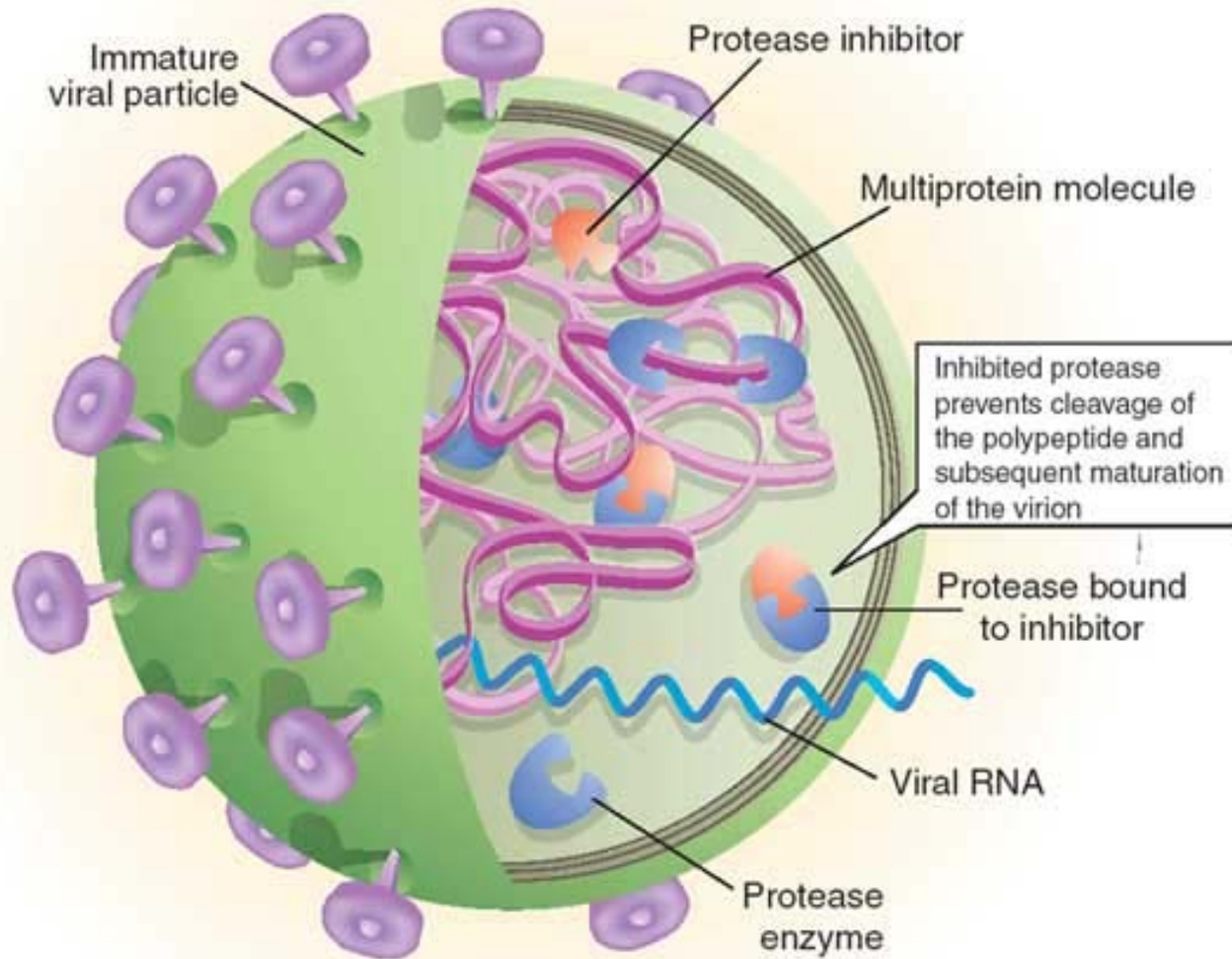
# CAN INFECTION BE EFFECTIVELY CONTROLLED?



