Vaccines and Immune Response

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NIH

Outline: Vaccines and Immune Response

- Vaccines
- History of Vaccination
 - A Great Success for Public Health
 - Challenges for the Future
- Vaccine Mechanism
 - Humoral immunity via antibodies from B-cells
 - Cellular immunity via killing and help from T-cells
- Vaccine Construction
 - Types of vaccines
- Measurement of Immune Response
- Statistical Analysis of Immune Response
- Poliomyelitis vaccine development

Vaccines

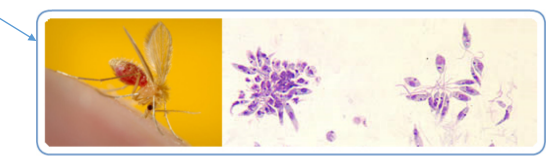
- A biological preparation that is used to induce immunity to a particular disease.
- One of the greatest public health inventions ever.
- A mystery revealed by the science method
- For me, a fascinating blend of biology, immunology, and statistical methodology.
- Lucky to be able to work in this area.

`Vaccination' in History

- People observed that, for select infections, people who recovered were subsequently immune
- Led to deliberate infections
 - China & Europe smallpox (variolation)
 - Dried pustules scratched on skin
 - Leishmanization
 - Lesion exudate scratched on buttocks
 - Chicken Pox parties
 - Zika infection for 10 year old girls?



FIGURES SHOWING VACCINATION PUSTULES From a Chinese work on Vaccination



Jenner and Vaccination

- Jenner was a 18th century English physician
- Observed that milkmaids got cowpox but not smallpox
- Gave cowpox to a 9 year old boy waited a while then ``challenged" with variolation of smallpox.
 - Success!
- Vaccination with cowpox compulsory in 1853
- Vaccination was controversial



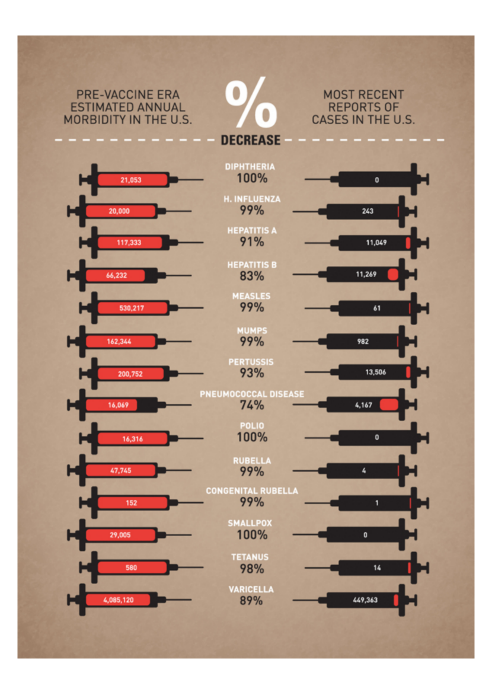
TABLE 1. Vaccine-preventable diseases

Disease	Year
Smallpox*	1798+
Rabies	1885+
Typhoid	1896+
Cholera	1896+
Plague	1897+
Diphtheria*	1923+
Pertussis*	1926+
Tetanus*	1927+
Tuberculosis	1927+
Influenza	1945&
Yellow fever	1953&
Poliomyelitis*	1955&
Measles*	1963&
Mumps*	1967&
Rubella*	1969&
Anthrax	1970&
Meningitis	1975&
Pneumonia	1977&
Adenovirus	1980&
Hepatitis B*	1981&
Haemophilus	
influenzae type b*	1985&
Japanese	
encephalitis	1992&
Hepatitis A	1995&
Varicella*	1995&
Lyme disease	1998&
Rotavirus*	1998&

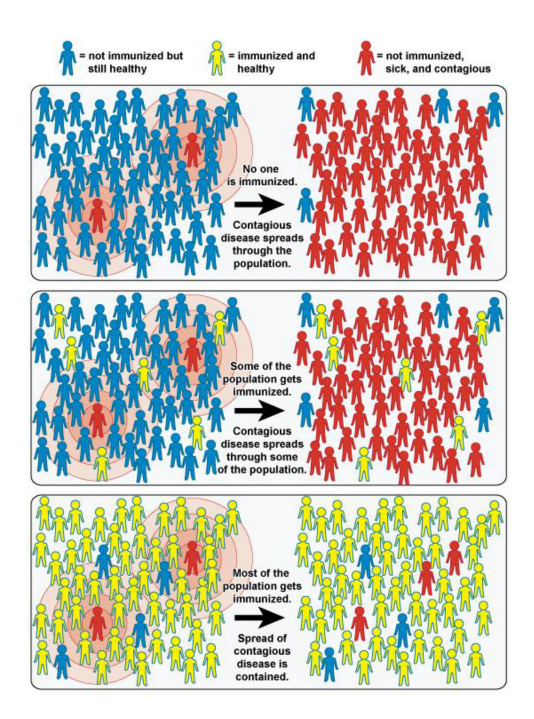
* Vaccine recommended.

+ Vaccine developed

& Vaccine licensed



Vaccination has enormously improved Public Health



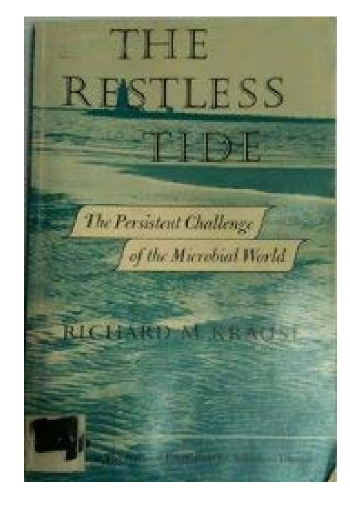
Vaccines work both directly and indirectly through herd immunity

Herd Immunity general immunity in a population based on acquired immunity by a high proportion of members

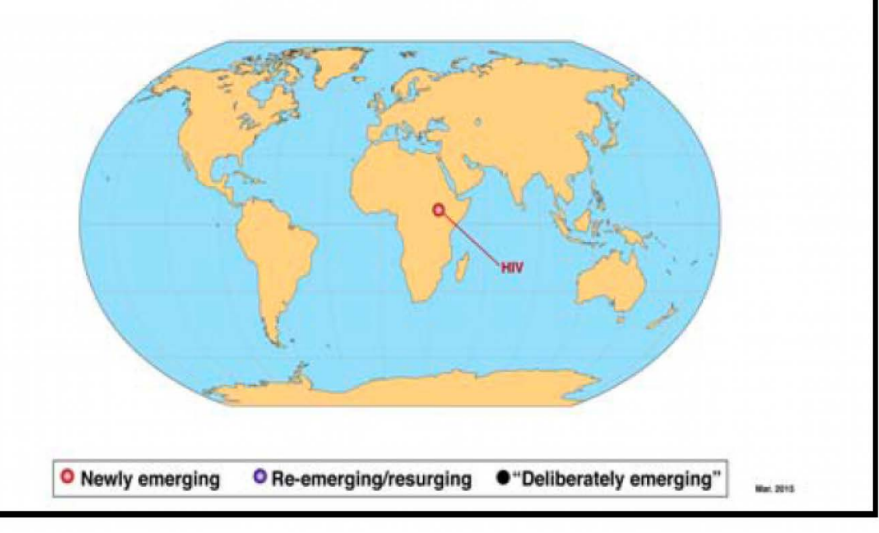


Misplaced complacency

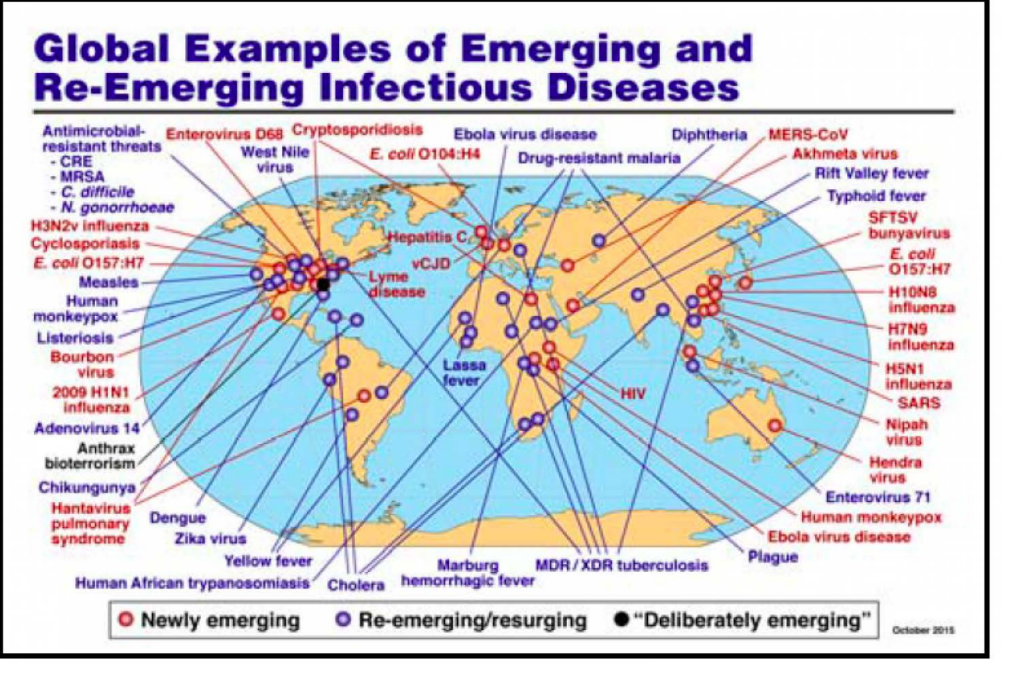
- By the late 1970s infectious diseases were widely thought to be tamed.
 - Vaccines
 - Antibiotics
- Pathogens continually evolve in response to selection pressure---drugs, habitat, human immunity
- More interconnection, habitat exposure climate change---mixes things up
- Bio-warfare



