VIROLOGIA

Viral vaccines



WHY VACCINES MATTER

NORTH KOREA: BEHIND THE CURTAIN

Why Vaccines Matter

- A LIFE OR DEATH SITUATION: While at never of the advance of the very open counteries and the again the information, rack of vaccination availability in poorer nations can be a death sentence.
 - **EXPENSIVE CHALLENGES:** Bangladesh and other countries face challenges like multiple strains of diseases, manufacturing and delivery of vaccines to remote areas.
- **BUSINESS VERSUS HUMANITARIAN INTERESTS:** While poorer countries are in greater need of vaccines, the surest economic return to vaccine manufacturers doesn't come from meeting the most critical need.

VACCINES

THE SEARCH FOR HAPPINESS

GEOGRAPHIC

What we can learn from Costa Rica, Denmark, and Singapore — the most joyful places on the planet

BY DAN BUETTNER

NOVEMBER 2017

Why Vaccines Matter

Here's a way to save hundreds of thousands of young lives: Give children in poor countries the shots that rich countries take for granted.

BY CYNTHIA GORNEY PHOTOGRAPHS BY WILLIAM DANIELS



VACCINATION: OUR PROVEN BEST DEFENSE AGAINST VIRUSES the vaccine concept

- Protects a recipient from a pathogenic agent
- Establishes an immunological resistance to an infection (immune memory)
- Breaks the chain of transmission

VACCINES:

the proven best defense against viruses

Impact of antiviral vaccines on diseases frequency: pre-vaccination era vs. 1998 (USA)

Annual reported cases:							
Disease	Dates	Pre-vaccination	Post-vaccination (1998)	Decrease (%)			
Smallpox	1900–04	48,164	0	100			
Poliomyelitis (paralysis)	1951–54	1,314	1*	100			
Measles	1958–62	503,282	89	100			
Mumps	1968	152,209	606	99.6			
Rubella	1966–68	47,745	345	99.3			
Congenital rubella syndrome	1958–62	823	5	99.4			



Profiles of successful vaccination campaigns



Decline in worldwide measles deaths due to vaccination



with vaccination

Milestones in antiviral vaccine development

Pre-1700s	Chinese doctors use powdered smallpox scabs to "immunize" intranasally. Mediterranean-area doctors use directed leishmania-infected sandfly bites to induce long-term protection from reinfection.
1721	Lady Montagu brings concept of variolation (inoculation with pus from recovering smallpox victim) from Turkey to England.
1798	Jenner publishes Variolae Vaccinae, the use of cowpox inoculation to protect against smallpox.
1885	Pasteur and collaborators introduce air-dried rabbit spinal cord as rabies vaccine.
1900	Walter Reed demonstrates that yellow fever is caused by a filterable virus.
1930–45	Introduction of vaccines for Japanese B encephalitis (1930), yellow fever (1935), and influenza (1936).
1946–75	Introduction of vaccines for polioviruses types 1–3 (Sabin attenuated strains and Salk inactivated virus); measles, mumps, and rubella viruses; tick-borne encephalitis virus; mouse brain, duck embryo, and tissue culture vaccines for rabies virus; inactivated influenza A and B viruses; and adenoviruses.
1975–present	Introduction of vaccines for hepatitis B virus, hepatitis A virus, varicella zoster virus (chickenpox), live, cold- adapted influenza virus, rotavirus, and human papillomavirus.

VACCINES:

the proven best defense against viruses

- Development of vaccines
- Eradicating a viral disease: is it possible?
- The eradication of smallpox (1978)
- The eradication of rinderpest virus (2011)
- What make eradication conceivable?
- The poliovirus case: should be vaccine eradication be next?