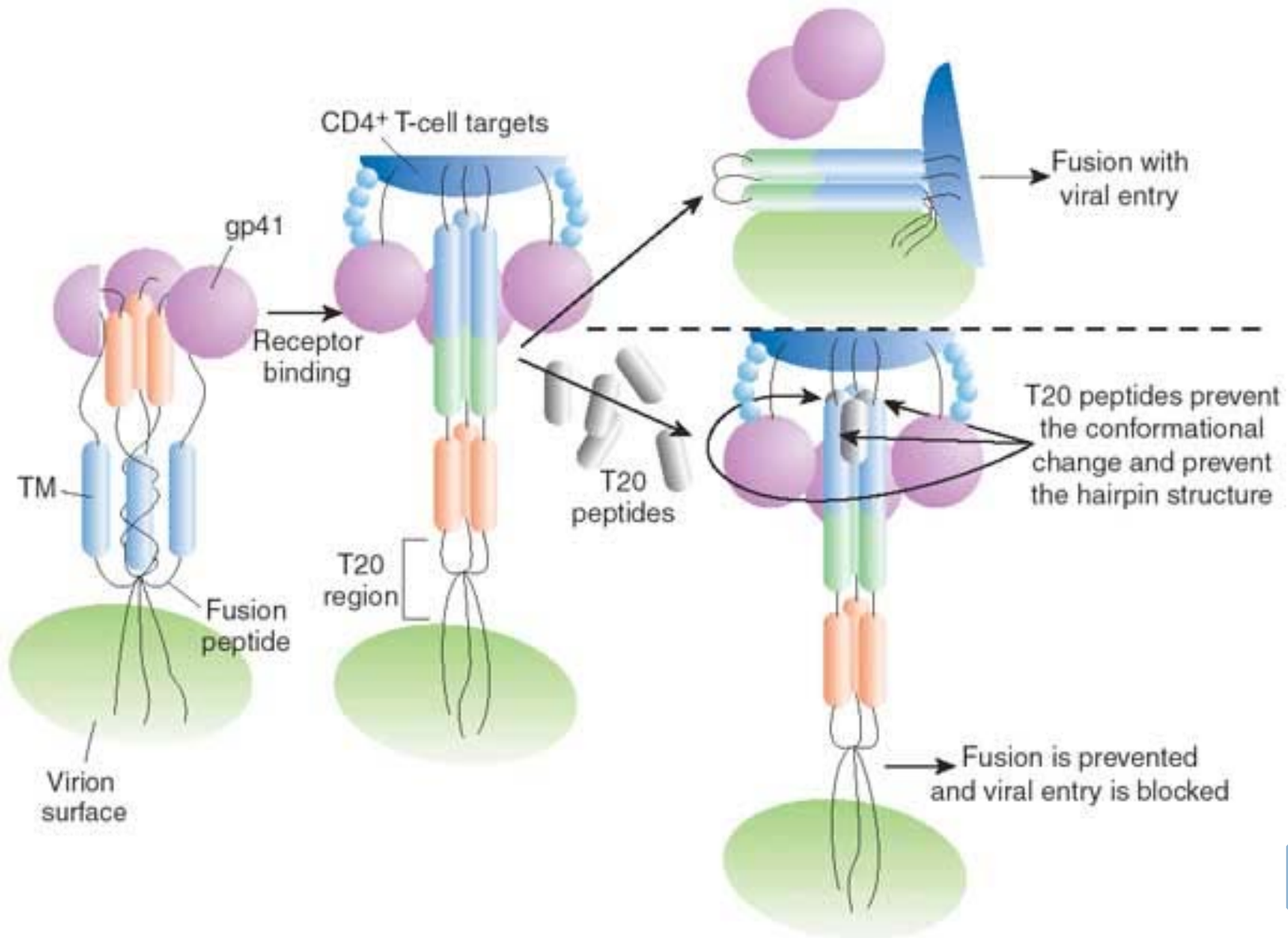
# MECHANISMS OF RT INHIBITORS



# MECHANISM OF PROTEASE INHIBITORS

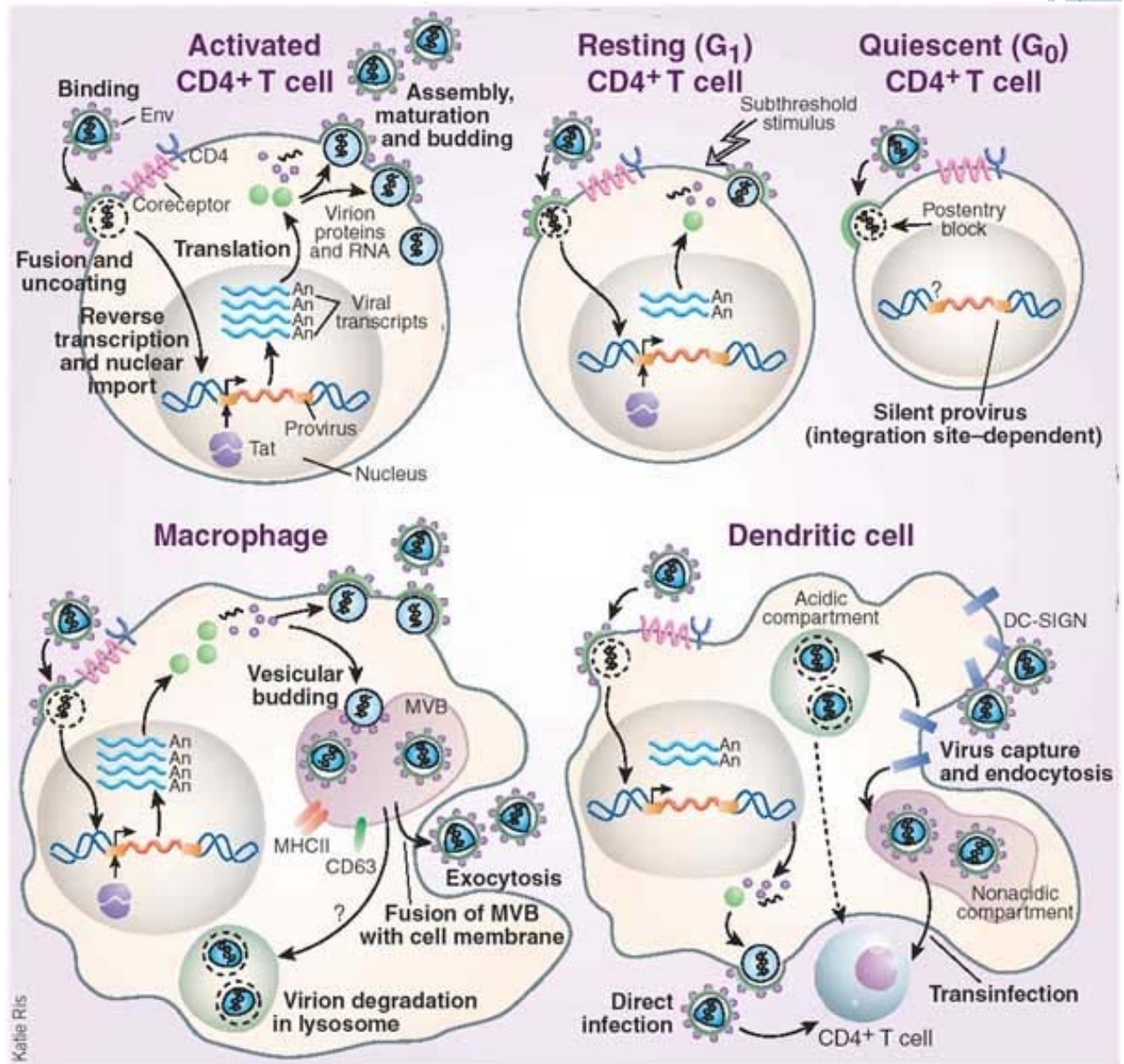


# FUSION INHIBITORS



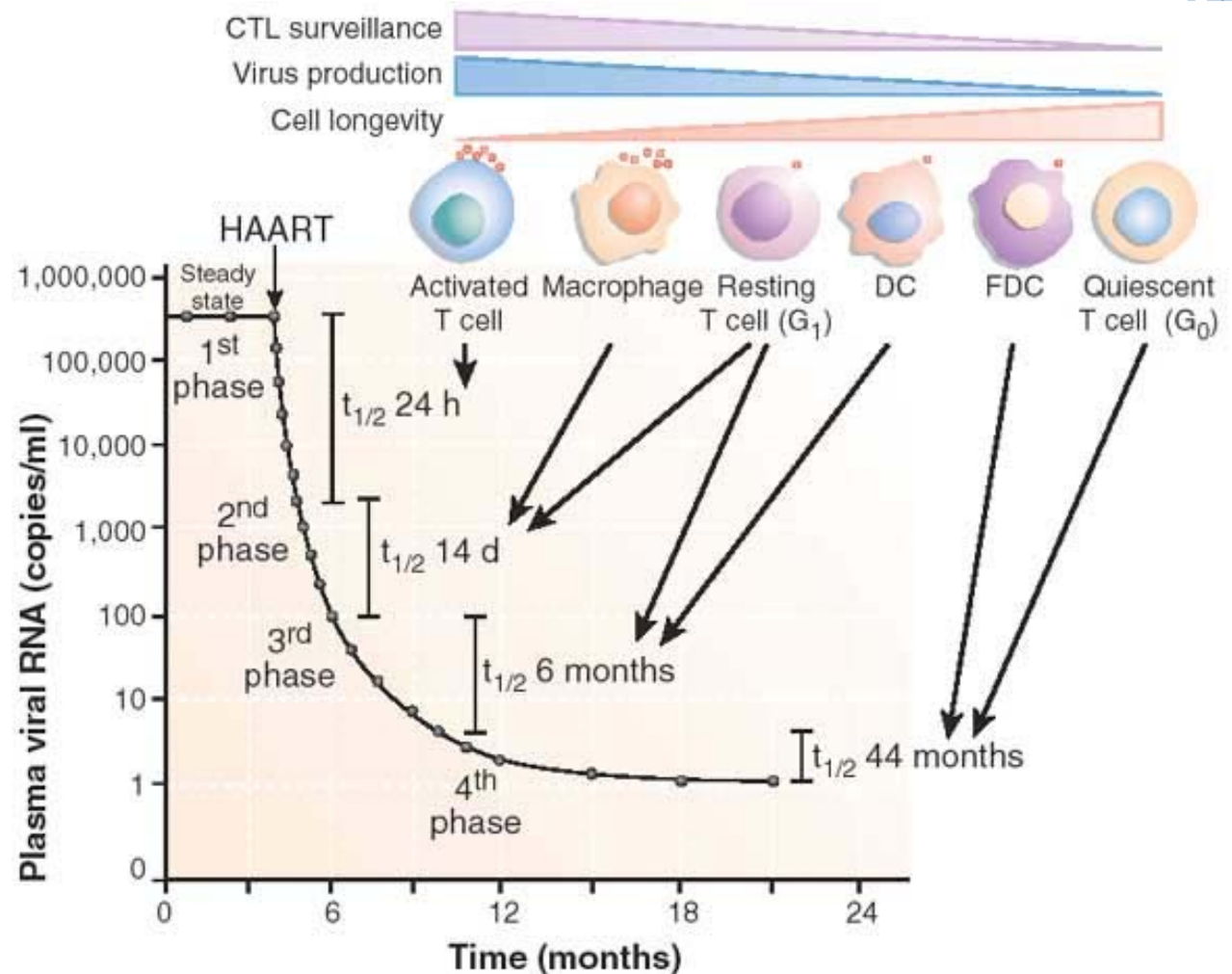
# LATENT RESERVOIRS OF VIRUS

- Multiple cell types can serve as latent reservoirs
- “Quiescence” of infected cells constrains the possibility total viral elimination

