Biomedical Engineering for Global Health

Lecture 10
HIV/AIDS vaccine development

Review of lecture 9

How do vaccines work?

Types of Vaccines:

Review of lecture 9

Are vaccines effective?

-Edward Jenner's experiment

-Name big success example: _____

How are vaccines tested?

Review of lecture 9

What are some challenges of vaccine development?

- -Developed countries
- -Developing countries

The big three:

, ,

Review of lecture 9

How do vaccines work?

Types of Vaccines:

- Non infectious: Inactivated, subunit & toxoid
- Live-attenuated
- Carrier
- DNA

Vaccine effectiveness

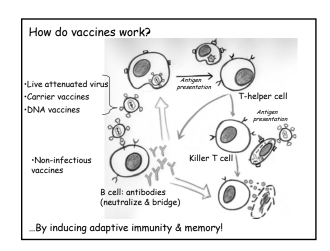
-From Edward Jenner to Smallpox erradication

Vaccine Safety:

-Clinical trials/VAERS

Challenges of vaccine development

- -Developed vs. developing world
- -The big three: <u>TB</u>, <u>Malaria</u>, <u>HIV</u>



Lecture map

HIV-1/AIDS

- History of epidemic
- The HIV-1 virus
- Clinical course of infection

The HIV vaccine

- History of HIV vaccines
- Challenges for vaccine development
- Types of vaccines
 - -VaxGen's qp120
 - -Sanofi Pasteur ALVAC: prime/boost strategy
 - Merk Ad5

Discussion:

- Specter article

Lecture map

HIV-1 /AIDS

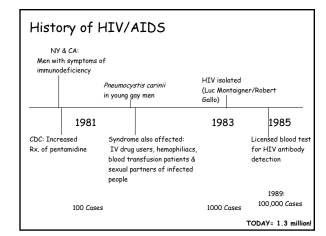
- History of epidemic
- The HIV-1 virus
- Clinical course of infection

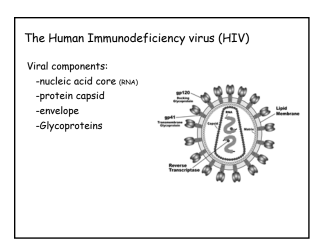
The HIV vaccine

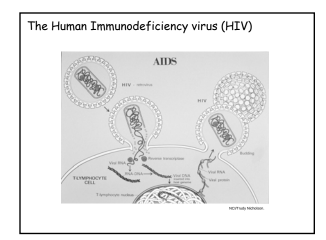
- History of HIV vaccines
- Challenges for vaccine development
- Types of vaccines
 - -VaxGen's qp120
 - -Sanofi Pasteur ALVAC: prime/boost strategy
 - Merk Ad5

Discussion:

- Specter article







Clinical course of HIV/AIDS

Acute: Infection of CD4+ cells (T-helper cells),

50% of memory cells lost! Loss of defense repertoire! High viral load

Symptoms 2-8wks: fever, pharyngitis malaise, weight loss

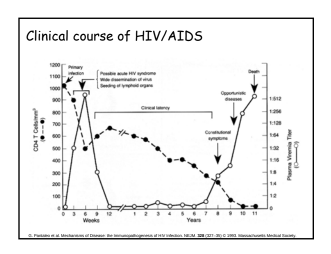
Chronic: Decreased CD4+ cells cannot support rate of replication
Innate and adaptive immune responses control expansion
Integrated provirus acts as latent virus reservoir:

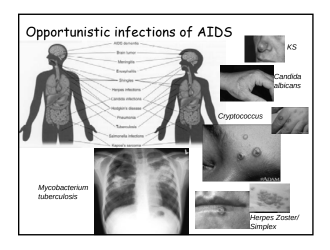
- no viral synthesis

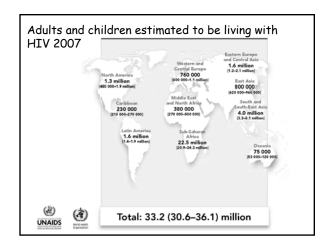
- reservoir <u>protected from antivirals and immune attack</u>
Mostly asymptomatic: fatigue & lymphoadenopathy

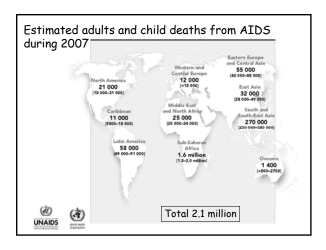
AIDS: Progressive loss of CD4+ (T helper) cells