Viral vaccines

Smallpox: a Historical Perspective

Smallpox: an historical perspective

Figure 2



A. Smallpox virus Copyright **1994 Veterinary Sciences** Division Queen's University Belfast

Figure 3



A. Edward Jenner



mummified head of Ramses V (died 1157 BCE) with rash that is probably the result of smallpox



B. Dr Jenner about to vaccinate a child



C. Infant with smallpox



D. Smallpox lesions on

skin of trunk. Photo taken in Bangladesh. CDC/James Hicks



Blossom the cow



CDC

F. The last known person in the world to have a natural case of smallpox. Variola minor in 23-year-old Ali Maow Maalin, Merka, Somalia



E. Powdered smallpox scabs were inhaled to protect against smallpox in Chinese medicine

Table 19.1Features of smallpox that enabled itseradication

Virology and disease aspects

No secondary hosts; it is a human-only virus

Long incubation period

Infectious only after incubation period

Low communicability

No persistent infection

Subclinical infections are not a source of spread

Easily diagnosed

Immunology

Infection confers long-term immunity

One stable serotype

Effective vaccine available

Vaccine is stable and cheap

Social political aspects

Severe disease with high morbidity and mortality

Considerable savings to developed, nonendemic countries

Eradication from developed countries demonstrated its feasibility

Few cultural or social barriers to case tracing and control

Viral vaccines

The poliomyelitis eradication:should vaccine eradication be next?



THE ROLE OF INACTIVATED POLIO VIRUS (IPV)

"The next step towards a sustainable polio-free world"









* as of 8 July 2006

Since the momentous launch of the Global Polio Eradication Initiative in 1988 during the World Health Assembly in Geneva, nearly five million children, who otherwise would have been paralyzed and incapacitated by polio, are walking, able and symptoms-free. More than 1.5 million deaths have been prevented.

The number of polio cases reported annually has decreased by 99% from 350,000 in 1988 to 2,000 cases in 2006.

Globally reported incidence of poliomyelitis in 2008. The Americas, Western Pacific, and European regions have been declared poliomyelitis free by the WHO. The number of cases has declined from an estimated 350,000 in 1988 to ca. 1,300 in 2008. At the same time, the number of countries in which poliovirus is endemic has decreased from >125 to 4.







As 2012, Polio remained endemic in four countries – Afghanistan, India, Nigeria and Pakistan – with a further four countries known to have (Angola, Chad and Democratic Republic of the Congo) or suspected of having (Sudan) re-established transmission of poliovirus. Several more countries had outbreaks in 2010 due to importations of poliovirus.





Wild poliovirus type 1 (N=51)
cVDPV1 (N=13)
cVDPV2 (N=2)

¹Excludes viruses detected from environmental surveillance.

Endemic country





¹Excludes viruses detected from environmental surveillance.

Viral vaccines

Vaccination in Italy



D.L. n.73, 7 giugno 2017

Vaccinazioni

I VACCINI, LA MIGLIOR DIFESA PER IL NOSTRO FUTURO

È IN VIGORE IL NUOVO DECRETO VACCINI

PERCHÈ I VACCINI SONO IMPORTANTI?



È IN VIGORE IL **NUOVO DECRETO VACCINI** PER L'ANNO SCOLASTICO 2017/2018. FACCIAMO CHIAREZZA



SCUOLA LE AVVENUTE VACCINAZIONI? Puoi presentare un'autocertificazione per dichiarare le vaccinazioni effettuate entro il **31 ottobre 2017** per la scuola

dell'obbligo o entro il 10 settembre 2017 per i nidi e la scuola dell'infanzia.

Entro il 10 marzo 2018 dovrai presentare copia del libretto delle vaccinazioni timbrato dalla ASL o il certificato vaccinale o un'attestazione dello stato vaccinale rilasciato dalla ASL

SE ALCUNE VACCINAZIONI EFFETTUATE NON DOVESSERO RISULTARE SUL LIBRETTO VACCINALE PERCHÉ FATTE, AD ESEMPIO, DAL MEDICO DI MEDICINA GENERALE. DAL PEDIATRA DI LIBERA SCELTA O PRIVATAMENTE, COSA DEVO FARE?