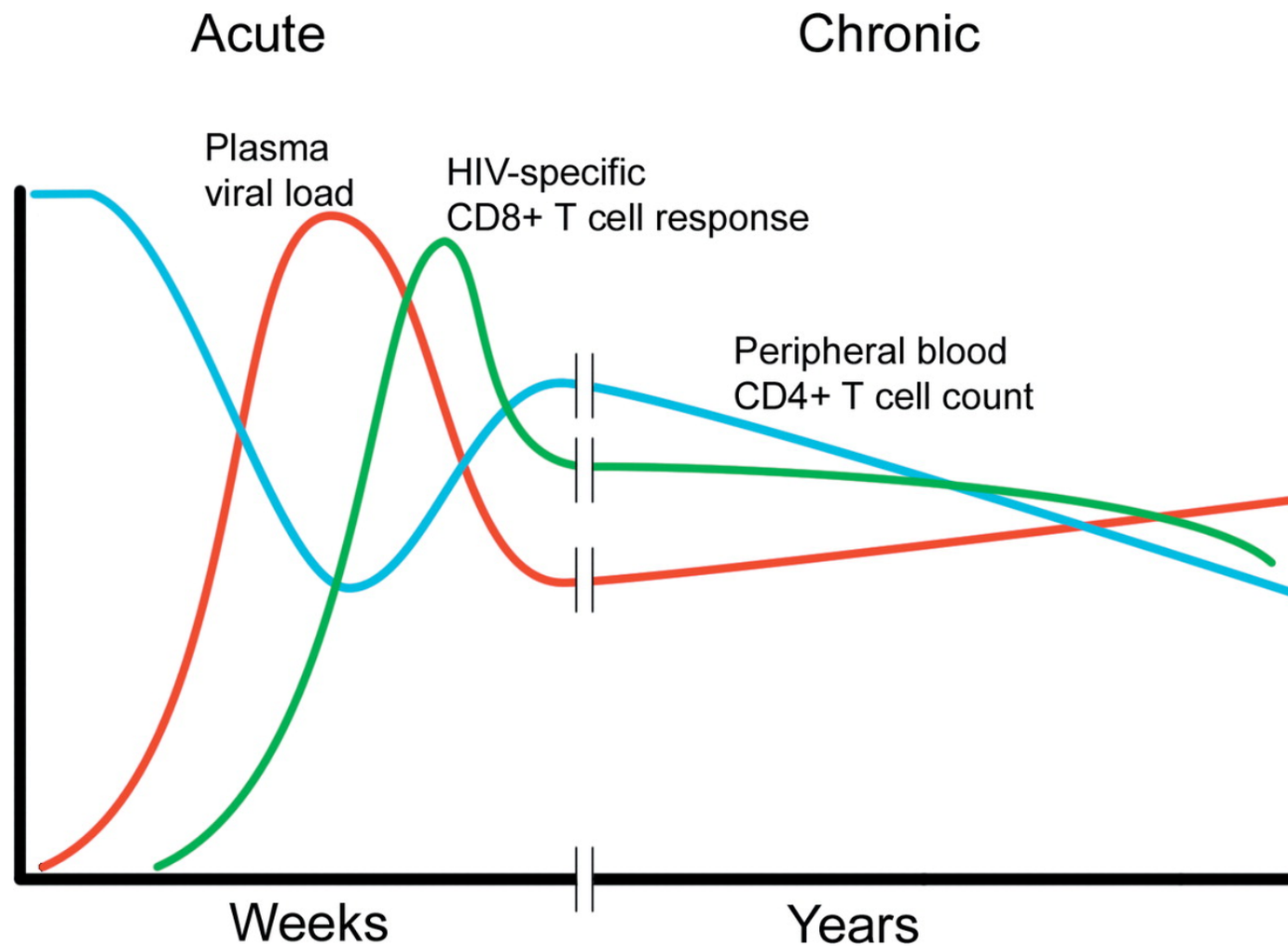


# CONTRIBUTION OF INDIVIDUAL RESERVOIRS

- Steady-state virus levels result from the relative contributions and turnover of each reservoir compartment
- After viral inhibition by HAART, plasma viral RNA decays in four distinct phases allowing a dissection of each reservoir's individual contribution



# CAN THE IMMUNE SYSTEM BE USED TO PREVENT OR CLEAR INFECTION?



# SUMMARY OF VACCINE TRIALS IN 2006

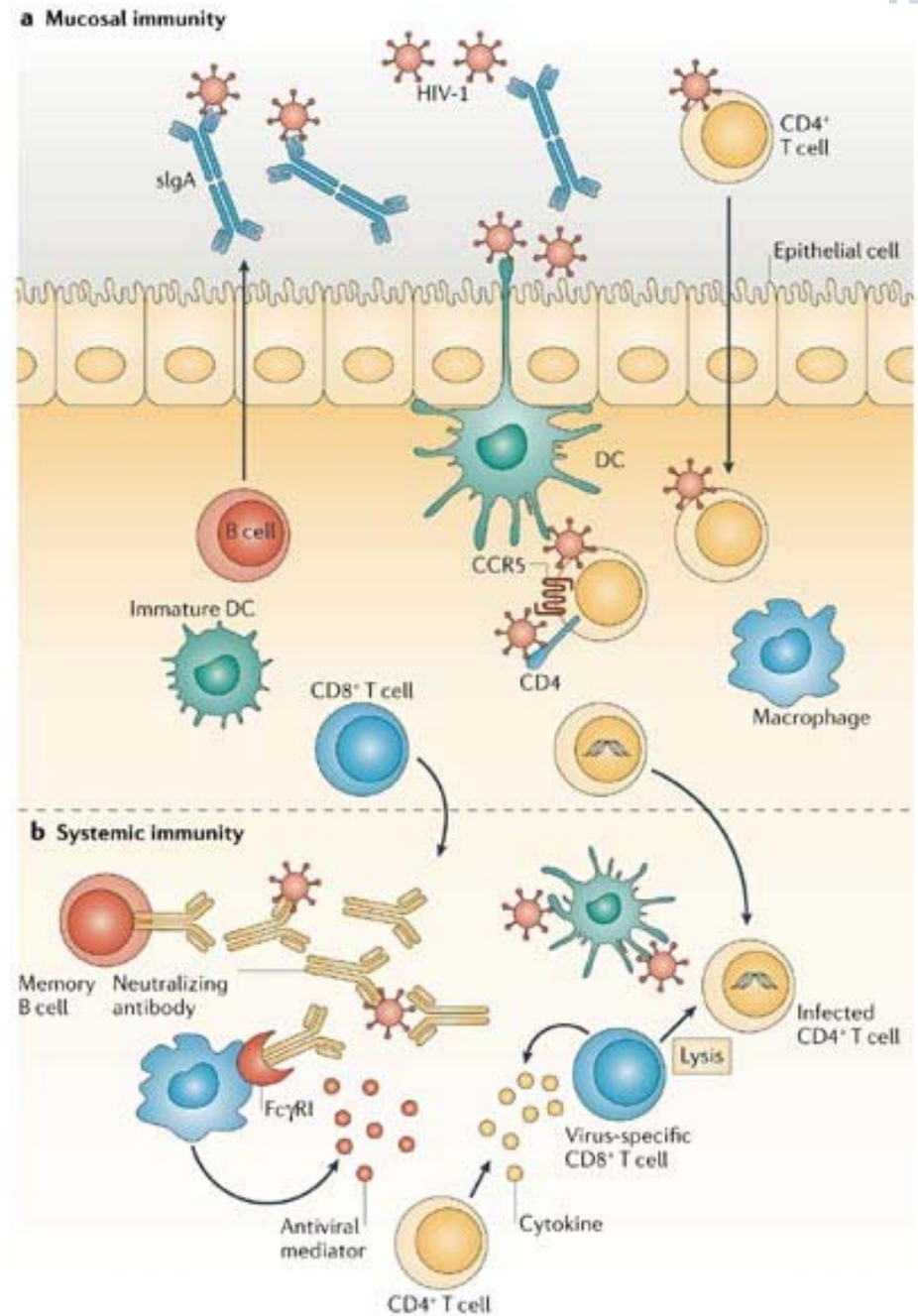
Vaccine candidate	Antigen (HIV-1 clade)	Manufacturer	Trial start date	Question being addressed
Prime with canarypox vector expressing HIV-1 genes	env (B, E), gag/pol (B)	Sanofi-Pasteur	October 2003	Will a gp120 protein vaccine that did not confer protection when used alone be useful in combination with a live, recombinant pox vector prime?
Boost with gp120 protein	gp120 (B, E)	Vaxgen		
Replication-defective adenovirus serotype 5 expressing HIV-1 genes	gag, pol, nef (B)	Merck	December 2004	Will an adenovirus-based vector vaccine confer a clinical benefit in individuals who become infected after vaccination?
Prime with plasmid DNA encoding HIV-1 genes	gag, pol, nef (B), env (A, B, C)	Vical, VRC	September 2005	Will a prime–boost strategy using DNA- and adenovirus-based vaccines encoding envelope proteins from three HIV-1 clades, as well as viral structural proteins, confer a benefit?
Boost with replication-defective adenovirus serotype 5 expressing HIV-1 genes	gag, pol (B), env (A, B, C)	GenVec, VRC		

Further information on [ongoing trials of preventative AIDS vaccines](#) can be found in the 2006 International AIDS Vaccine Initiative report. env, envelope; gag, group-specific antigen; gp120, glycoprotein 120; nef, negative factor; pol, polymerase; VRC, Vaccine Research Center, National Institutes of Health, Maryland, USA.