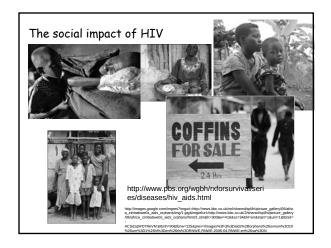
= profound defect on cellular immunity increased viral load & opportunistic infections and cancer











Lecture map HIV-1/AIDS - History of epidemic - The HIV-1 virus - Clinical course of infection The HIV vaccine - History of HIV vaccines - Challenges for vaccine development - Types of vaccines - VaxGen's gp120 - Sanofi Pasteur ALVAC: prime/boost strategy - Merk Ad5 Discussion: - Specter article

History of HIV vaccines

- 1984:
 - Robert Gallo discovers virus that causes HIV
 - Margaret Heckler, Secretary of HEW, predicts we will have vaccine within 2 years
- · 1997:
 - President Clinton declares, "an HIV vaccine will be developed in a decade's time."
- · 2003
 - President Bush asks congress to appropriate \$15B to combat the spread of HIV in Africa and the Caribbean
- · Today: Where is the vaccine?

Challenges of HIV vaccine

- 1. Many forms of HIV
 - HIV-1: Many subtypes: 9 clades
 - HIV-2 Western Africa
- 2. Each sub-type may require different vaccine
- 3. HIV mutates rapidly: error-prone reverse transcriptase
- Surface glycoproteins not readily available for antibodies:
 - Coated in sugary molecules: N-linked glycans
 - · Change shape after attachment step
- 5. HIV infects, suppresses and destroys key cells of the immune system

Design Goals for HIV Vaccine

- · Must produce both:
 - Antibody mediated immunity (B cells)
 - Immune system must see virus or viral debris
 - Cell mediated immunity (killer T cells)
 - HIV viral proteins must be presented to immune system on $\ensuremath{\mathsf{MHC}}$ receptors

Types of Vaccine

- · Non-infectious vaccines
 - Stimulate B-cells
 - Killed virus
 - Subunit
 - Toxoid
- · Live attenuated vaccines
 - Stimulate both B-cells and killer T-cells
- · Carrier vaccines
 - Stimulate both B-cells and killer T-cells
- · DNA vaccines:
 - Stimulate both B-cells and T-cells

Methods tried for HIV vaccine development

Table 1 Methods used for the development of currently licensed vaccines and their failure to yield an HIV/AIDS vaccine

Type of vaccine

Examples

Convolute for

Problem for HIV vaccine development

Too risky—he attenuated Malesia

Manura

Manura

Walcella

Whole

Inactivated polio

Inactivated polio

Inactivated polio

Inactivated polio

Inactivated polio

Inactivated polio

Inactivated Hepetoto. A notibody

Inactivated and poeter subunit vaccines porteet primarily by eliciting

artibiodies. Both these approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for

HIV/AIDS. In the approaches have failed to elicit protective artibiodies for hiv/AIDS. In the approaches have failed to elicit protective artibiodies for hiv/AIDS. In the approaches have failed to elicit protective artibiodies for hiv/AIDS. In the approaches have failed to elicit protective artibiodies for hiv/AIDS. In the approaches have failed to elicit protective artibiodies for hiv/AIDS. In the approaches have failed to elicit protective artibiodies for hiv/AIDS. In the approaches have failed to elicit protective artibiodies for hiv/AIDS. In the approaches have failed to elicit protective artibiodies for hiv/AIDS. In the approaches have failed to elicit protective artibiodies for hiv

(From Robinson H.L., Clin. Pharmacol. Ther. 2007, 82: 686-693)

Live attenuated viral vaccine

- · Most likely to stimulate necessary immune response
- · Too dangerous!
 - Virus mutates constantly
 - If it undergoes mutation that restores its strength, would be devastating
- · Monkey experiments:
 - <u>All</u> vaccinated animals developed AIDS and died (although more slowly than those infected with unaltered virus)

Non infectious vaccines

- Whole virus: May not inactivate all virus

Animal studies:

Stimulates Ab which block a small # of HIV viruses
Does not stimulate cell mediated immunity

- Viral subunit: envelope glycoprotein: VaxGen

Animal studies:

Not successful: protection only vs. virus with exact same envelope proteins

- Phase I/II: Are memory B cells enough to protect vs. HIV?
 Modest Ab response vs. limited spectrum of HIV strains
 No cell-mediated immune response
- Phase III: placebo, 2ble blind trials: Antibodies in 90% of vaccinated people, yet no protection (2005-2006: volunteer 2500 IV drug users Thailand, 5000 American gay men at risk for HIV-1)

Carrier vaccines

Use harmless viral vectors to transport HIV-1 genes into human cells. If booster is needed, different carrier must be used

 $\textit{ALVAC} \colon \textit{Canarypox}$ virus expressing 3 HIV proteins

<u>Prime/boost strategy</u>:

Combination ALVAC/ VaxGen

Phase I/ II: Safe and immunogenic: Ab, CD4+ & few CD8+ cells Phase III: Thailand study: 16,000 patients, \$120 million

Merk Ad5: Adenovirus5 expressing 3 HIV proteins

Phase I: Safety and immunogenecity: elicits CD8+ responses

Phase II: currently ~3000 volunteers in US and

Caribbean

Problem: In developing countries ~80% pre-existing immunity to Ad5!