Global Examples of Emerging and Re-Emerging Infectious Diseases



Dr. Fauci's slide circa 1985



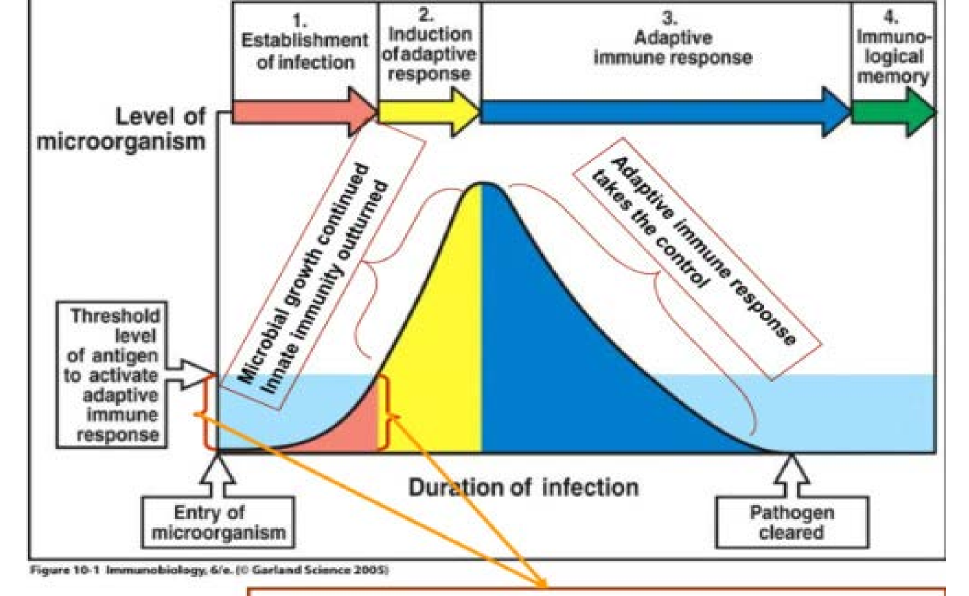
Dr. Fauci's slide Today

Vaccine Challenges Today

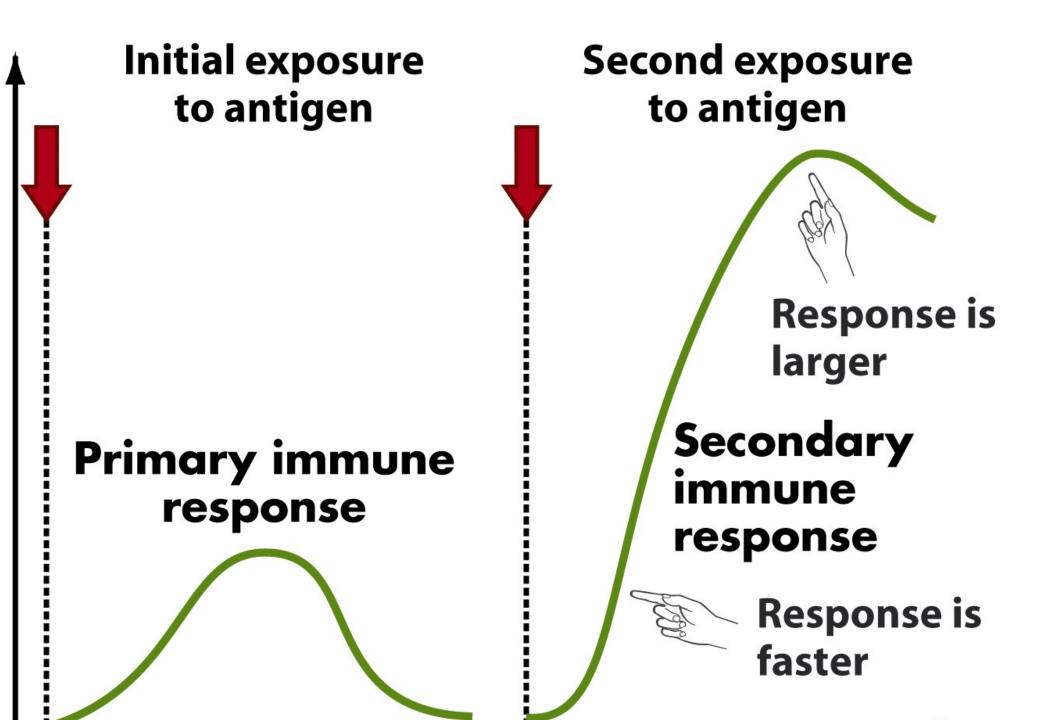
- Tough infectious diseases
 - HIV
 - Malaria
 - Tuberculosis
- Rapid response to new emerging infectious diseases
 - West Nile
 - Ebola
 - Zika
- Outside the box
 - Cigarettes: (NicVAX) Opiods
 - Cancer: construct a unique vaccine for each patient based on their tumor
 - Mosquitoes as flying syringes

How do vaccines work?

- The immune system's memory of a prior infection allows a rapid response that can quickly destroy a pathogen's expansion
- Vaccines exploit this by introducing a fake germ that induces the immune response, but doesn't cause disease
- When the real germ appears it is rapidly destroyed before causing disease



Innate immune response function at this stage to control most of the infection. Otherwise, we would have been suffering from different infections all the time.



Adaptive Immune response to a pathogen

- B-cells and T-cells are immune cells.
 - 70,000 to 2,000,000 cells/ml in blood
- A cell recognizes a unique sequence of amino acids = part of a germ

I attack

invaders outside

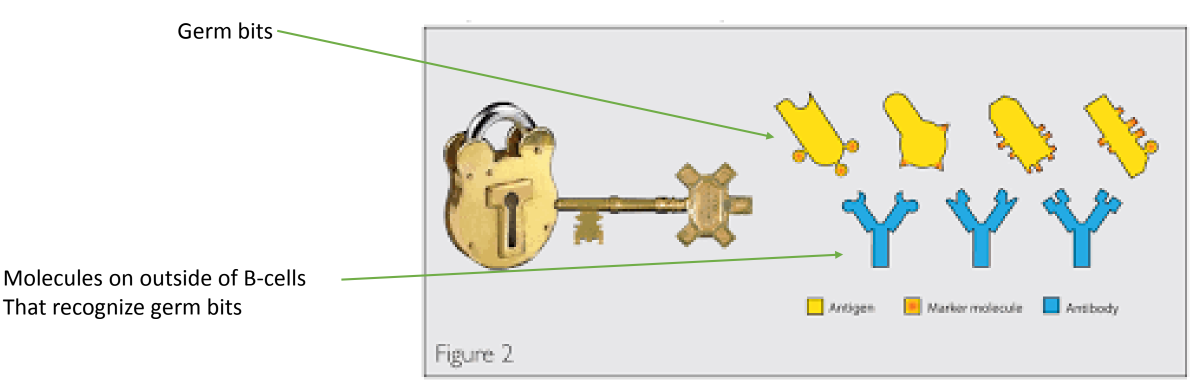
the cells.

l attack

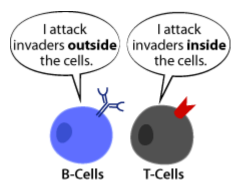
invaders inside

the cells.

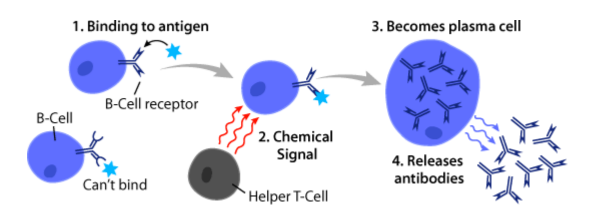
T-Cells



B-Cells



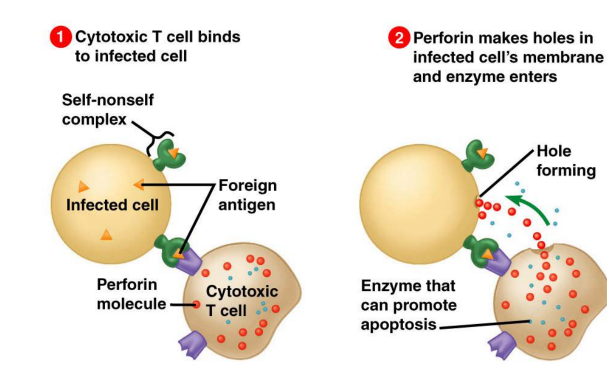
- B-cells work by identifying bits of extra-cellular germs
- Get signals from helper T-cells to expand and churn out antibodies

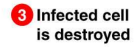


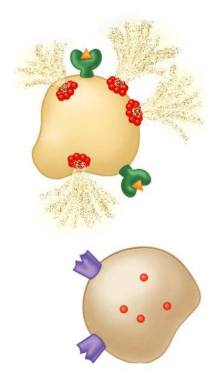
- Antibodies block germs or 'tag' them for killing
- After infection, some progeny become 'memory' B-cell for rapid response

T-cells

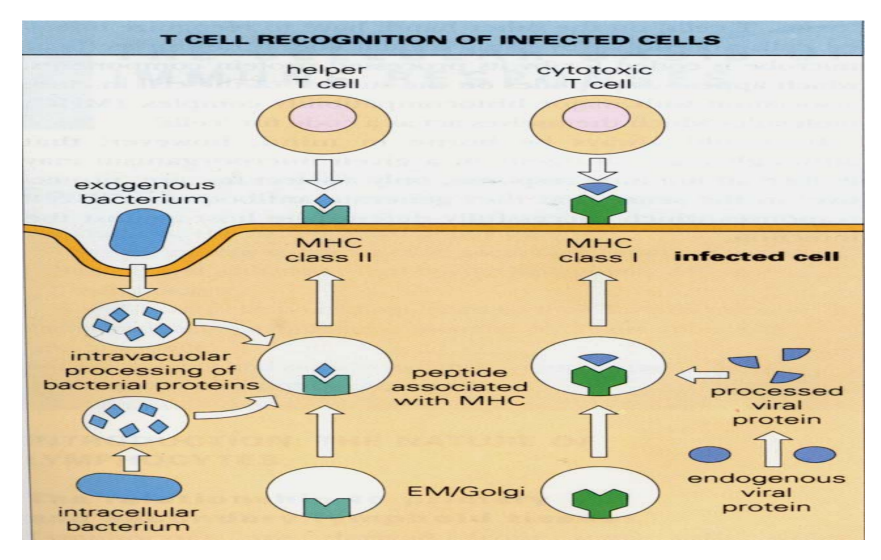
• T-cells work by identifying germ infected cells and killing them







T-cell recognition, the details



Immune Memory





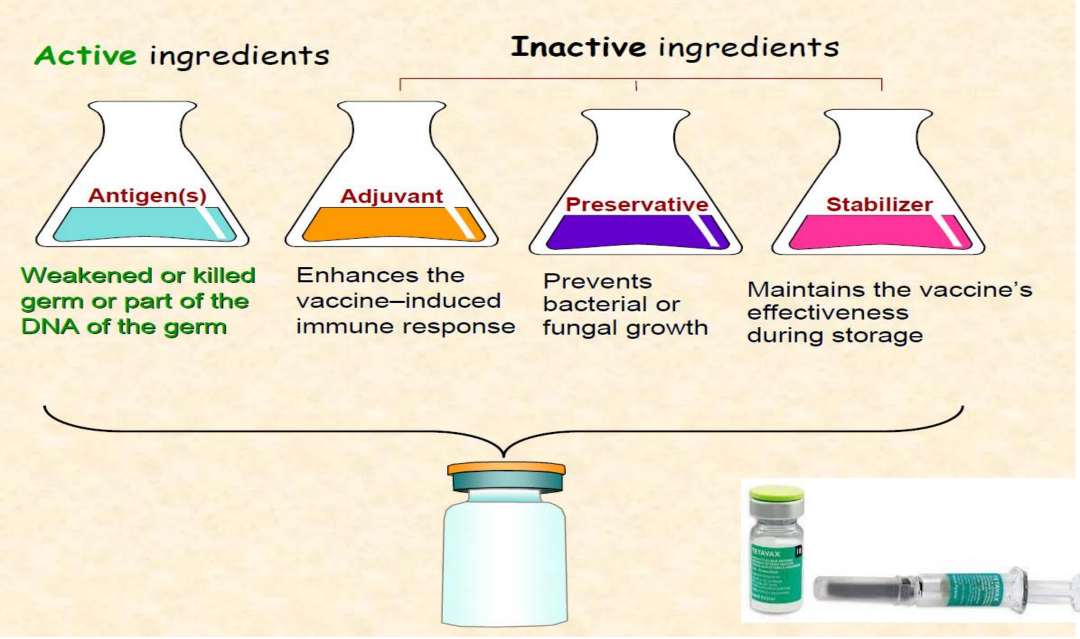
Once a lymphocyte has recognised Some cells then become long antigen a foreign antigen it expands lasting >20 years 'memory' cells. to eliminate the infection Effector Memory Infection Initial exposure Second exposure to antigen to antigen expansion Lymphocytes Antibody concentration Response is larger Pathogen Secondary **Primary immune** immune response response Response is faster Time Memory cells respond very quickly to

subsequent exposure to antigen

Types of vaccines (antigens)

- Goal is to trick the immune system to responding to a benign proxy for the germ. . . How?
- KILLED: Grow the germ and kill it
- ATTENUATED: Grow the germ in a hostile environment
 - Cold adapted strains can't handle body heat (think polar bears in the Sahara)
 - Radiation (think slow moving zombies)
- SUBUNIT: Grow the germ and snip out part of it
- VECTOR: Modify an benign virus to deliver germ bits

What's in a typical vaccine?