Devi recarti alla ASL per ottenere la registrazione sul libretto

IMPOSSIBILITÀ A VACCINARSI

SE MIO FIGLIO NON PUÒ VACCINARSI PERCHÉ È MALATO, COSA DEVO FARE?

Se tuo figlio si trova in condizioni di salute che non gli consentono di vaccinarsi in maniera definitiva, devi richiedere al pediatra di libera scelta o al medico di medicina generale di tuo figlio un'attestazione per giustificare la mancata somministrazione. Se invece tuo figlio è malato in modo temporaneo, puoi posticipare la data della vaccinazione fino alla sua quarigione, presentando un'attestazione del pediatra di libera scelta o del medico di medicina generale di tuo figlio.

SE MIO FIGLIO (DA 0 A 6 ANNI) NON HA EFFETTUATO UNA VACCINAZIONE **OBBLIGATORIA** ENTRO IL 10 SETTEMBRE 2017, PUÒ FREQUENTARE L'ASILO NIDO O LA SCUOLA DELL'INFANZIA?

Si, tuo figlio può frequentare regolarmente, purché tu dimostri di aver prenotato la vaccinazione alla ASL, che provvederà ad eseguire la vaccinazione (o ad iniziarne il ciclo, nel caso preveda più dosi) entro la fine dell'anno scolastico.

SE MIO FIGLIO È, INVECE, NELLA FASCIA DI ETÀ TRA 6 E 16 ANNI E NON HA EFFETTUATO UNA DELLE VACCINAZIONI OBBLIGATORIE. POTRÀ FREQUENTARE LA SCUOLA?

Sì, potrà frequentare, ma, sarai contattato dalla ASL per un colloquio informativo. Ove tu non provveda, comunque, a far vaccinare tuo figlio, ti verrà applicata una sanzione pecuniaria.

MENI





VACCINALE





Vaccinazioni

Vaccinazioni obbligatorie da zero a 16 anni

Il decreto vaccini ha aumentato il numero di vaccinazioni obbligatorie per i minori da zero a 16 anni, estendendole da 4 a 10 in base all'anno di nascita. La vaccinazione per la varicella è obbligatoria soltanto per i nati a partire dal 2017. Di seguito una tabella riepilogativa:

Vaccinazione \ Anno	2001	2002	2003	2004	2005	2006	2007	2008	2009	2010	2011	2012	2013	2014	2015	2016	2017
anti- poliomielitica	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	х	Х
anti-difterica	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х
anti-tetanica	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х
anti-epatite B	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	х	Х
anti- pertosse	Х	Х	Х	х	х	Х	Х	Х	Х	Х	Х	х	Х	Х	Х	х	Х
anti- Haemophilus tipo b	х	х	х	х	х	х	х	х	Х	х	х	х	х	х	х	х	х
anti-morbillo	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х
anti-rosolia	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х
anti-parotite	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х	Х
anti-varicella																	Х



1

Vaccinazioni

Il calendario vaccinale del Piano Nazionale di Prevenzione Vaccinale 2017-2019

Vaccino	0gg-30gg	3° mese	4° mese	5° mese	6° mese	7° mese	11° mese	13° mese	15° mese	6° anno	12°-18° anno	19-49 anni	50-64 anni	> 64 anni	Soggetti ad aumentato rischio	
DTPa**		DTPa		DTPa			DTPa			DTPa***	dTpalPV	1 dose	dTpa**** ogni	10 anni	(1)	
IPV		IPV		IPV			IPV			IPV	urpan v					
Epatite B	EpB-EpB*	Ер В		Ер В			Ер В								(2)	
Hib		Hib		Hib			Hib								(3)	
Pneumococco		PCV		PCV			PCV							PCV+PPSV	(4) ^^	
MPRV								MF	PRV	MPRV					(6) ^	
MPR									pure PR	oppure MPR					(5) *****	
Varicella									+ V	+ V					(6)^	
Meningococco C								Me	n C [§]		Men ACWY coniugato				(7)	IPV
Meningococco B*^		Men	B Men	в	Men B			Men B								Ep B Hib
HPV											HPV°: 2-3 funzione di et	3 dosi (in tà e vaccino)			(8)	DTPa
Influenza														1 dose all'anno	(9) °°	dTpa dTpa-IPV
Herpes Zoster														1 dose#	(10)	MPRV
Rotavirus					o tre dos di vaccin											MPR V
Epatite A															(11)	PCV

