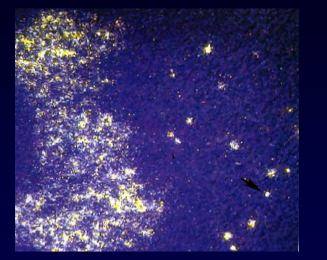


HIV Integration

Primary HIV Infection: Pathogenetic Steps

- Virus dendritic cell interaction
 - Infection is typically with R5 (M-tropic) strains
 - Importance of DC-SIGN
- Delivery of virus to lymph nodes
- Active replication in lymphoid tissue
- High levels of viremia and dissemination
- Downregulation of virus replication by immune response
- Viral set point reached after approximately 6 months

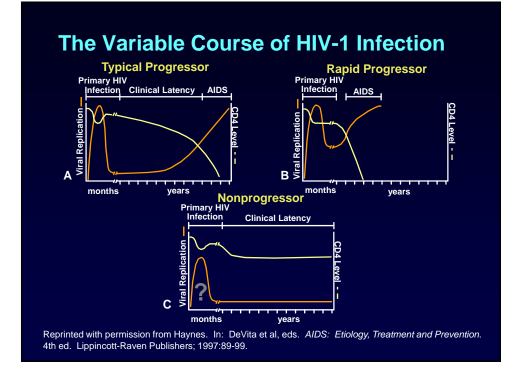
PHI: Early Seeding of Lymphoid Tissue



Schacker T et al: J Infect Dis 2000;181:354-357

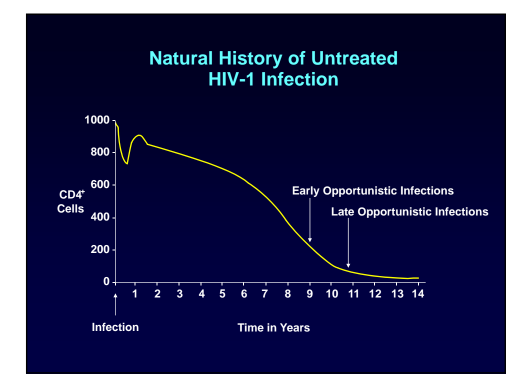
Primary HIV Infection: Clinical Characteristics

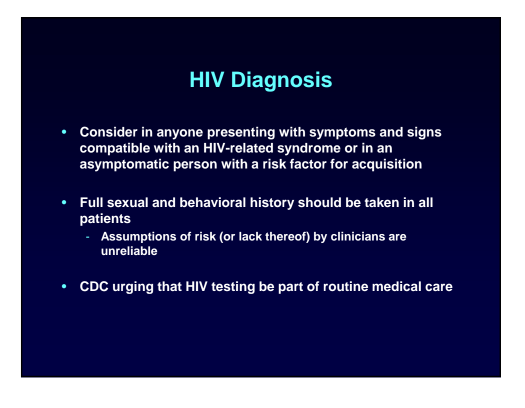
- 50-90% of infections are symptomatic
- Symptoms generally occur 5-30 days after exposure
- Symptoms and signs
 - Fever, fatigue, myalgias, arthralgias, headache, nausea, vomiting, diarrhea
 - Adenopathy, pharyngitis, rash, weight loss, mucocutaneous ulcerations, aseptic meningitis, occas. oral/vaginal candidiasis
 - Leukopenia, thrombocytopenia, elevated liver enzymes
- Median duration of symptoms: 14 days



Primary HIV Infection: Determinants of Outcome

- Severity of symptoms
- Viral strain
 - SI (X4) vs. NSI (R5) viruses
- Importance of GI tract associated lymphoid tissue (GALT)
- Immune response
 - CTL response
 - Non-CTL CD8 responses
 - Humoral responses?
- Viral set point at 6-24 months post-infection
- Other host factors
 - Chemokine receptor and HLA genotype
- Gender and differences in viral diversity?
- Antiviral therapy
 - Near vs. long-term benefit?