Letvin *Nature Reviews Immunology* 6, 930–939 (December 2006) | doi:10.1038/nri1959

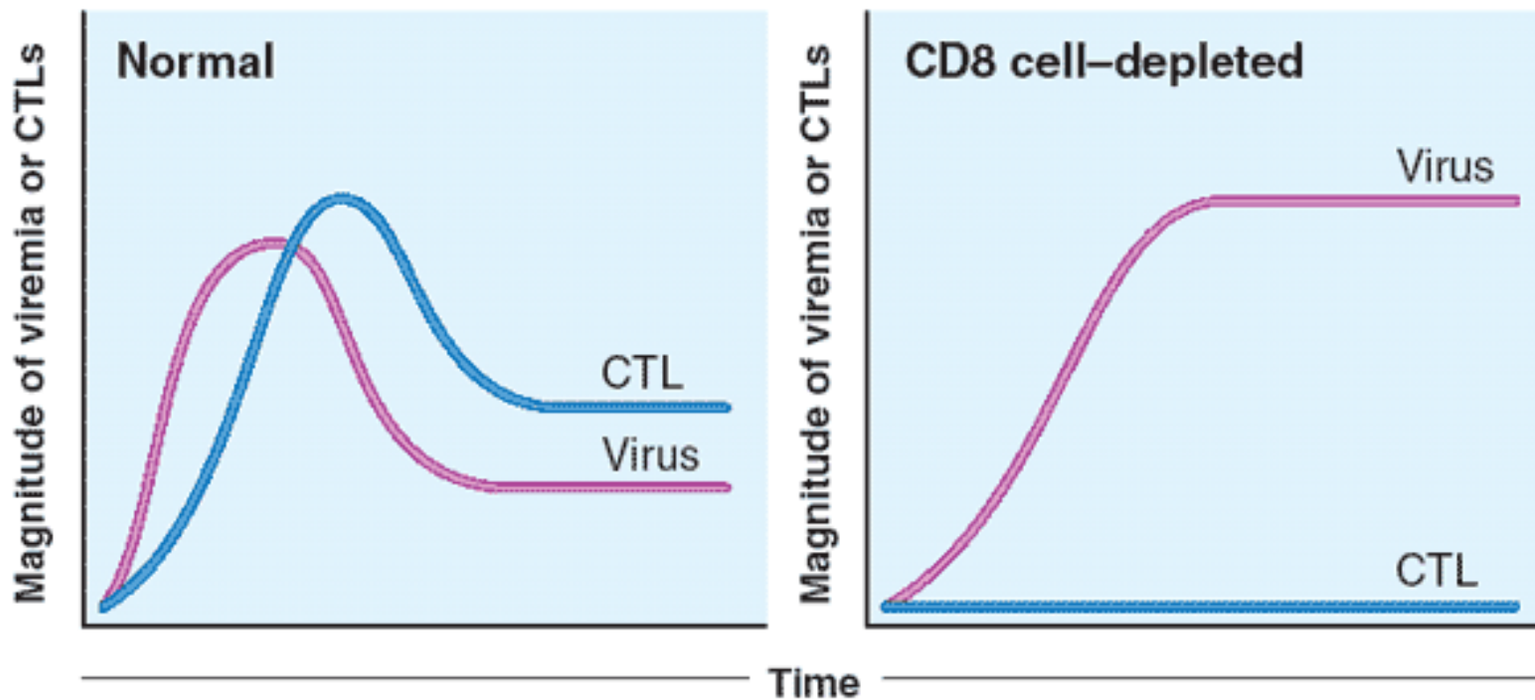
# MECHANISMS OF IMMUNE PROTECTION

- “Standard” immunological protection mechanisms, including antibody, clearance by phagocytic cells and Fc receptors, and cytotoxic killing of infected cells all function to limit infection and control long-term viral loads
- The loss of effective immune control is what leads to the development of AIDS, therefore the immune response in principle is an effective tool for viral control and clearance



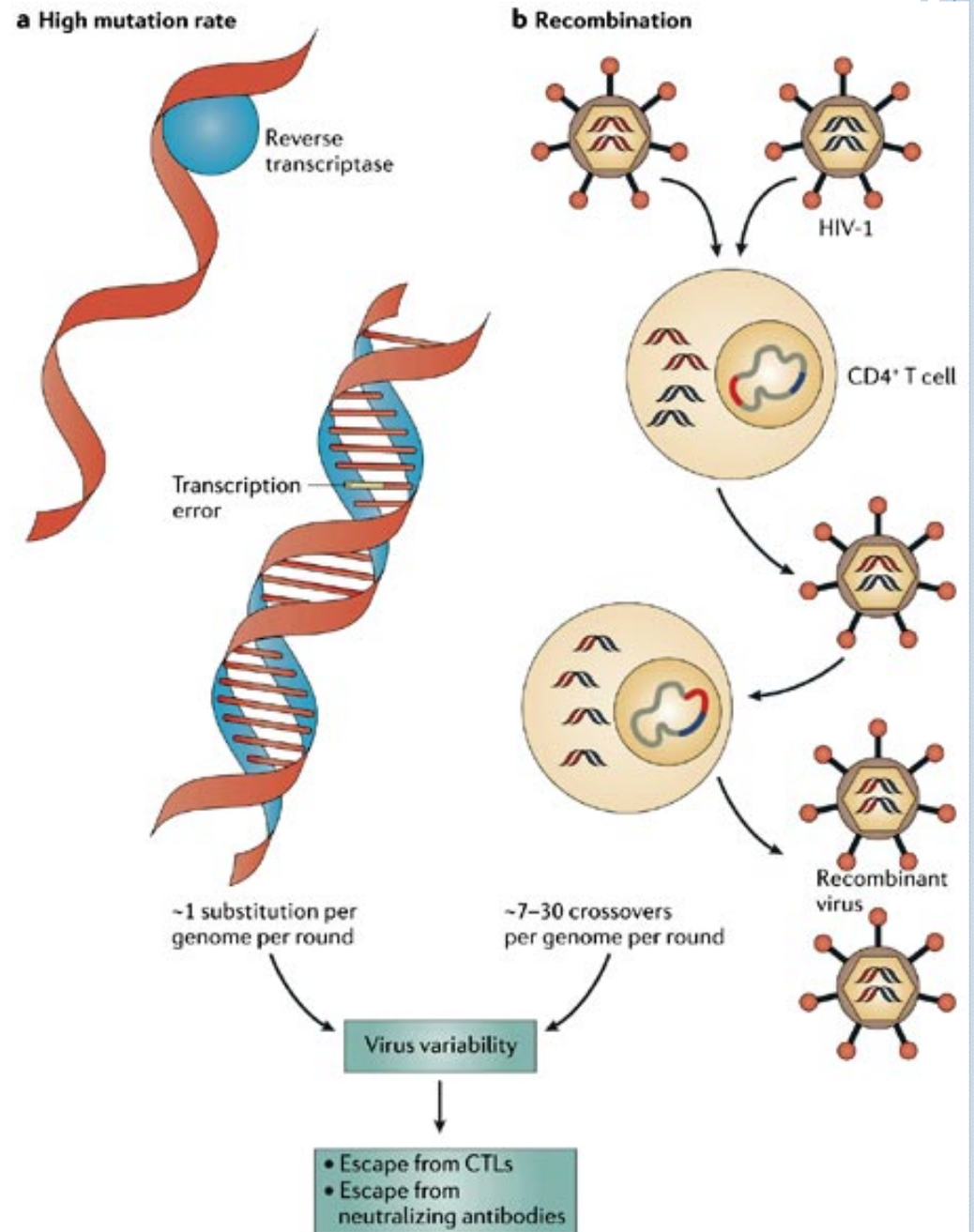
## CD8 T CELLS PROVIDE SIGNIFICANT VIRAL CONTROL DURING THE CHRONIC PHASE OF INFECTION