Co-somministrare nella stessa seduta
Somministrare in seduta separata
Vaccini per categorie a rischio

IPV	=	vaccino antipolio inattivato
Ер В	=	vaccino contro il virus dell'epatite B
Hib	=	vaccino contro le infezioni invasive da Haemophilus influenzae tipo b
DTPa	=	vaccino antidifterite-tetano-pertosse acellulare
dTpa	=	vaccino antidifterite-tetano-pertosse acellulare, formulazione per adulti
dTpa-IP	v =	vaccino antidifterite-tetano-pertosse acellulare e polio inattivato,
		formulazione per adulti
MPRV	=	vaccino tetravalente per morbillo, parotite, rosolia e varicella
MPR	=	vaccino trivalente per morbillo, parotite, rosolia
v	=	vaccino contro la varicella
PCV	=	vaccino pneumococcico coniugato
PPSV	=	vaccino pneumococcico polisaccaridico
MenC	=	vaccino contro il meningococco C coniugato
MenB	=	vaccino contro il meningococco B
HPV	=	vaccino contro i papillomavirus
Influenza	a =	vaccino contro l'influenza stagionale
Rotaviru	IS =	vaccino contro i rotavirus

Viral vaccines

Vaccine Basics

VACCINATION: basis for protective immunity



NATURAL PASSIVE IMMUNIZATION: transfer of antibody from mother to infant



VACCINATION: vaccines can be *active* or *passive*

- Active instilling into the recipient a modified form of the pathogen or material derived from it that induces immunity to disease (long term protection).
- Passive instilling the products of the immune response (antibodies or immune cells) into the recipient (short term protection)

Zmapp, the best known passive vaccine







Then a gene is removed from the new antibody...





leaves make large quantities ZMAPP of the drug

- Raised in mice immunized with virus-like particles •
- Chimerized into human IgG1 scaffold •
- Produced in tobacco plants •

VACCINATION:

requirements for an effective vaccine

Safety

Side effects must be minimal Induction of protective immune response

Practical issues

The vaccine must not cause disease

- Vaccinated individuals must be protected from illness due to pathogen
- Proper innate, cellular, and humoral responses must be evoked by vaccine
- Cost per dose must not be prohibitive
- The vaccine should be biologically stable (no genetic reversion to virulence; able to survive use and storage in different surroundings)
- The vaccine should be easy to administer (oral delivery preferred to needles)
- The public must see more benefit than risk

THE HERD IMMUNITY CONCEPT



Herd immunity threshold (HT): $1-1/R_0$ R₀ reproduction number R₀ smallpox: 5-7 HT: 80-85% R₀ measle: 12-18 HT: 93-95%



THE HERD IMMUNITY CONCEPT

The correlation between herd immunity and the potential for outbreaks.