Immune Response Assays

- Measurement and analysis of the immune response to vaccination is a key feature of vaccine development
 - Identify which aspects of the immune system prevent disease
 - Identify and enhance certain aspects of the immune system
 - Identify a 'correlate of protection' e.g. Antibody > threshold => protection)
- Many ways to measure the immune system
- Measurement involves biological systems, assays can be twitchy
 - Qualification, Validation
 - Lots of statistical issues

Assays



- Antibodies: ELISA
 - Enzyme linked immuno-sorbent assay.
 - measures how the antibodies bind to their target antigen
- Antibodies: TZM-bl
 - Genetically engineered HIV infectable HeLa cells from Henrietta Lacks
 - light up with firefly biolumenescence when infected with HIV
- T-cells: ICS
 - intracellular cytokine staining measures secreted chemical signals from T-cells
 - e.g. ``come here'' ``do this'' ``have some poison"
- Antibodies: Standard Membrane Feeding Assay
 - Evaluate vaccine induced birth control for sexual-stage malaria parasites
- Antibodies: Binding Antibody Multiplex Assay
 - Efficiently evaluate binding of multiple antibody types at multiple sites

Measurement of Antibodies

- Antibodies are produced by B-cells and circulate in blood
- An antibody binds to a unique string of 5-8 amino acids on the germ
 - Can neutralize the germ by preventing it from infecting a cell
 - Many antibodies can glom on germ and signal other cells to attack it
- Need to measure antibodies for the pathogen of interest
- Two ways to measure
 - Binding assay --- antibody sticks to its cognate antigen
 - Functional assay --- antibody prevents germ from infecting a cell

Antibody

Antiaens

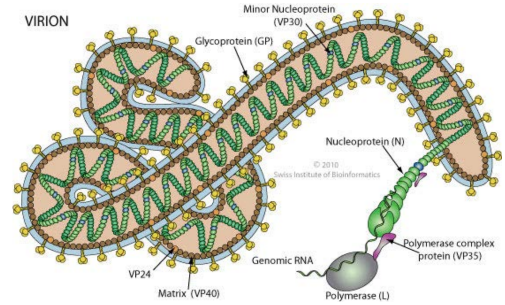
Antiaen-bindina site

Ebola Vaccines

- Experiments have suggested antibody response was correlated with survival in vaccinated monkeys exposed to disease
- Vaccine studies have looked at antibody response to see if vaccine is inducing an immune response
- Antibody responses measured in Ebola vaccine trials in West Africa

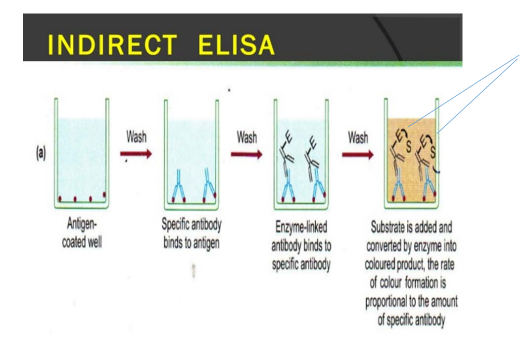
Antibody to Ebola virus

- Rec.
- Outer shell of Ebola virus is a glycoprotein GP (mixture of sugars and protein)
- Vaccines express bits of the GP to induce immune response
- Elisa assay used to measure the antibodies to GP

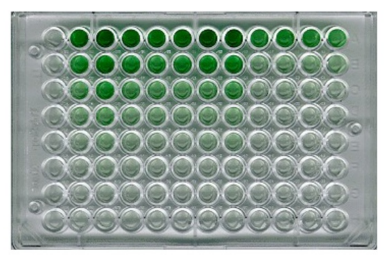


Binding antibody ELISA

- Enzyme-Linked ImmunoSorbent Assay performed in a 96-well plate
- Measures abundance of antibodies for a specific antigen



Enzyme-Linked Immunosorbent Assay (ELISA)

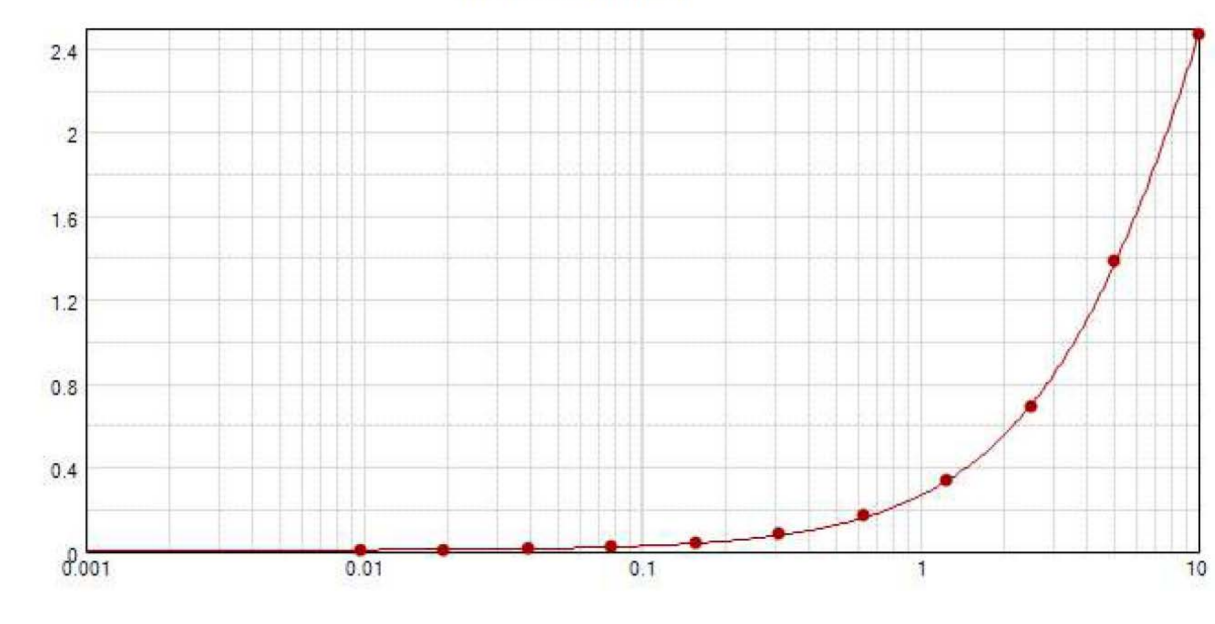


96-well plate

REFERENCE STANDARD of Each Well has a know concentration

	1	2	Э	4	5	6	7	0	9	10	11	12
А	2.671 0.048	1.509 0.042	0.767	0.416	0.217	0.121 0.038	0.082 0.037	0.061 0.037	0.054 0.040	0.048	0.049	0.130 0.037
В	2.353 0.046	1.347 0.048	0.697	0.341 0.038	0.201 0.038	0.113 0.038	0.075 0.038	0.059 0.038	0.049 0.038	0.045 0.037	0.043 0.039	0.111 0.039
С	1.075 0.041	1.066 0.042	0.690 0.043	0.616 0.040	1.446 0.043	1.222 0.042	0.426 0.040	0.548	0.323	0.350	2.507 0.048	1.136 0.042
D	0.666	0.574 0.040	0.428 0.039	0.290 0.040	0.929 0.043	0.788 0.041	0.245 0.040	0.267 0.039	0.176	0.189 0.039	1.577 0.044	0.579 0.040
E	0.316 0.039	0.306 0.039	0.191 0.039	0.190 0.039	0.515 0.040	0.342 0.040	0.156 0.039	0.199 0.052	0.117 0.038	0.117 0.039	0.873 0.041	0.294 0.039
F	0.161 0.039	0.150 0.039	0.119 0.041	0.125 0.038	0.246 0.039	0.172 0.039	0.087	0.097 0.039	0.082	0.238	0.436 0.040	0.166 0.039
G	0.101 0.038	0.109	0.075 0.041	0.082	0.143 0.039	0.111 0.042	0.073 0.039	0.106	0.060	0.073	0.288 0.039	0.099 0.039
н	0.065	0.067 0.038	0.066	0.064 0.038	0.098 0.038	0.103 0.039	0.056	0.054	0.054	0.053 0.038	0.154 0.039	0.070 0.038

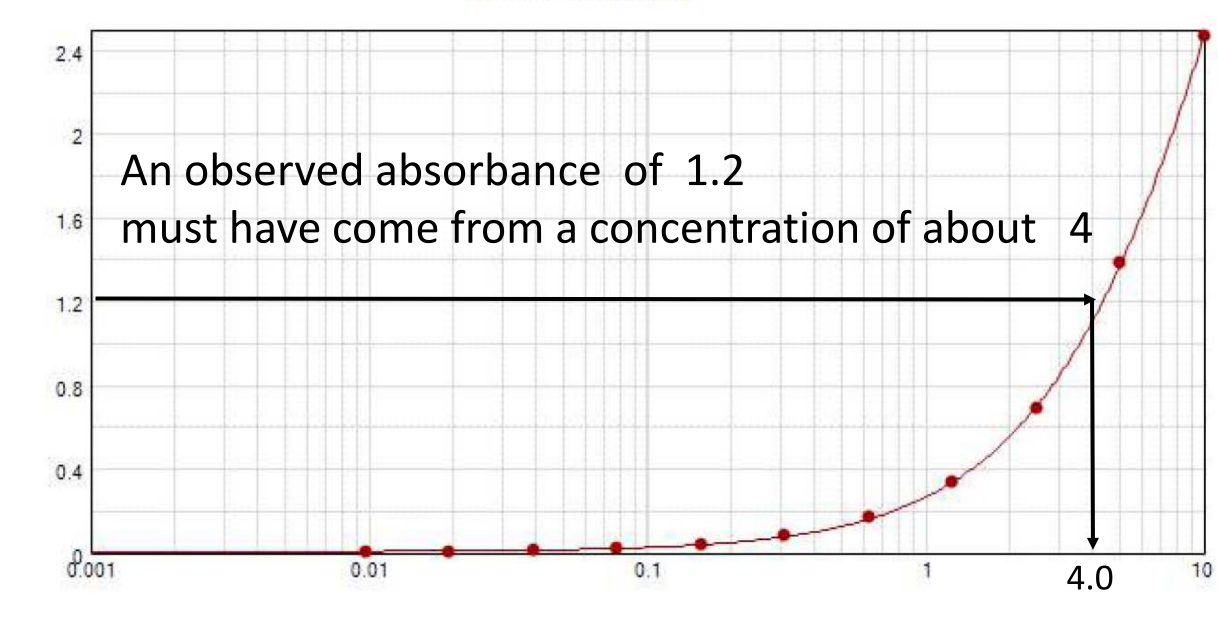
Absorbance



Relative Concentration (ELISA units/mL)