- CD8 depletion in SIV-infected animals leads to rapid increase in viral titers and pathogenesis of disease

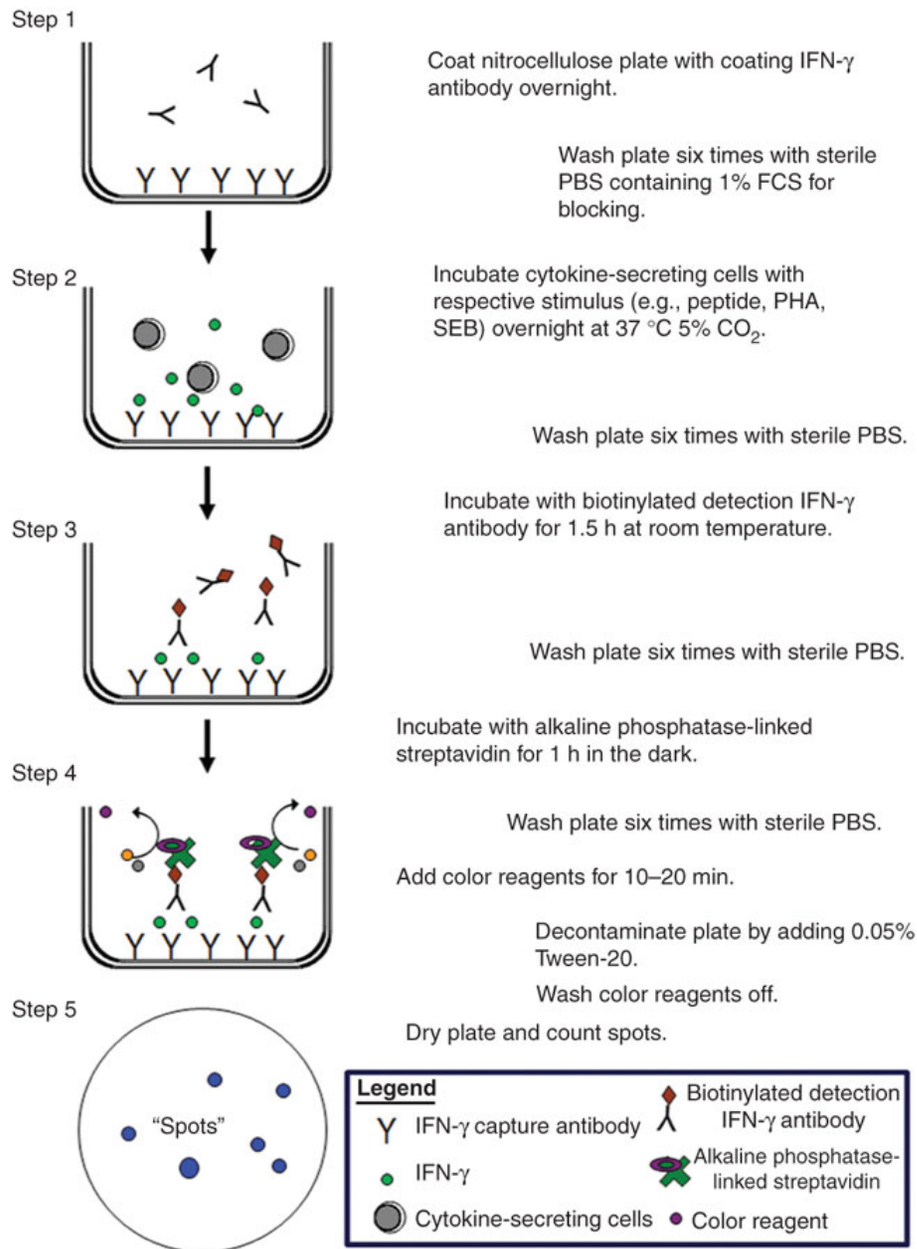


# VIRAL IMMUNE ESCAPE MECHANISMS

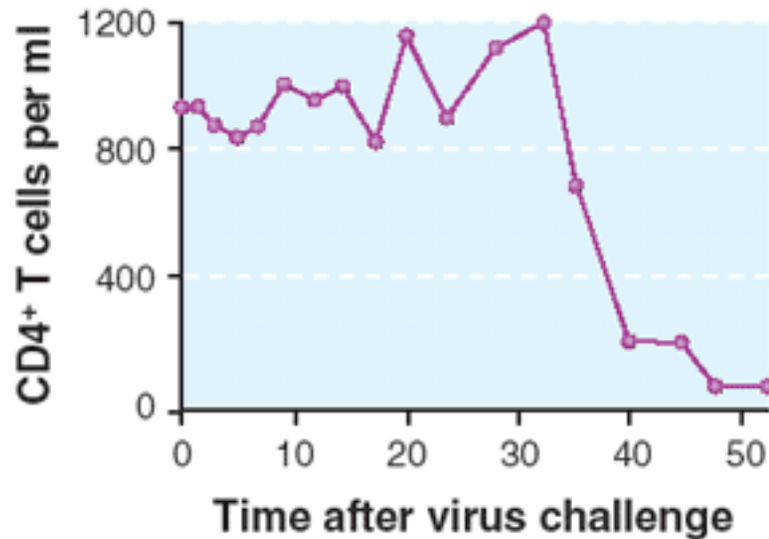
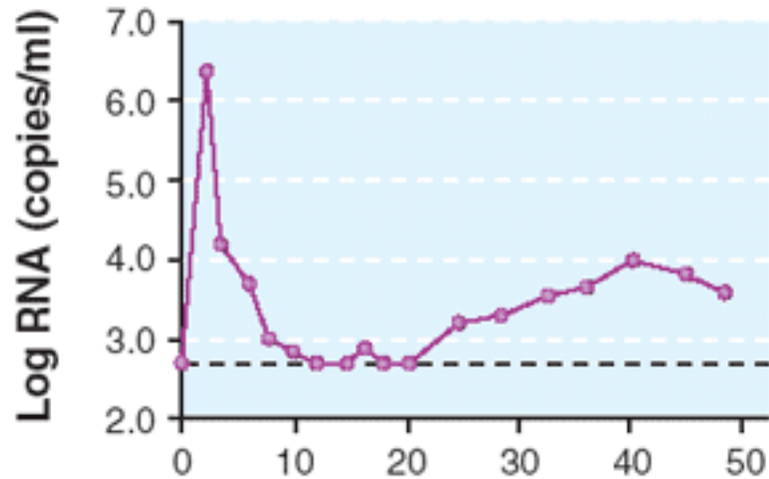
- “Antigenic drift” from the very high rate of mutation of the RT enzyme allows rapid escape from individual antibody and CTL responses
- Epitopes are constrained by structural/functional requirements



# HOW DO WE ASSAY FOR T CELL RESPONSES IN HIV INFECTED INDIVIDUALS?



# IMMUNODOMINANT EPITOPE ESCAPE CAN LEAD TO LOSS OF VIRAL CONTROL



## Gag p11C (181–189) sequences

	C	T	P	Y	D	I	N	Q	M	
Week 0	-	-	-	-	-	-	-	-	-	(15/15)
Week 14	-	-	-	-	-	-	-	-	-	(8/8)
Week 20	-	I	-	-	-	-	-	-	-	(10/10)
Week 24	-	I	-	-	-	-	-	-	-	(11/11)
Week 28	-	I	-	-	-	-	-	-	-	(11/11)
Week 36	-	I	-	-	-	-	-	-	-	(11/11)
Week 44	-	I	-	-	-	-	-	-	-	(10/10)