As the % of vaccinated individuals dip below 90%, a corresponding rise in the number of acute cases is observed



When 80% of population is immunized with measles, 76% of population is immune

VACCINATION: viral vaccines in the USA

Disease or virus	Type of vaccine	Indications for use	Schedule
Adenovirus	Live attenuated, oral	Military recruits	One dose
Hepatitis A	Inactivated whole virus	Travellers, other high-risk groups	0, 1, and 6 mo
Hepatitis B	Yeast-produced recombinant surface protein	Universal in children, exposure to blood, sexual promiscuity	0, 1, 6, and 12 mo
Influenza	Inactivated viral subunits	Elderly and other high-risk groups	One dose seasonally
Influenza	Live attenuated	Children 2–8 yr old, not previously vaccinated with influenza vaccine	Two doses at least 1 mo apart
		Children 2–8 yr old, previously vaccinated with influenza vaccine	One dose
		Children, adolescents, and adults 9-49 yr old	One dose
Japanese encephalitis	Inactivated whole virus	Travelers to or inhabitants of high-risk areas in Asia	0, 7, and 30 days
Measles	Live attenuated	Universal vaccination of infants	12 mo of age; 2nd dose, 6 to 12 yr of age
Mumps	Live attenuated	Universal vaccination of infants	Same as measles, given as MMR
Papilloma (human)	Yeast- or SF9-produced virus-like particles	Females 9–26 yr old	Three doses
Rotavirus	Live reassortant	Healthy infants	2, 3, and 6 mo or 2 and 4 mo of age depending on vaccine
Rubella	Live attenuated	Universal vaccination of infants	Same as measles, given as MMR
Polio (inactivated)	Inactivated whole viruses of types 1, 2, and 3	Changing: commonly used for immunosuppressed where live vaccine cannot be used	2, 4, and 12–18 mo of age, then 4 to 6 yr of age
Polio (live)	Live, attenuated, oral mixture of types 1, 2, and 3	Universal vaccination; no longer used in United States	2, 4, and 6–18 mo of age
Rabies	Inactivated whole virus	Exposure to rabies, actual or prospective	0, 3, 7, 14, and 28 days postexposure
Smallpox	Live vaccinia virus	Certain laboratory workers	One dose
Varicella	Live attenuated	Universal vaccination of infants	12 to 18 mo of age
Varicella-zoster	Live attenuated	Adults 60 yr old and older	One dose
Yellow fever	Live attenuated	Travel to areas where infection is common	

VACCINATION: how to make a vaccine?



VACCINATION: traditional vaccines

Traditional vaccines are of two types:

- Inactivated (killed) (whole cells, viruses, inactivated toxin proteins (toxoids)
- Attenuated (live) -viral or bacterial strain

VACCINATION:

commonly used traditional antiviral vaccines

- Attenuated (replication competent) vaccines
 - Poliomyelitis
 - •Mumps
 - Measles
 - Rubella
 - Yellow fever
 - Varicella
- Inactivated vaccines
 - Influenza
 - Poliomyelitis
 - Rabies


Inactivated influenza vaccine

- >7000 deaths/yr in Italy due to influenza virus
- Vaccine: virus grown in embryonated chicken eggs, formalin-inactivated or detergent or chemically disrupted virions.
- Hundreds million of doses manufactered each year.
- 60% effective in healthy children and adults <65 yr.
- Protection correlates with antibodies to HA and NA.
- Envelope proteins change each year; new strains must be selected in the first months for manifacture.
- Use reassortants with most RNA segments from high-yielding strain, HA, NA from selected strain.

HA antigenic drift: influenza virus

Annual timeline for creating an influenza virus vaccine



*World Health Organization Global Influenza Surveillance Network *WHO Collaborating Centres

[‡]US Centers for Disease Control and Prevention [§]US Food and Drug Administration







26 febbraio 2016

Raccomandazioni dell'OMS per la composizione del vaccino antinfluenzale per la stagione 2016-2017 (Emisfero Nord)

Dal 22 al 24 febbraio 2016 si è svolto a Ginevra il *meeting* annuale dell'OMS per l'aggiornamento della composizione del vaccino antinfluenzale per la stagione 2016/2017.

Le raccomandazioni emanate sono il risultato dei dati di sorveglianza virologica forniti da tutti i Centri Nazionali di riferimento (NIC), afferenti alla rete internazionale dell'OMS, attualmente composta da oltre 140 laboratori.