DNA vaccines

- Strategy:
 - Inject large amounts of DNA which codes for viral protein
 - Elicits immune response against that protein
- · Successful in animal trials
 - Generate killer T cell response
- Can we find a single protein that will elicit immune response against many HIV strains?
- Currently in Phase I : Oxford-Nairobi Prostitute Vaccine (Prime/boost: naked DNA modified <u>vaccinia Ankara virus</u> as HIV gene carrier)

HIV trials in progress: 2006

Phase	Candidate vaccine	Start	Vo	lunteers
	Synthetic peptide V3 (B) (United Biomedical)	1994	30	Healthy
MI	gp120 (B) (VaxGen)	1995	33	IDU
1	gp120 (B) (Chiron)	1995	54	Healthy
1	HIV-1 immunogen (Remune)	1996	30	HIV+ve
II.	HIV-1 immunogen (Remune)	1997	297	HIV+ve
MI	gp120 (B/E) (Chiron)	1997	380	Healthy
L/III	gp120 (B/E) (VaxGen)	1998	92	Recovering IDI
MI	ALVAC-HIV(vCP1521) + gp120 B/E (Aventis Pasteur and Chiron)	2000	65	Healthy
	and ALVAC-HIV(vCP1521) + gp160 B/E (Aventis Pasteur)		65	Healthy
MI	ALVAC-HIV(vCP1521) + gp120 B/E (Aventis Pasteur and VaxGen)	2000	125	Healthy
III	gp120 B/E (VaxGen)	1999	2500	IDU

(From Rerks-Ngarm et al. ;AIDS, 2006, 20: 1471-1479)

HIV trials in progress: 2007

Trial phase	Vaccine type	HIV Insert	HIV-1 sequences in vaccine	Trial sites	Volunteers	Associated company
Phase III	ALVACIge120 ¹⁷	1 canary pox expressing Gag, hol, gp120 followed by boosting with 2 gp120 proteins	B and E	Thelland	Fully enrolled 16,000 volunteers efficacy trial	Sanofi Pasteur, VauGen
Phase II proof of concept	AdS ^{re}	3 AdS vectors expressing Gag, Pol, and Nef	•	Americas, Caribbean, South Africa	4,500 volunteers are testing same and cross-clade protection in the presence of low and high levels of pre- existing immunity	Merck
	DNAUNAS**	6 DNAs expressing Gag, Pol, Net, and clades A-C Enus followed by boesting with 4 AdS vectors expressing Gag-Pol and clades A-C Envs	AC	USA, Caribbean, Central and South Africa	720 volunteers will test for cross-clade protection	Developed by the NIH vaccine research center
Preparing to enter phase II	COMUNIVA ⁴⁰	DNA expressing Gag, Pol, and Env boosted by MVA expressing Gag, Pol, and Env	8	Americas		GeoVax, developed at the Emory Vaccine Center and NIH
	DINAMINA	7 DNAs expressing sequences from clade A and B Gag, clade B Not, clade B Rnv, clade B Env, chimeras of clade A and C Envs and 1 MNA expressing a clade A/E recombinant from Thalland	A-C and E	Europe, Afric	,	Developed by Karolinska Institute, Walter Reed Army Institute for Research, and NH
	DNA/MYAC ⁶³	2 DNAs expressing gp120 and a Gag-Pol-Nef fusion protein boosted by a single MVA expressing the same proteins	Chinese B/C recombinant	China		Developed by EuroVac

Dangers of Vaccine Trials

- Most researchers feel first HIV vaccines will not be more than 40-50% effective
 - Will vaccinated individuals engage in higher risk behaviors?
 - Vaccine could cause as much harm as it prevents
- Future vaccines cannot be tested against placebo, would be unethical

Summary of lecture 10

The HIV-1 virus

- Life cycle
- Clinical course of disease: acute, chronic,AIDS

The HIV vaccine

- -5 challenges for vaccine development
- -Possible vaccine alternatives
- -Current HIV vaccines in advanced clinical trials: VaxGen, ALVAC, AD5 $\,$
 - -Dangers of vaccine trials