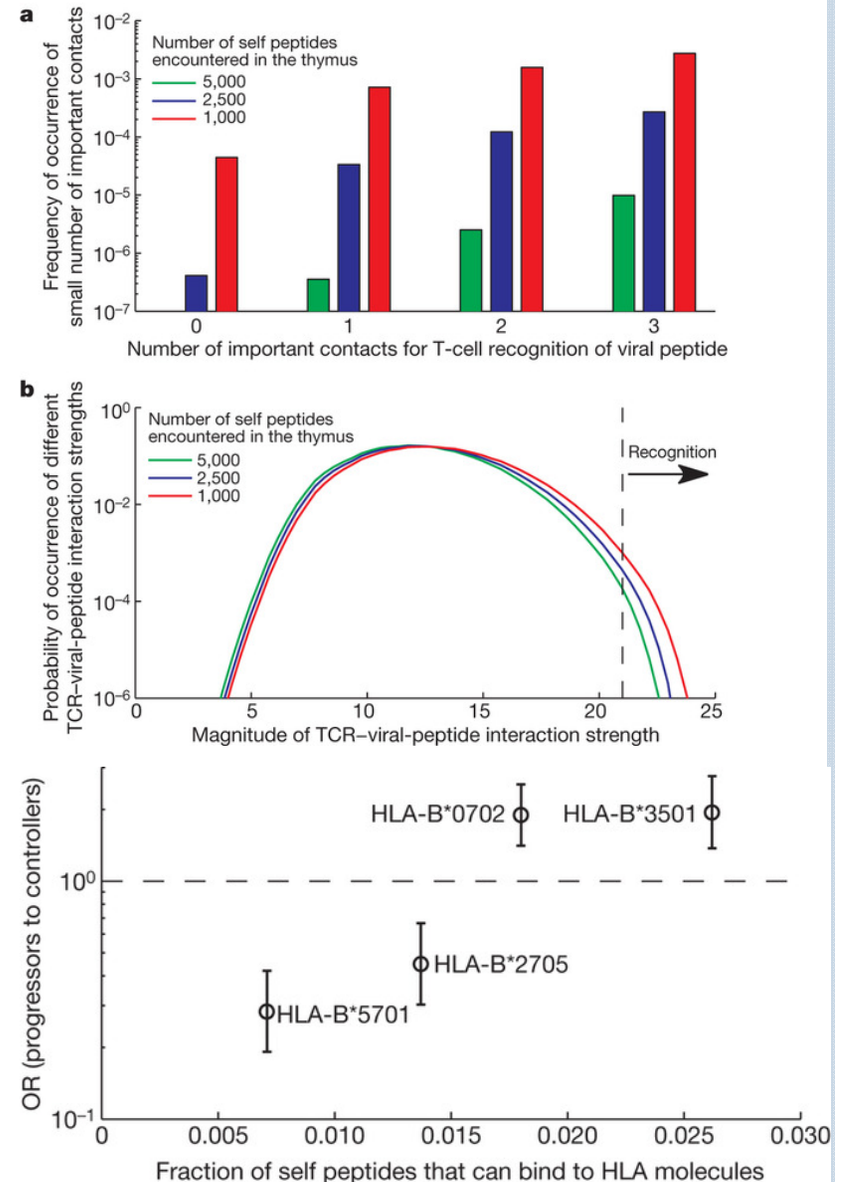


# RECENT REPORTS RELATING MHC HAPLOTYPE TO HIV CONTROL

Nature 465, 350–354 (20 May 2010) Effects of thymic selection of the T-cell repertoire on HLA class I-associated control of HIV infection

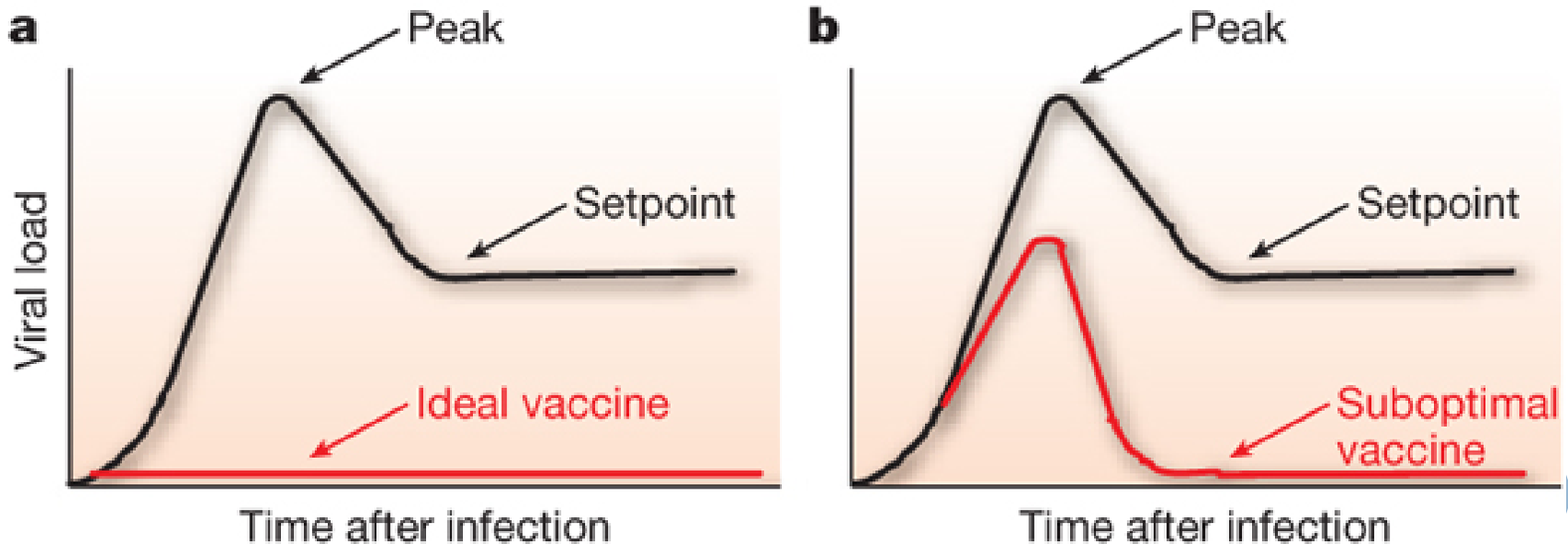
Andrej Košmrlj, Elizabeth L. Read, Ying Qi, Todd M. Allen, Marcus Altfeld, Steven G. Deeks, Florencia Pereyra, Mary Carrington, Bruce D. Walker & Arup K. Chakraborty

- Relating the breadth of the TCR repertoire (how many different T cell receptors does the body make?) to the MHC haplotype (the more self peptides available for negative selection, the narrower (and less “cross-reactive” the TCR repertoire)
- Less cross-reactive TCR repertoires are then associated with poor control