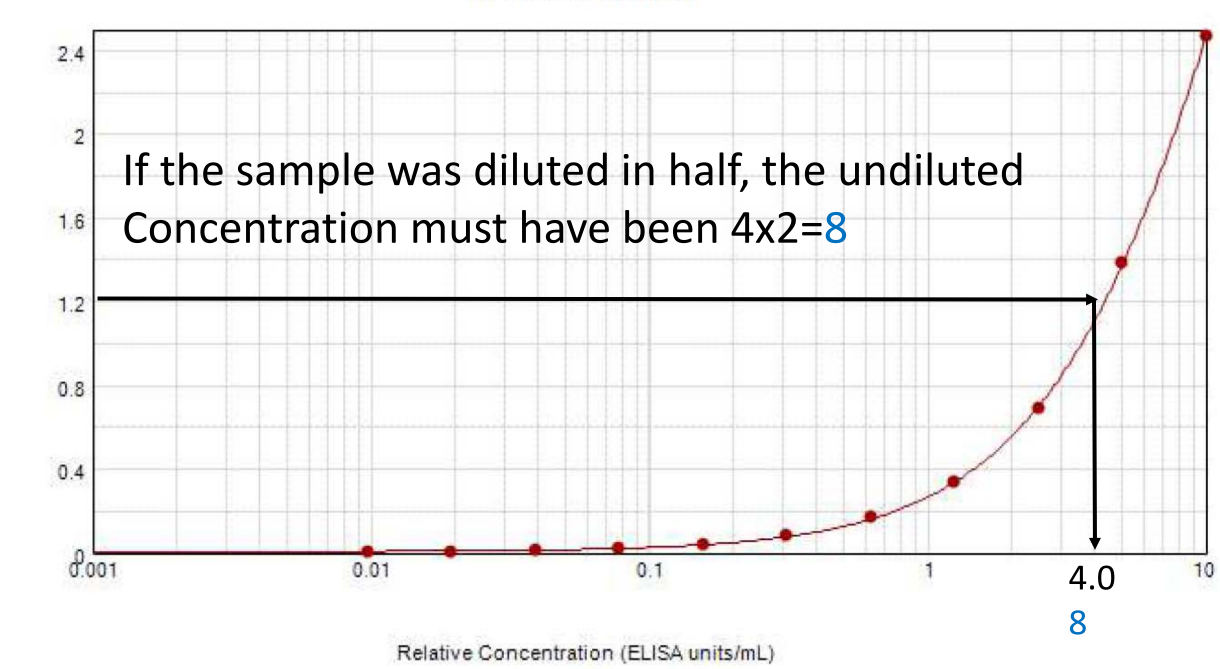
Reference Standard

Reference Standard

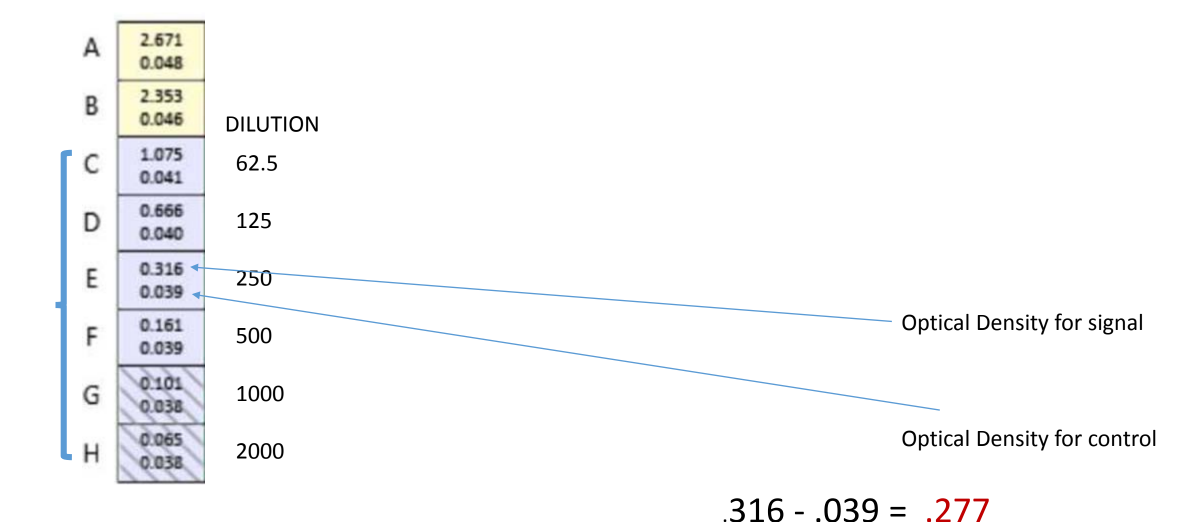


Relative Concentration (ELISA units/mL)

Reference Standard

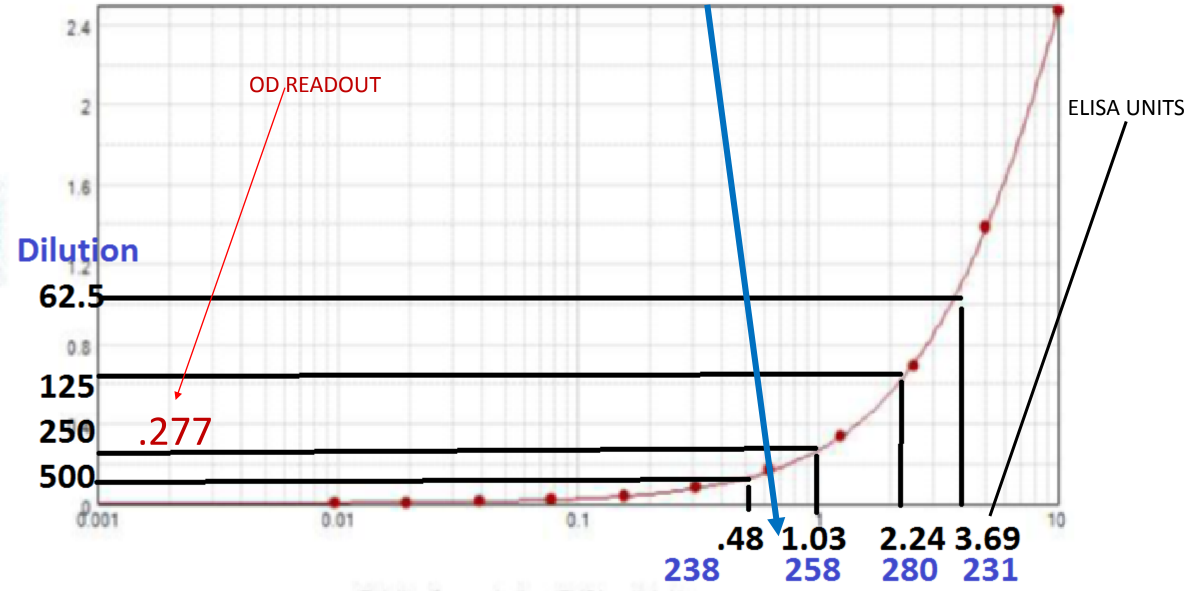


Optical Density for Subject 1's sera at different dilutions



P E R S O N # 1

1.04* 250 = 258 => ELISA UNITS FOR UNDILUTED



Deletive Persententian (EUC) units (m))

Readout for person 1

Sample	Wells	Values	Result	Dilution	Adj.Result	
1	C1	1.035	3.69	62.5	230.84	٦
2	D1	0.626	2.24	125.0	279.77	
3	E1	0.277	1.03	250.0	257.76	
4	F1	0.122	0.48	500.0	238.16	
5	G1	Masked	Masked	1000.0	Masked	
6	H1	Masked	Masked	2000.0	Masked	

AVG = 251.63

Masked values are too low or aberrant to be credible and excluded.

- lower readout than negative control

- sample coefficient of variation (S/ \overline{X}) improves a lot with their elimination

Positivity Criteria

- Vaccine studies like to report the response or 'take' rate
 - A relic from variolation when a 'take' could be observed if pustule formed where scratched?
 - Take => you'll be protected?
- Responder definition 1
 - Take Y from unvaccinated controls & determine mean + 2 std = c
 - If readout Y > c => responder
- Responder definition 2
 - Take placebo group 1 month change $Y_1 Y_0$ & determine mean + 2 std = c
 - If vaccine $Y_1 Y_0 > C => Responder$



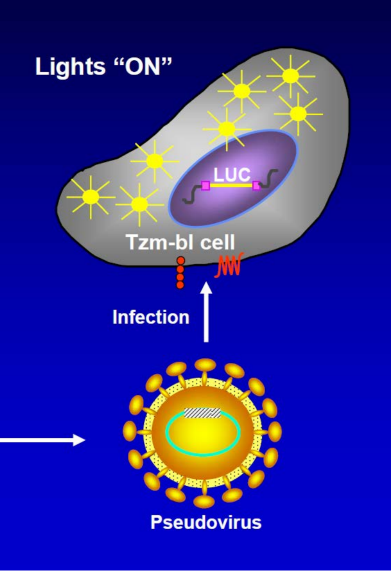
- Use t-test to determine if the mean Y differs between groups
- Use Fisher's exact test to determine if the response rate differs between groups