Laboratory Diagnosis of Established HIV Infection: Antibody Detection

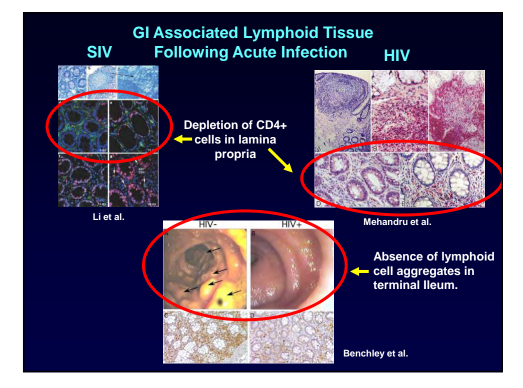
- Screening
 - Serum ELISA
 - Rapid blood or salivary Ab tests
- Confirmation
 - Western blot
 - In some settings, confirmation of one rapid test is done by performing a second, different rapid test
- Written consent for HIV Ab testing must be obtained and be accompanied by pre- and post-test counselling
 - Consent process may change to make it simpler and easier but proper counselling remains crucial

Laboratory Diagnosis of Acute HIV-1 Infection

- Patients with acute HIV infection may present to a health care facility before full antibody seroconversion
 - ELISA may be negative
 - ELISA may be positive with negative or indeterminate Western blot
- Plasma HIV-1 RNA level should be done if acute HIV infection is suspected
- Follow-up antibody testing should be performed to document full seroconversion (positive ELISA and WB)

Established HIV Infection: Pathogenesis

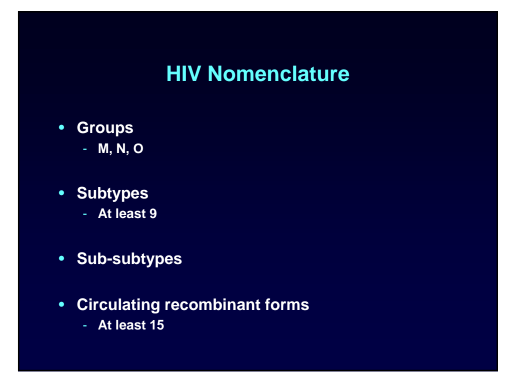
- Active viral replication present throughout course of disease
- Major reservoirs of infection exist outside of blood compartment
 - Lymphoreticular tissues
 - » Gastrointestinal tract (GALT)
 - Central nervous system
 - Genital tract
- Virus exists as multiple quasispecies
 - Mixtures of viruses with differential phenotypic and genotypic characteristics may coexist
- At least 10 X 10⁹ virions produced and destroyed each day
- T_{1/2} of HIV in plasma is <6 h and may be as short as 30 minutes
- Immune response, chemokine receptor status and HLA type are important codeterminants of outcome

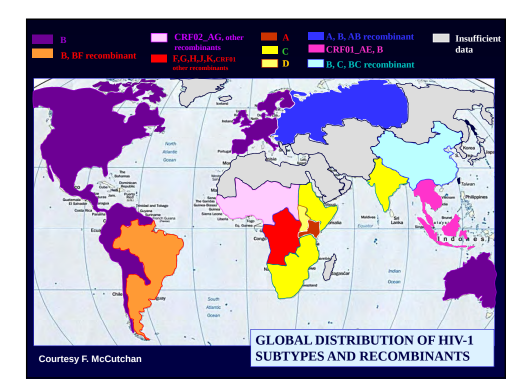


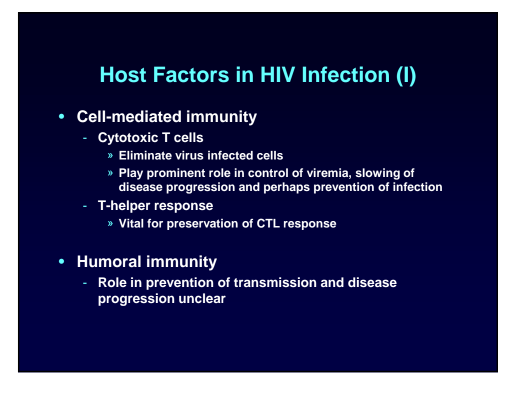
Determinants of Outcome: Selected Viral Factors