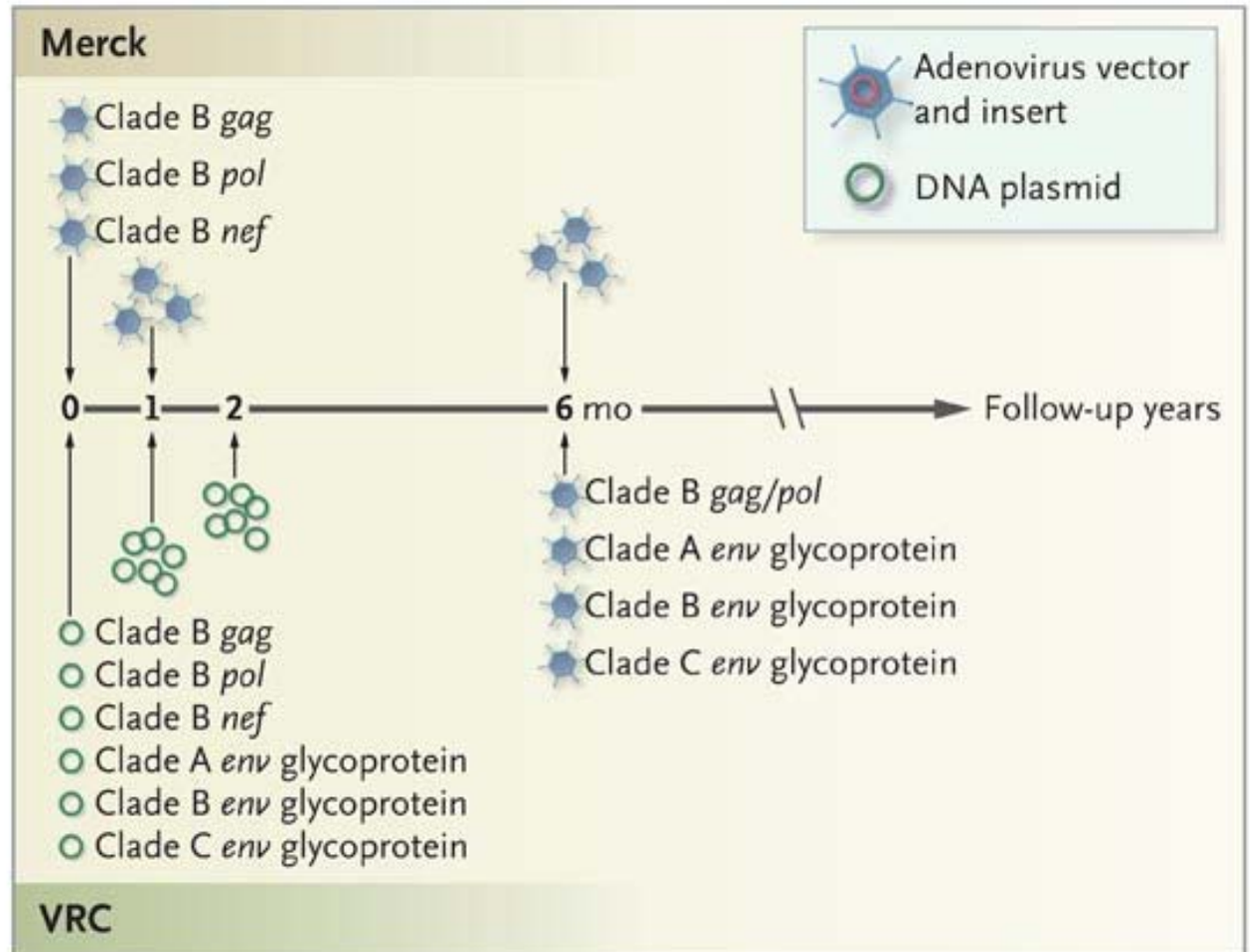
## VACCINE EXPECTATIONS

- Since viral load “set point” is a key predictor of disease progression and pathogenesis, even a suboptimal vaccine could be of use in highly endemic areas to protect against disease and spread (we’ll talk more about this when we get to malaria)



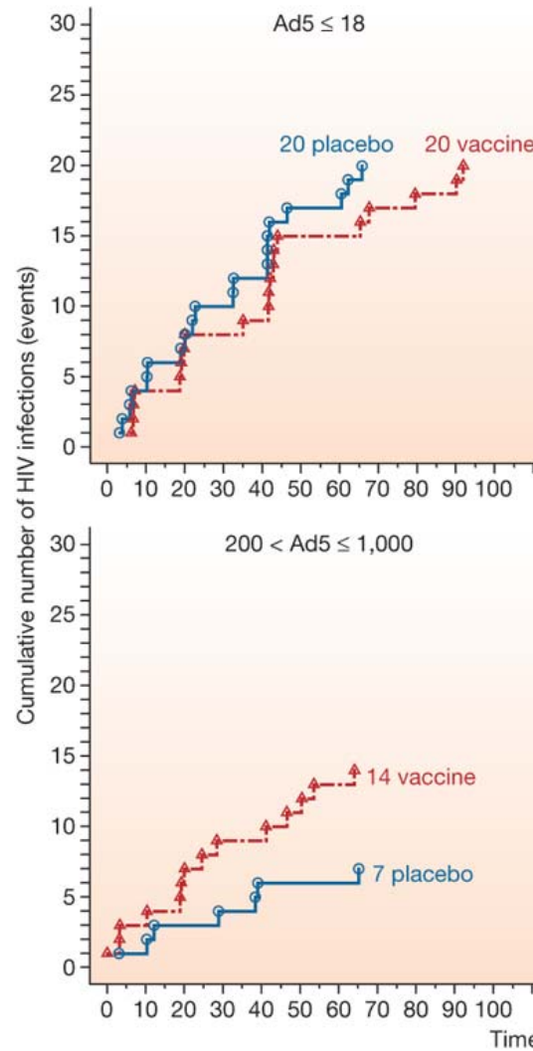
# THE MERCK VACCINE

- Use of a viral vector has been shown experimentally to boost cellular responses, by delivering more antigen with the proper innate/PAMP signals



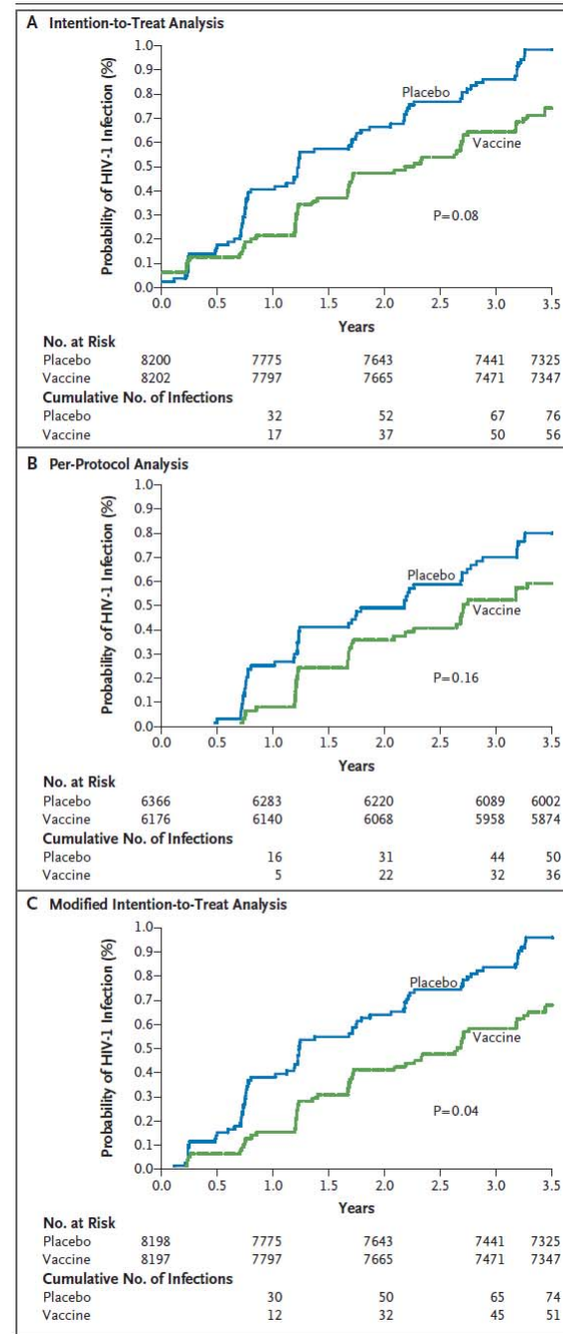
# MERCK VACCINE FAILURE

- Not only did the Merck Vaccine fail to protect, there appeared to be an enhancement of infection in vaccinees who had relatively higher pre-existing antibody titers to the viral vector
- This failure led to the cancellation of other vaccine trials based on a similar approach
- HVTN-505 just halted in April 2013—also Ad5 based (41 vacc inf, 30 placebo)