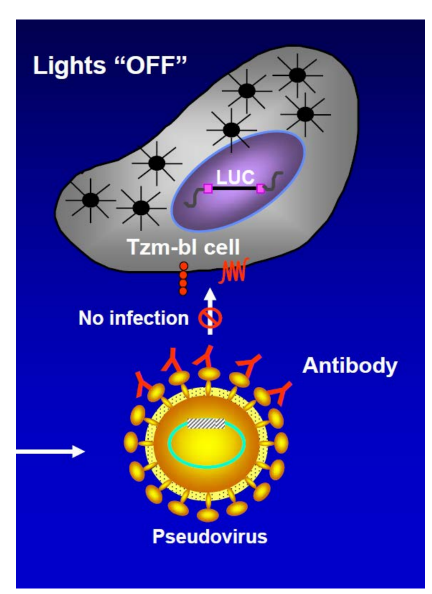
Functional antibody assay: TZM-bl

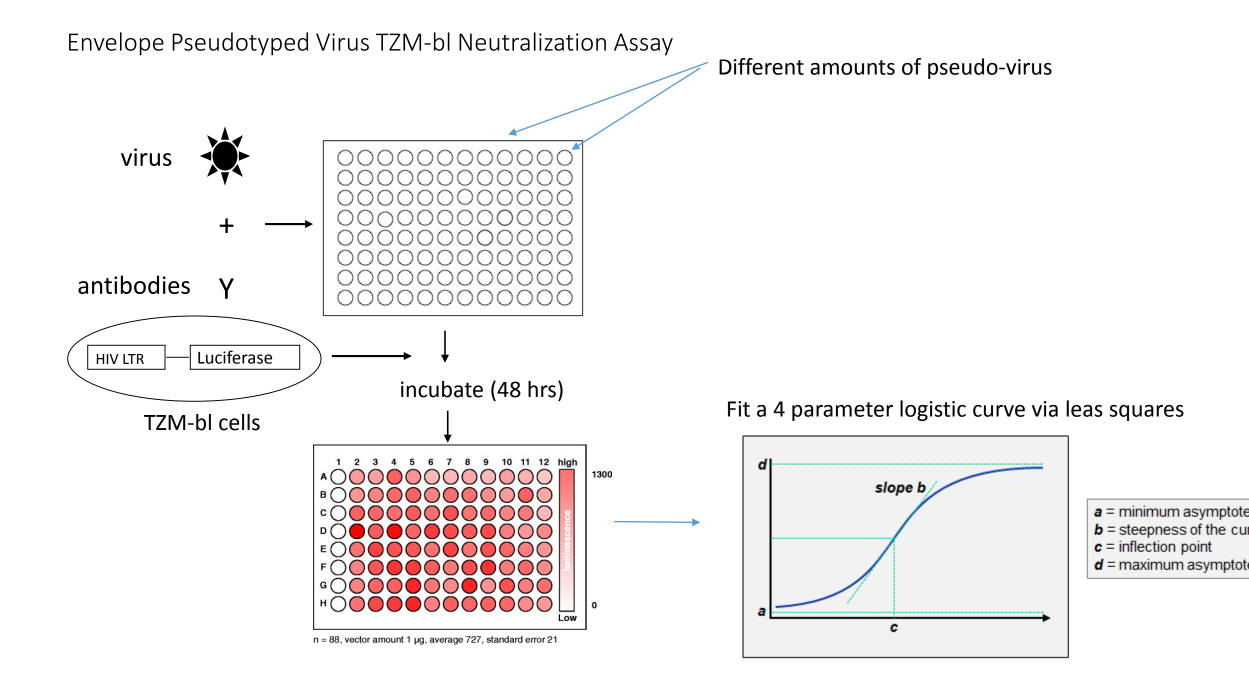
- TZM-bl are genetically engineered HIV-infectable cells that contain a gene for luciferase which makes fireflies glow
 - Lucifer light-bearer (lucem ferre)
- If HIV infects a TZM-bl cell HIV replication within the cell turns on the luciferase gene.
- Mix serum from vaccinees, TZM-bl cells, and HIV-like virions in wells at various concentrations

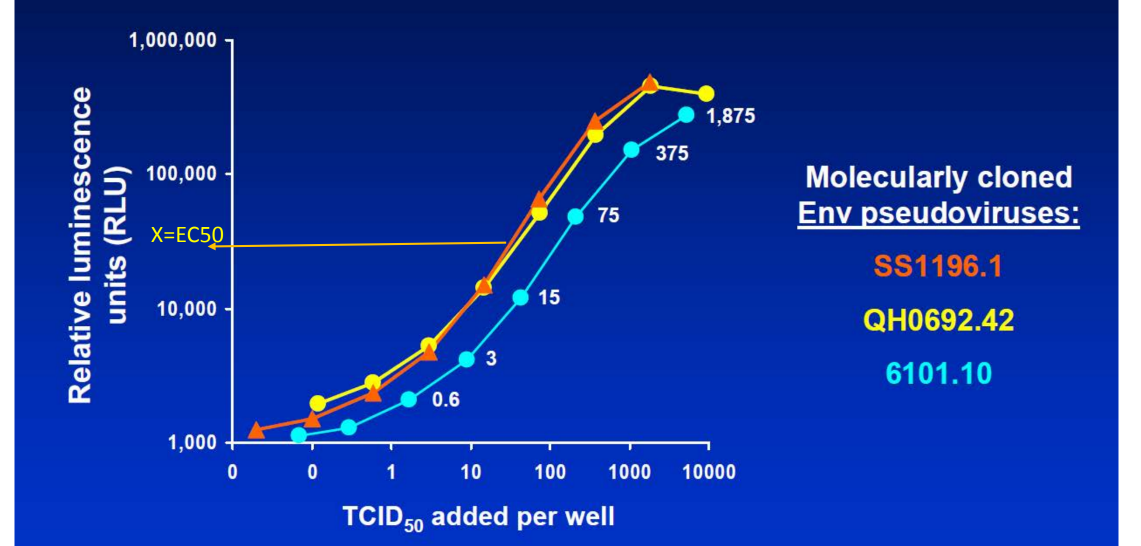


How the TZM-bl Assay works









Analysis of TZM-bl readout

- Interest in how much antibody is needed for protection
- Gave 21 animals different amounts of antibody
 - Measured amount in blood using TZM-bl assay X
 - Animals were 'challenged' with SHIV and infection status recorded
 Y = 1 if infected
 Y = 0 uninfected
- Used maximum likelihood to estimate a, b of probit regression

 $P(Y = 1) = \Phi(a + b X)$

Standard Normal Cumulative Distribution Function

TZM-bl Infection Old assay

3.2582	0	2.0899
3.6101	0	2.0899
	0	1.6021
3.1715	0	1.6021
2.9314	0	1.1761
2.9256	1	1.2553
3.0066	0	1.2553
2.5736	0	1.1139
2.6198	0	1.0792
	1	0.8451
1.9655	0	0.7782
2.4441	1	0.8451
1.9834	1	0.4771
	1	0.6021
	0	0.6021
	0	0.9031
	0	0.699
	1	0.6021
	1	0.4771
	1	0.4771
	1	0.4771

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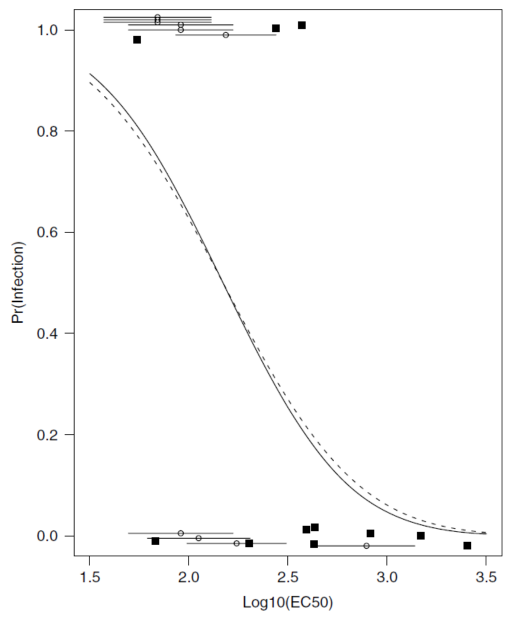


Table 1
Pseudo-likelihood estimates for the ID_{50} and ID_1 . Confidence
intervals are obtained by profiling the pseudo-likelihood ratio
$test\ statistic.$

		95% profile CI					
Parameter	Estimate	Lower	Upper				
$\overline{\mathrm{ID}_{50}}$	2.17	1.46	2.62				
ID_1	3.32	2.67	8.04				

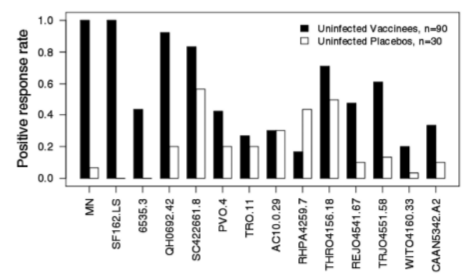
Target for a vaccine may be to achieve 3.32 units of Antibody

TZM-bl assay in a human vaccine trial

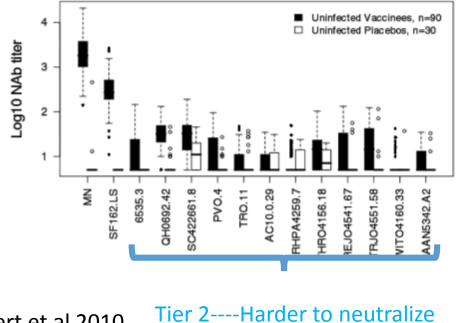
- VAX004 was the first phase III HIV vaccine trial
 - Used 2 gp120 proteins from 2 HIV-1 strains
 - Overall there was no efficacy
- Use the TZM-bl assay to characterize the immune response of an inefficacious vaccine.
 - Evaluate 14 types of HIV virus: HIV-1_{MN}, SF162.LS, and 12 'tier 2' viruses

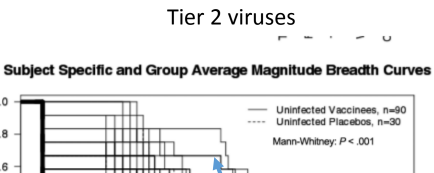
Genomic RNA

Neutralization Response Rates



Neutralization Response Levels





1.0

0.8

0.6

0.4

0.2

0.0

to Response

Positive response rate

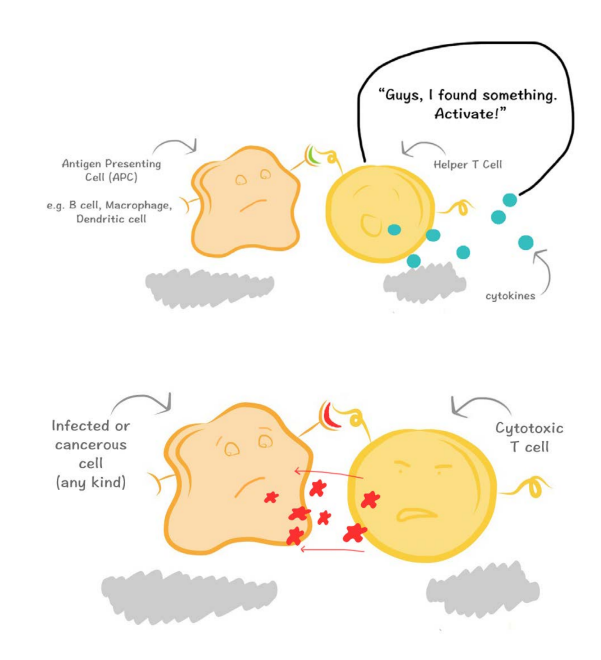
1.0 1.5 2.0 Log10 NAb titer This vaccine had a TZM-bl value X of 1.6 or better for 8 of 12 tier 2 viruses

Gilbert et al 2010

Α

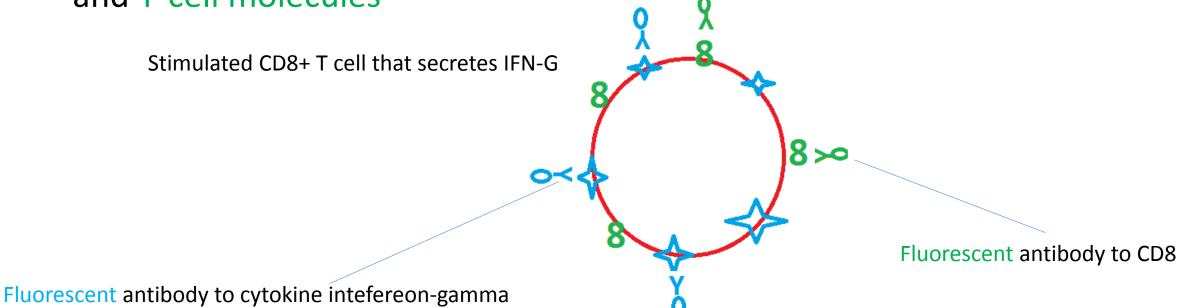
Measurement of T-cells

- T-cells secrete chemicals when confronted with a cell that displays part of *their* germ
- The chemicals either
 - Signal other cells to come and destroy
 - Kill an infected cell
- Assays 'annoy' T cells with e.g. HIV antigen and look for chemicals, if present, T-cell recognized HIV....