Escape from immune response

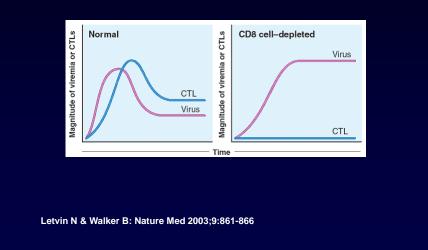
- Under immune selective pressure (cellular and humoral), mutations in *gag*, *pol* and *env* may arise
- Attenuation
 - *nef* deleted viruses associated with slow or long-term nonprogression in case reports and small cohorts
- Tropism
 - R5 to X4 virus conversion associated with increased viral pathogenicity and disease progression
- Subtypes
 - Potential for differential risks of heterosexual spread or rates of disease progression

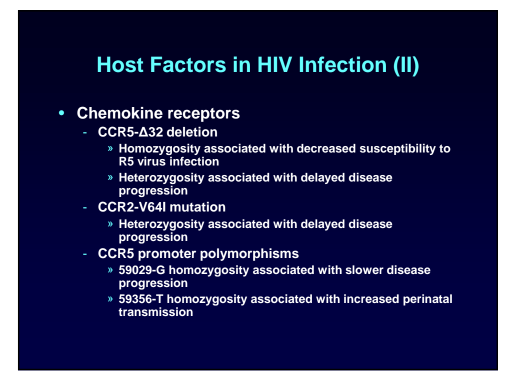






Role of CTL's in Control of Viremia





Host Factors in HIV Infection (III)

• Other genetic factors

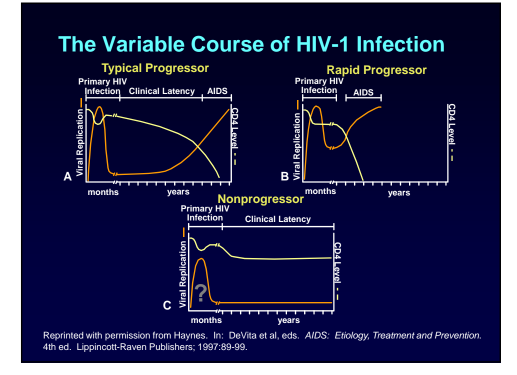
- Class I alleles B35 and C ω 4
 - » Associated with accelerated disease progression
- Heterozygosity at all HLA class I loci » Appear to be protective
- HLA-B57, HLA-B27, HLA-Bω4, HLA-B*5701
 - » Associated with long-term non-progression
- HLA-B14 and HLA-C8
 - » ?Associated with long-term nonprogression

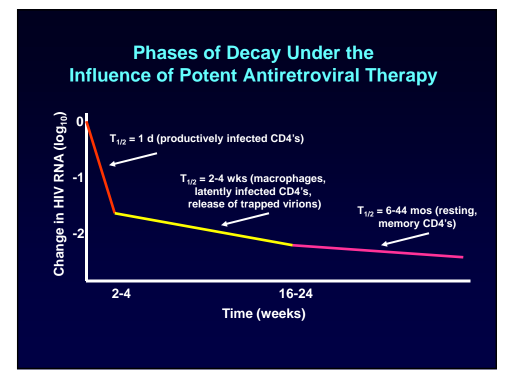
Mechanisms of CD4+ Cell Death in HIV Infection

- HIV-infected cells
 - Direct cytotoxic effect of HIV
 - Lysis by CTL's
 - Apoptosis
 - » Potentiated by viral gp120, Tat, Nef, Vpu

• HIV-uninfected cells

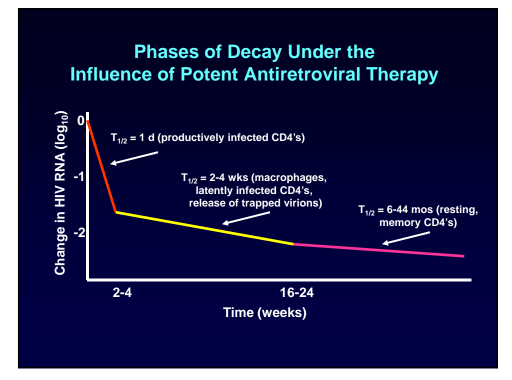
- Apoptosis
 - » Release of gp120, Tat, Nef, Vpu by neighboring, infected cells
- Activation induced cell death