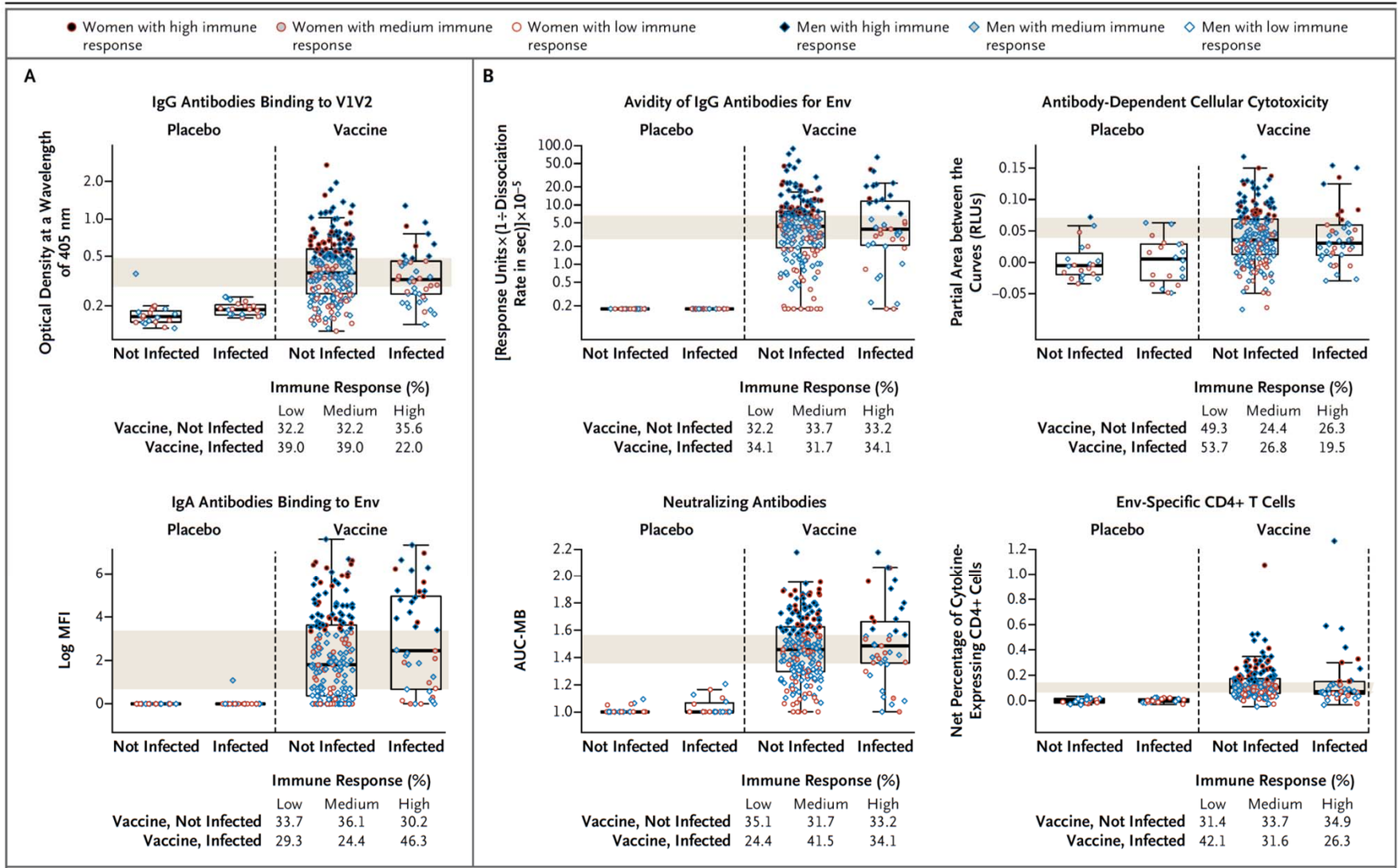


# A PROTECTIVE VACCINE? RV144 TRIAL

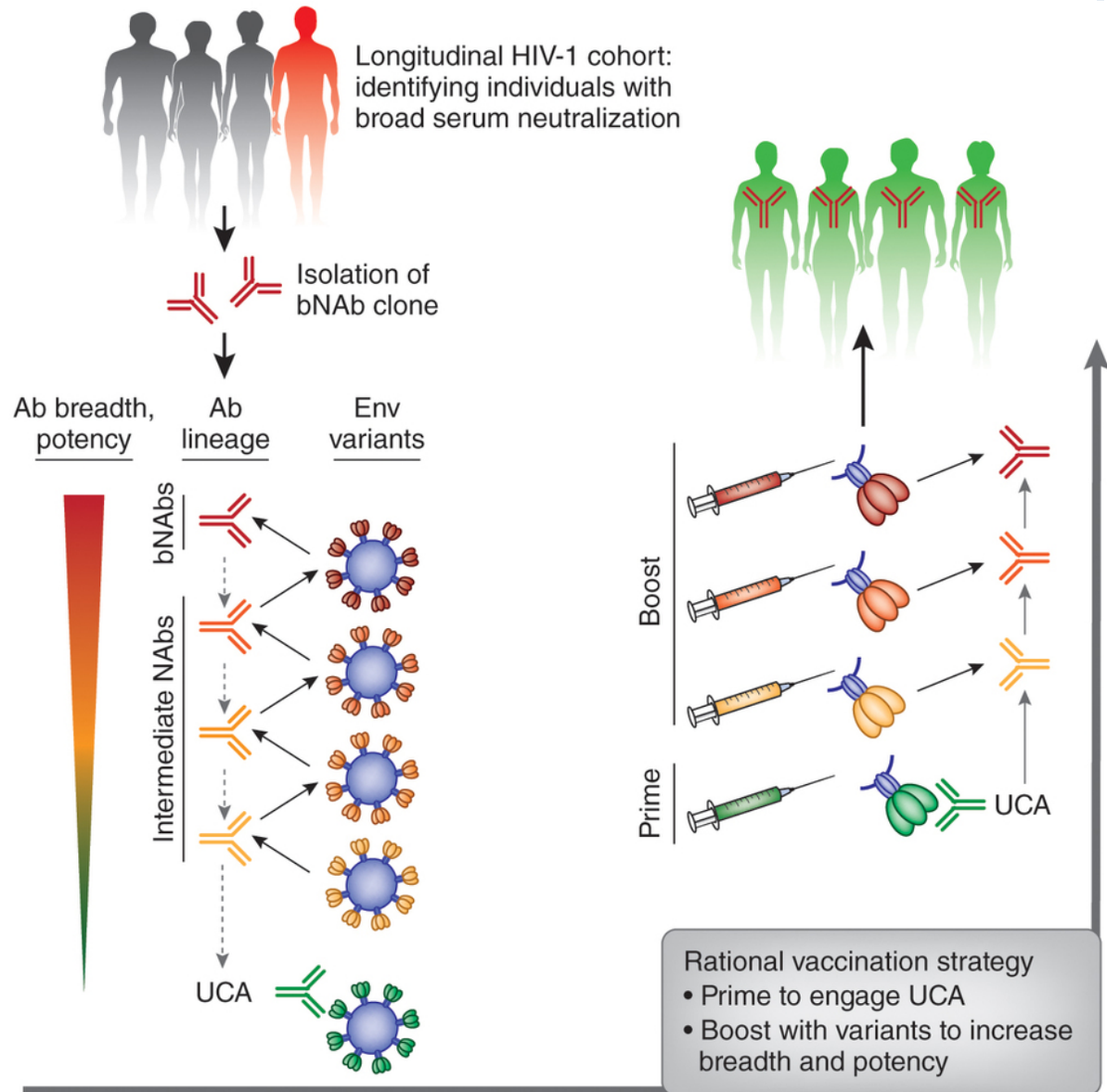
- ALVAC/AIDSVAX Prime boost-boost vaccine (canarypox followed by protein boost, gp120 based)
- 16,402 vaccinees
- Vaccine efficacy was 31.2%
- No mitigation of viral load in those that did become infected



# IMMUNE CORRELATES OF HIV RISK



# BROADLY NEUTRALIZING ANTIBODY APPROACHES FOR HIV VACCINE



## POINTS FOR DISCUSSION

- HIV is a unique pathogen in that it targets the immune system directly—playing “offense”—killing or dysregulating the cells that specifically target it and “defense”, employing more conventional immune escape mechanisms
- Despite this, the immune response, both antibody and CTLs, provide an important level of control over the virus for an extended period of time, keeping the reservoir relatively stable
- Vaccines could in principle employ similar strategies, but drugs are still the most effective treatment tool