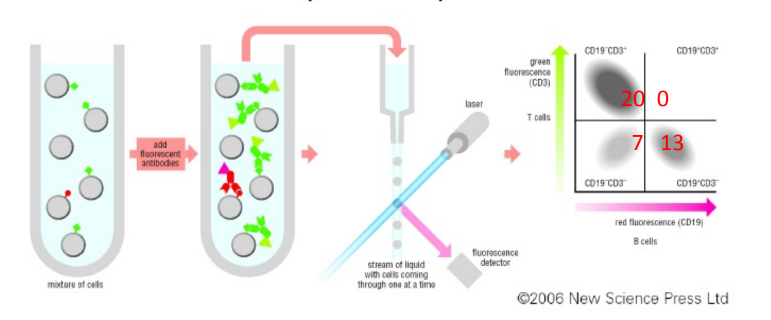


Flow cytometry measurement of T cells

- Sample circulating T-cells from blood of a vaccine trial subject
- Annoy the T-cells with HIV-peptides and see if they secrete chemicals
- Make fluorescently labeled antibodies to attach to chemicals and T-cell molecules



Flow cytometry measurement of T cells Use of monoclonal antibodies recognizing CD antigens by flow cytometry



Count # of cells in each quadrant CD19 - CD3 - 7 CD19+ CD3 - 13 CD19- CD3 + 20 CD19+ CD3 + 0

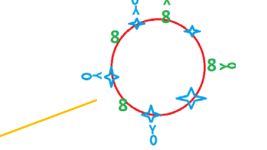
Flow cytometry: measure the amount of a protein on the surface (or inside) individual cells; measure the numbers of particular types of cells in blood (etc.)

Different parts of HIV

Table 1 Sample Data for one blood sample from one patient, CD8 cells only, in response to background and peptide stimulation (columns). The rows represent the set of 2^5 mutually exclusive outcomes of positive and negative results for the five cytokines

PatientID	Combo	Background	ENV	GAG	NEF	POL	TRV		
1481	7 + g + b + 2 + a +	0	1	53	11	10	53		
1481	7 + g + b + 2 + a - a - b + b + 2 + a - b + a - b + b + 2 + a - b + b + a - b +	0	0	21	11	13	34		
1481	7 + g + b + 2 - a + b + a + b + a + b +	5	20	298	77	97	198		
1481	7 + g + b + 2 - a - a	18	67	658	311	228	370		
1481	7 + g + b - 2 + a + b	0	0	0	0	0	0		
1481	7 + g + b - 2 + a - a - b - 2 + a - b - a - b - a -	1	0	1	0	0	0		
1481	7 + g + b - 2 - a + b - 2 -	1	0	0	1	2	3		
1481	7 + g + b - 2 - a - a	1	7	23	15	4	29		
1481	7 + g - b + 2 + a + a + b + 2 + a + b + a +	1	0	0	0	0	0		
1481	7 + g - b + 2 + a - b	5	2	3	4	8	8		
1481	7 + g - b + 2 - a +	4	9	7	0	4	5		
1481	7 + g - b + 2 - a - a - b + 2 - a - b + 2 - a - b + 2 - a - b + 2 - a - b + 2 - a - b + 2 - a - a - a - a - a - a - a - a - a -	222	380	458	393	380	297		
1481	7 + g - b - 2 + a + a + b - 2 + a + b -	0	0	0	0	0	0		
1481	7 + g - b - 2 + a - b - a - a - a - a -	28	36	12	19	34	33		
1481	7 + g - b - 2 - a + b - 2 -	0	1	1	0	0	2		
1481	7 + g - b - 2 - a - b - a - b - 2 - a - b - a - b - 2 - a - b - 2 - a - b - 2 - a - b - 2 - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - a - b - a - a - b - a - a - b - a - a	242	357	253	269	209	248		
1481	7 - g + b + 2 + a + a + b + 2 + a + b +	0	0	1	1	0	2		
481	7 - g + b + 2 + a - b + a - b +	3	3	3	0	0	0		
1481	7 - g + b + 2 - a +	6	5	5	2	2	21		
1481	7 - g + b + 2 - a - a	13	29	46	33	31	58		
1481	7 - g + b - 2 + a + a + b - 2 + a + b -	0	0	0	0	0	0		
481	7 - g + b - 2 + a - b - a - b - a - b -	0	3	0	0	0	3		
481	7 - g + b - 2 - a + b	0	0	0	1	0	1		
481	7 - g + b - 2 - a - a	8	45	23	29	89	100		
481	7 - g - b + 2 + a + a + a + a + a + a + a + a + a	0	0	0	0	0	0		
481	7 - g - b + 2 + a - b + a -	1	12	0	3	5	5		
481	7 - g - b + 2 - a +	4	5	2	1	2	2		
481	7 - g - b + 2 - a - b	1269	1898	1476	1242	1443	984		
481	7 - g - b - 2 + a + a + b - 2 + a + b -	1	1	0	1	0	0		
1481	7 - g - b - 2 + a - b - 2 +	113	424	44	89	87	88		
1481	7 - g - b - 2 - a +	64	85	24	25	32	39		
1481	7 - g - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - b - a - b - 2 - a - a - b - a - a - b - a - a - b - a - a	199351	219871	207308	204533	157181	192849		

Counts of different CD8+ Killer T-cells



CD8+ T-cells that secrete IFN-g but nothing else

Simple Analysis of ICS T-cells

PatientID	Combo	Background	ENV	GAG	NEF	POL	TRV
1481	7 - g + b - 2 - a - a - b - 2 - a - b - a - b - 2 - a - b - a - b - 2 - a - a - b - 2 - a - b - 2 - a - b - 2 - a - b - 2 - a - b - 2 - a - b - 2 - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - b - 2 - a - a - a - a - a - a - a - a - a	8	45	23	29	89	100

Patient 1481's Readout for CD8 positive T-cells that recognize TRV: 100/195379 - 8/201401 = .00047 = Y

Can do a univariate t-test of outcome Y in vaccinees versus placebos

Can do a multivariate t-test to see if the vector of outcomes differ between vacinees and placebos

Mixture model Analysis of ICS T-cells

	Positive (for IF-gamma alone)	Negative (for IF-gamma alone)					
Stimulated	100	195279					
Unstimulated	8	201401					

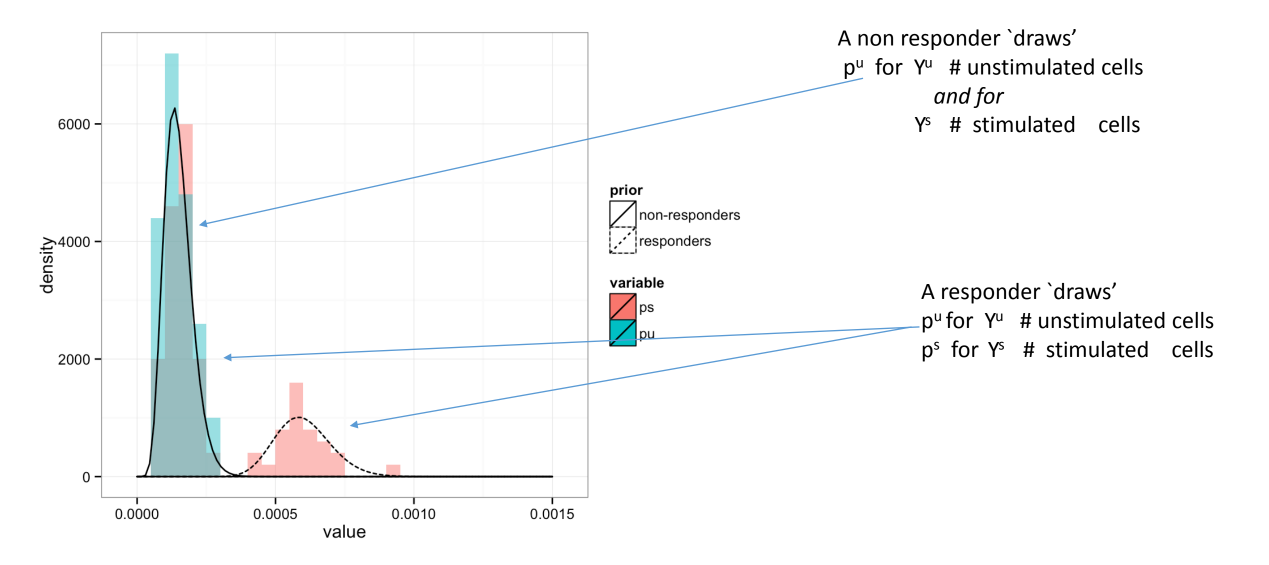
- Let p^s be the true probability a stimulated cell is positive
- Let p^u be the true probability an unstimulated cell is positive
- Y^s = 100 ~ Binomial (p^s, 195379)
- Y^u = 8 ~ Binomial (p^u,201409)
- Assume p^u follows Beta(a^u, b^u) distribution

p^s follows Beta(a^u, b^u) if non-responder i.e. R=0

p^s follows Beta(a^s, b^s) if responder i.e. R=1

- Mixture

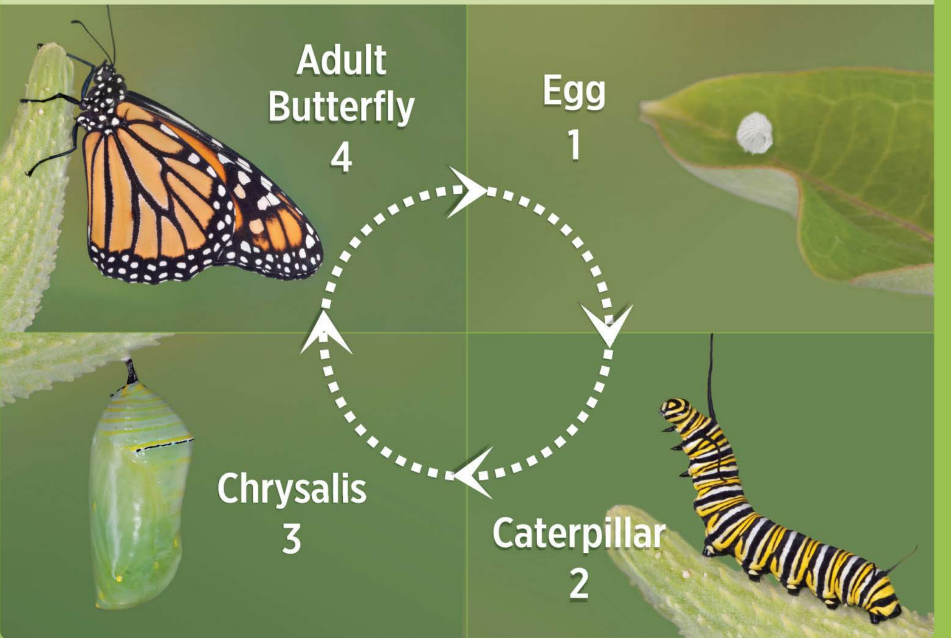
Beta distributions for p for all subjects



MIMOSA Analysis

- Each person is a non-responder (R=0) or responder (R=1)
- Can estimate Pr(R=1) = Pr(p^u < p^s) if large enough, classify as a responder
- Can be more sensitive and specific than using Fisher's exact test For a subject ``Fred"
 - Borrows strength from other subjects via use of commonly estimated Beta(a^u, b^u) and Beta(a^s, b^s)
 - Blends it with 2x2 table for Fred's cell counts
- Fisher's exact test only uses Fred's table to classify Fred.