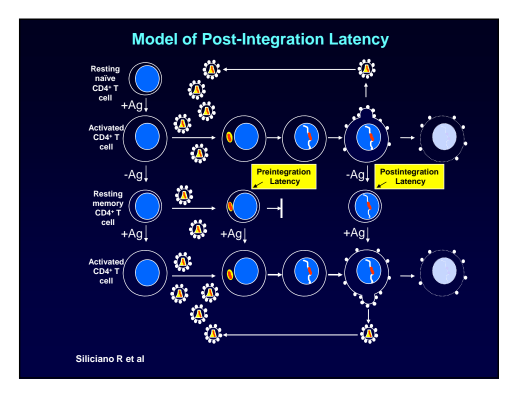




Therapeutic Implications of First and Second Phase HIV RNA Declines

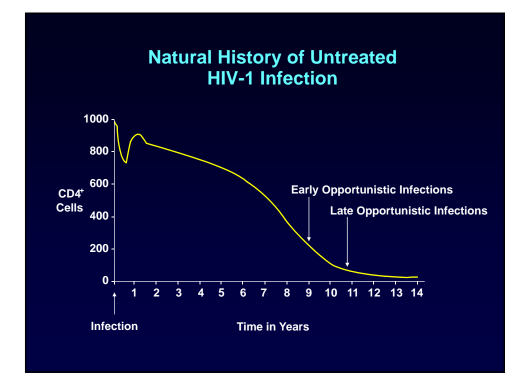
- Antiviral potency can be assessed in first 7-14 days
 Should see 1-2 log declines after initiation of therapy in
 - persons with drug susceptible virus who are adherent
- HIV RNA trajectory in first 1-8 weeks can be predictive of subsequent response
 - Durability of response translates into clinical benefit

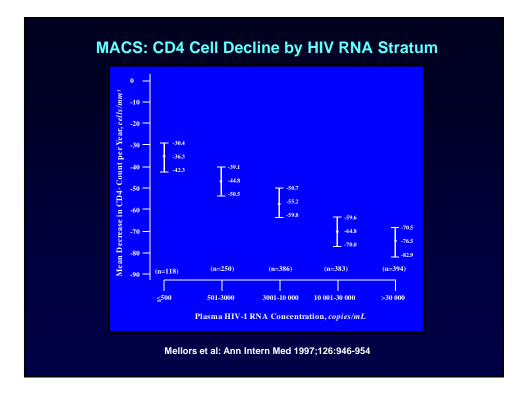




Therapeutic Implications of Third Phase of HIV RNA Decay: Latent Cell Reservoir

- Viral eradication not possible with current drugs
- Archive of replication competent virus history is established
 - Drug susceptible and resistant
- Despite the presence of reservoir(s), minimal degree of viral evolution observed in patients with plasma HIV RNA levels <50 c/ml suggests that current approach designed to achieve maximum virus suppression is appropriate





CD4 and HIV-1 RNA (I)

- Independent predictors of outcome in most studies
- Near-term risk defined by CD4
- Longer-term risk defined by both CD4 and HIV-1 RNA
- Rate of CD4 decline linked to HIV RNA level in untreated persons



- Good but incomplete surrogate markers
 - For both natural history and treatment effect
- Thresholds are arbitrary
 - Disease process is a biologic continuum
 - Gender specificity of HIV RNA in early-mid stage disease needs to be considered
- Treatment decisions should be individualized
 - Baseline should be established
 - Trajectory determined

"Non-AIDS" Conditions

- Since 2006, a number of "non-AIDS" conditions have been described to be associated with uncontrolled HIV-1 viremia, even in persons with relatively well preserved CD4 cell counts (e.g., >350/mm³)
 - Cardiovascular events
 - Hepatic disease
 - Renal disease
 - Malignancies
- Direct effect of HIV-1 on organ systems, associated immune activation and/or other mechanisms may be involved
- Active area of investigation
- Redefining HIV-related disease progression and influencing decision of when to start ART