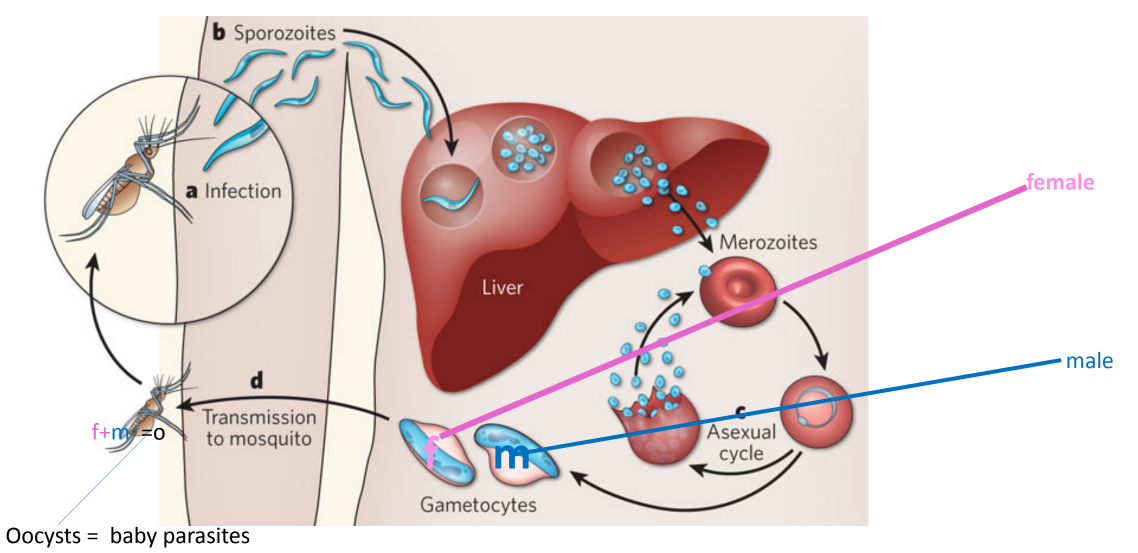
THE LIFE CYCLE OF A MONARCH BUTTERFLY



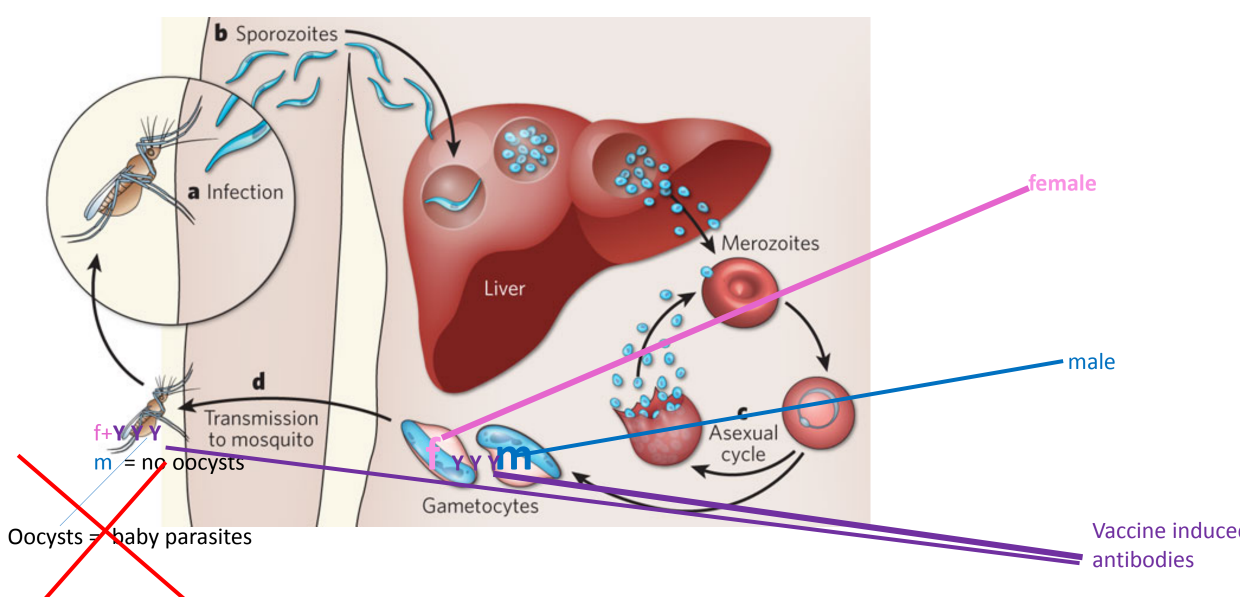
Kafka



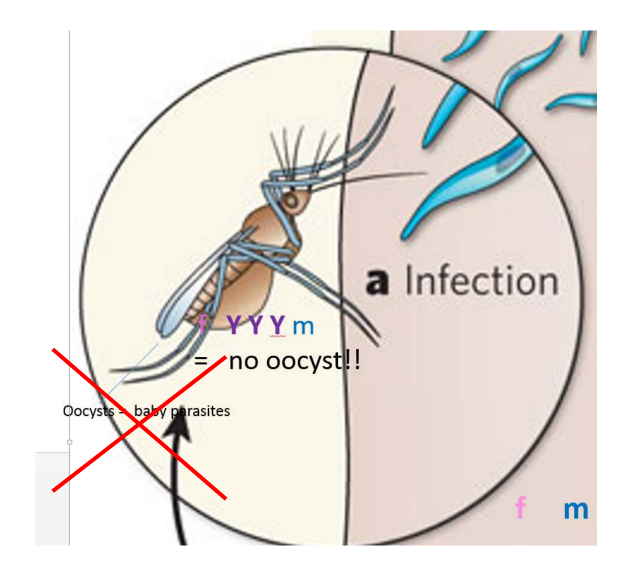
Malaria Life Cycle



Antibodies break the life cycle



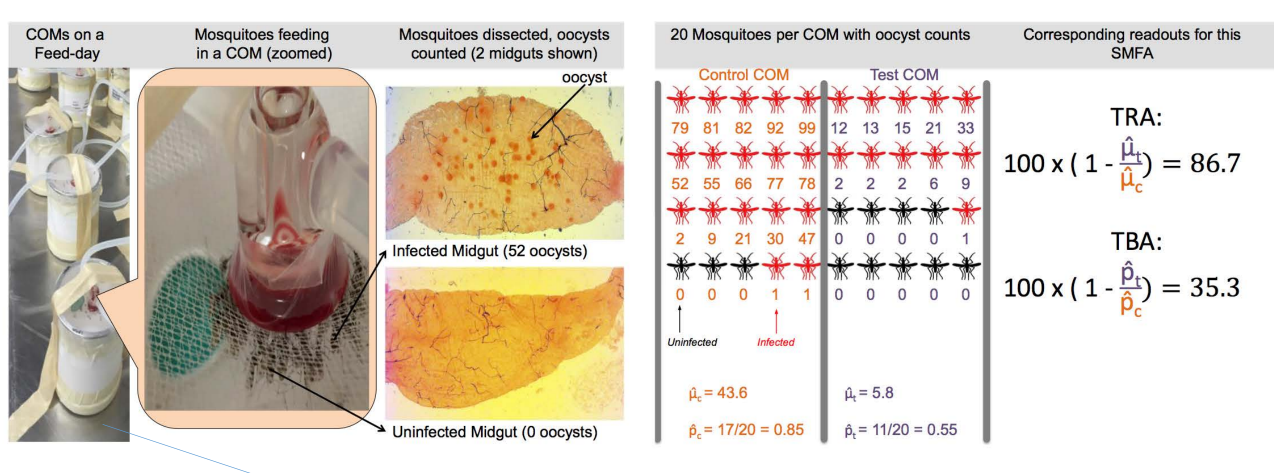
Antibodies: Barrier method of parasite birth control



Standard Membrane Feeding Assay

- Gametocyte (f/m) to oocyst to sporozoite in mosquito
- Vaccinated humans produce antibodies Y Y
- Mosquito sucks up blood and
 - mfYYfmYY ffYYm
- Antibodies prevent oocyst progeny from gametocyte parents f m
- Use a Membrane Feeding Assay to see how well antibodies block oocyst development

Assay Readouts for SMFA



Container

Assay Analysis

- SMFA assays are extremely variable in the oocyst counts
- TBA is like perfection----i.e. zero oocysts in most mosquitoes
 - A day with lots of oocysts hard to be perfect
 - A day with very few oocysts easy to be perfect
- Analysis
 - Rank vaccines by TRA (reduction in mean counts)



- To evaluate perfection (TBA) statistically standardize to 2 oocysts per mosquito on control
- Allows fair comparisons from container to container, day to day, lab to lab

Assay Analysis

i j k day container mosquito

• Use a zero-inflated negative binomial mixture model for the counts

$$\mathsf{P}(\mathsf{Y}_{\mathsf{ijk}} = \mathsf{y}_{\mathsf{ijk}}) = \begin{cases} \pi + (1 - \pi) \left(1 + \frac{\lambda_{ij}}{\theta} \right)^{-\theta}, & y_{ijk} = 0\\ (1 - \pi) \frac{\Gamma(y_{ijk} + \theta)}{\Gamma(y_{ijk} + 1)\Gamma(\theta)} \frac{\left(\frac{\lambda_{ij}}{\theta}\right)^{y_{ijk}}}{\left(1 + \frac{\lambda_{ij}}{\theta}\right)^{y_{ijk} + \theta}}, & y_{ijk} = 1, 2, \dots \end{cases}$$

• Get model based estimates of $\hat{\mu}_{c}$ $\hat{\mu}_{t}^{+}$ for an assay with μ_{c} set to 2

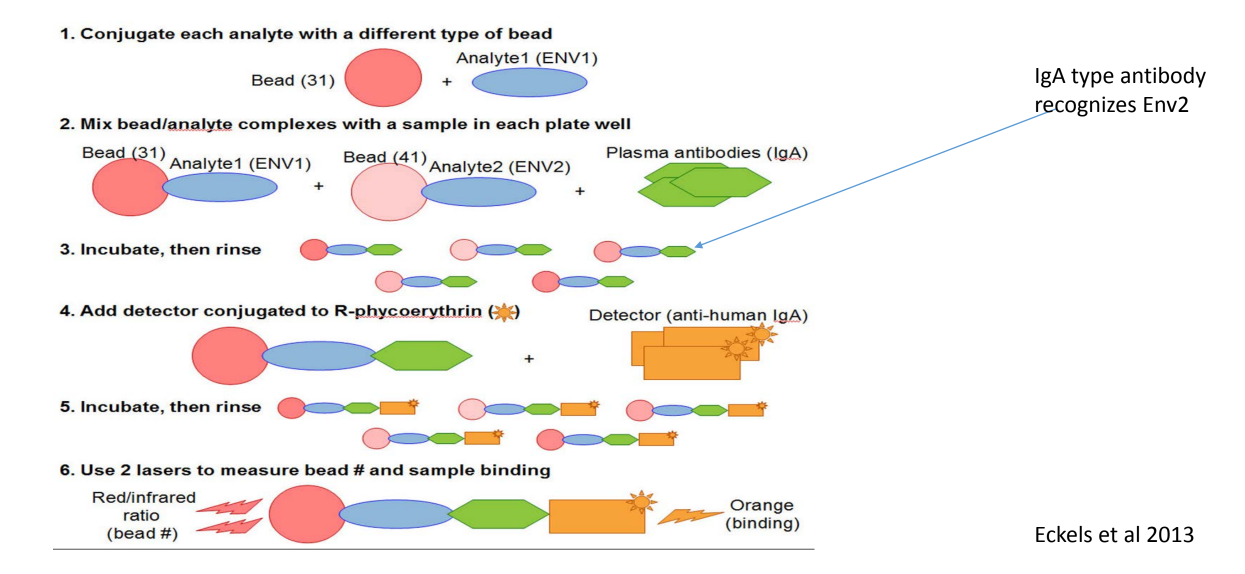
 $\boldsymbol{\hat{p}}_{c} = \boldsymbol{\hat{p}}_{t}^{:}$

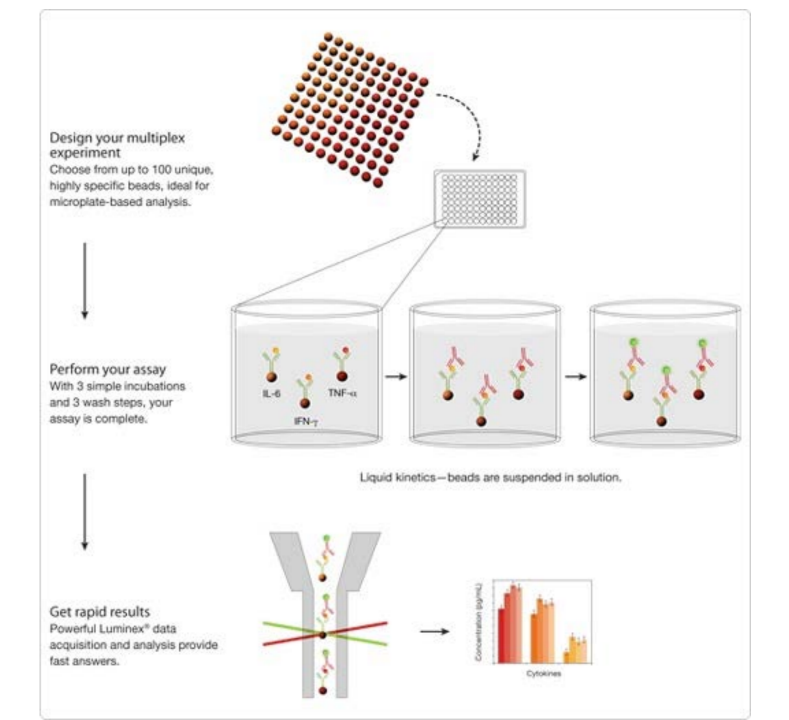
Binding Antibody *Multiplex* Assay

- Simultaneous evaluation of multiple parameters within each well
 - Multiple antigens
 - Multiple types of antibodies
- e.g. IgA type antibody recognizes gp120 region of HIV
- Used in immune correlates analysis of an HIV vaccine to efficiently evaluate many antigen/antibody types

Haynes et al 2012

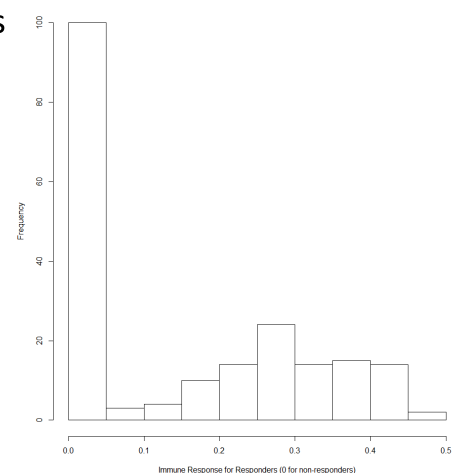
BAMA details





Analysis of Immune Response

- The immune response can be decomposed as
 - R = 1 if respond 0 otherwise
 - Y = magnitude of response in responders
- How to compare a mixture distribution between vaccinees and placebos?



Proportion of non-Responders (Y=0) and Y in Responders (Y>0)

Hu-Proschan two-part test

- Let Z_p be the test that the proportions of responders are the same in vaccine and placebo groups.
- Let Z_{γ} be the test that the average immune response Y is the same in the vaccine and placebo groups.
- Form an overall test

$$Z = \frac{Z_{P} + Z_{Y}}{\sqrt{2}}$$

• Weighted versions of the test can be incorporated.

Poliomyleitis caused by poliovirus

- Poliomyleitis is a viral disease that can infect the central nervous system and cause lasting disabilities in a small number of infected individuals.
- Polio infection is most common in children but adults are at risk too
 - Franklin Roosevelt developed polio
- Polio was greatly feared.
 - Outbreaks are unpredictable
 - Paralyzed children are a visual reminder
- National Foundation for Infantile Paralysis was formed in 1938 to develop a vaccine.





Key developments

- Virus was isolated in infected subjects 1908
- Identification of three serotypes of polovirus, each serotype has a distinctive surface and a specific antibody works against a specific type.
- Confirmation that neutralizing (blocking) antibodies protect against disease
 - At risk children who received antibodies from polio survivors saw 80% reduction in paralytic poliomyelitis compared to children with gelatin
- Growth of virus in cell culture



Vaccine developments

- Inactivated polio vaccine (IPV):
 - Three serotypes grown in cell culture and then killed by formalin
 - Developed by Jonas Salk, injected
 - Can't cause disease
- Oral polio vaccine (OPV):
 - Three serotypes were weakened by repeated passage in cold non-human cells
 - Replicates in the gut. Very rarely causes disease or mutates to a more virulent form
 - Developed by Sabin, swallowed

1954 Polio Field Trial of Salk Vaccine

- Salk Vaccine was promising but unproven.
- A field trial was essential. Earlier killed vaccines had some unkilled virus that lead to disease
- Intense publicity about the vaccine. Trial needed to be done in a single season
- Rate of paralytic polio by region was highly variable.

Key Features of Trial

- Two studies
 - Blinded placebo controlled individually randomized study in 84 areas in 11 states. Children in grades 1-3 randomized.
 - Observational trial 127 areas in 33 states. Children in grade 2 vaccinated grades 1 and 3 received nothing. Helped public support
- Conducted in spring and summer of 1954
 - Enrollment took long---vaccinations into mid June
 - Antibodies measured



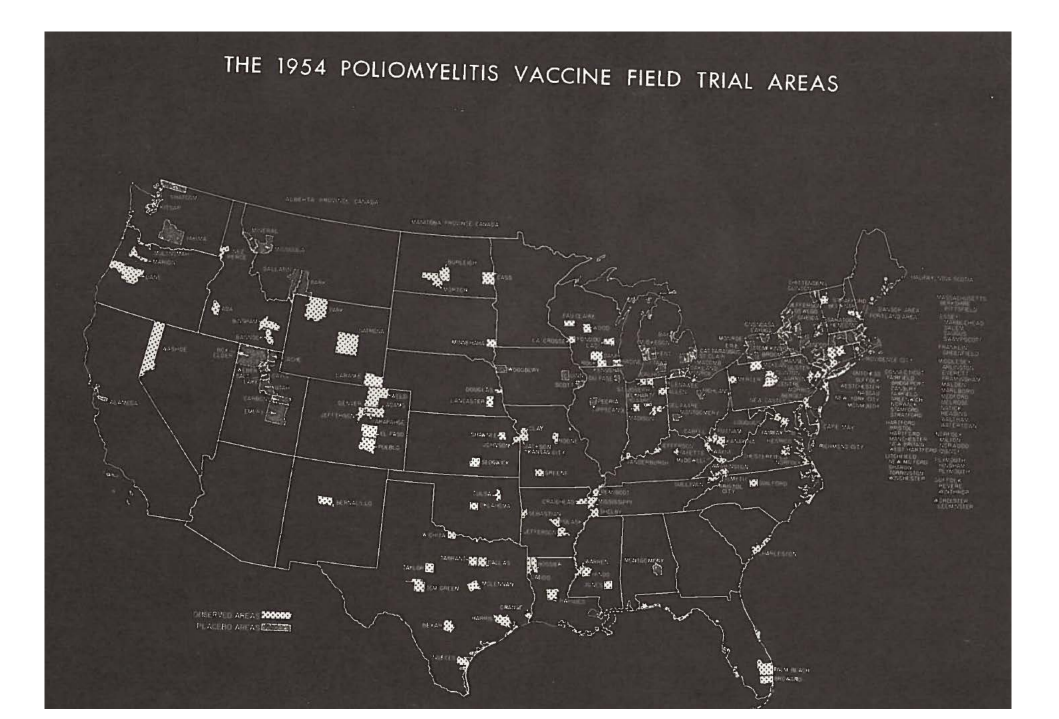




Table 5

DIAGNOSTIC CLASS BY VACCINATION STATUS OF STUDY CASES PLACEBO AND OBSERVED AREAS

	Chin Am	Total Study Cases			Poliomy	Doubtful		Not					
Vaccination Status	Study Population			Total		Paralytic		Nonparalytic		Poliomyelitis		Poliomyelitis	
		Number	Rate	Number	Rate	Number	Rate	Number	Rate	Number	Rate	Number	Rate
All Areas - Total	1, 829, 916	1,012	55	858	47	682	37	176	10	66	4	88	5
Placebo Areas - Total	749, 236	428	57	355	47	267	36	88	12	24	3	49	7
Vaccinated	200, 745	81	40	56	28	33	16	23	11	10	5	15	7
Placebo	201, 229	162	81	138	69	110	55	28	14	7	3	17	8
Incomplete Vaccinations	8,484	2	24	2	24	2	24	-	- 1	-	-	-	-
Incomplete Placebo Injections	8, 577	6	70	6	70	4	47	2	23		-	2 -	-
Not Inoculated	330, 201	177	54	153	46	118	36	35	11	7	2	17	5
Observed Areas - Total	1,080,680	584	54	503	47	415	38	88	8	42	4	39	4
Vaccinated	221, 998	75	34	55	25	38	17	17	8	12	5	8	4
Controls	725, 173	440	61	391	54	331	46	60	8	24	3	25	3
Incomplete Vaccinations	9,904	4	40	4	40	4	40		-		-		-
Second Grade Not Inoculated	123, 605	65	53	53	43	42	34	11	9	6	5	6	5

1-16/55 = .71 Vaccine efficacy

After the 1954 Field Trial

- Cutter incident of Salks inactivated polio vaccine (IPV)
 - One manufacturer didn't properly kill the virus
 - 260 cases were caused: 94 vaccinees, 126 family, 40 community
- Sabin's oral attenuated vaccine (OPV) worked well in Soviet Union
 - Licensed in US 1960
 - Widely used in US 1961-89, simpler & worked better than IPV but
 - Causes paralysis in 1 of 2.9 million vaccinations
- By 2000 US had switched from OPV to IPV



Global Polio Eradication



- Campaign started in 1988, WHO UNICEF & Rotary Foundation, now supported by BMGF & Hutch.
- Afghanistan & Pakistan two remaining countries with endemic polio
 - Challenge: vaccination is a western plot to sterilize
 - Challenge: sham Hep B vaccination campaign used to confirm Osama bin Laden's identity
- Oral polio vaccine (OPV) is highly effective but causes some polio making eradication difficult.
- Plan is to switch from OPV to killed (inactivated) IPV with last wild polio case

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