Qui di seguito viene riportata la nuova composizione vaccinale 2016/2017:

A/California/7/2009 (H1N1)	Presente anche nel vaccino 2015/2016
A/Hong Kong/4801/2014 (H3N2)	Nuova variante
B/Brisbane/60/2008 (lineaggio B/Victoria)	Nuova variante

VACCINATION: comparison of the predicted immune response to live and killed viruses used as vaccines





Comparison of immune responses to live and killed viruses

VACCINATION: problems with traditional vaccines

- Disadvantages of inactivated vaccines:
 - they themselselves can cause severe reactions.
 - the organism or the toxin may not be completlely killed or inactivated.

- in some cases producing a sufficient quantity of infectious agent is extremerly costly or even impossible (HBV, HPV)

Severe adverse events associated with antiviral vaccine administration

Vaccine implicated	Event
Smallpox ("lymph" vaccines)	Sepsis due to bacterial contamination
Yellow fever (1942)	Vaccine lot contaminated with hepatitis B virus, leading to 28,000 cases of hepatitis B
Inactivated polio (1955)	Incomplete inactivation of virus, leading to 204 cases of paralytic disease (Cutter incident)
Inactivated measles (1960s)	Atypical (severe) disease upon exposure to natural measles infection
Attenuated polio (Sabin)	Vaccine-associated paralysis (~1 per 2.4 million persons vaccinated) due to reversion to more pathogenic strain
Live vaccines (vaccinia, polio, measles)	Dissemination and death in immunocompromised individuals
Attenuated live measles	Unexplained mortality in girls who received high titer formulations (relative risk of death doubled to age 5); more than 20 million doses of vaccine distributed
Inactivated influenza A	Apparent risk of Guillain Barré Syndrome (~1 per 1 million persons vaccinated), only in some years
Rotavirus (1999)	Intussusception—bowel folding and obstruction (vaccine withdrawn)

VACCINATION:

producing attenuated human viruses



Passage history of the Oka-strain VZV vaccine

Isolated by Takahashi et al. in 1974 from three-year-old Japanese boy (family name Oka)

Initial growth in human lung fibroblast cells

Twelve passages in primary guinea pig fibroblasts

Two passages in WI-38 cells (human diploid fibroblasts)

Three to six passages in MRC-5 cells (human diploid lung cells)

Licensed for use in Japan and Korea in late 1980s, and in USA in 1995

Live attenuated vaccines: passage histories of live attenuated measles virus vaccines derived from original isolate of Edmonston



VACCINATION: live attenuated oral poliovirus vaccines

A Derivation of Sabin type 3 attenuated poliovirus

Type 3 P3/Leon/37 (isolate from fatal paralytic case)

21 passages in vivo (intracerebrally in monkeys)
8 passages in vitro (monkey testicle cultures)
39 passages in vitro (monkey kidney cultures)
3 plaque purifications (monkey kidney cultures)
3 passages in vitro (preparative, monkey kidney cultures)

P3/Leon 12a1b KP3/56 Sabin vaccine strain

B Determinants of attenuation in the Sabin vaccine strains

Virus	Mutation (location/nucleotide position)
P1/Sabin	5'-UTR (480) VPI (1106) VPI (1134) VP3 (3225) VP4 (4065)
P2/Sabin	5'-UTR (481) VPI (1143)
P3/Sabin	5'-UTR (472) VP3 (3091)

VACCINATION: live attenuated oral poliovirus vaccines



Attenuation of poliovirus neurovirulence: the Sabins9 OPV licensed in 1961

VACCINATION: problems with traditional vaccines

- Disadvantages of attenuated vaccines:
 - reversion to the virulent state (polio).

- growth in tissue culture cells or in animals poses the risk of introducing hidden viruses from host cells (early lots of polio vaccines-SV40).

- even attenuated pathogens can produce severe disease in people with immune system deficiences.

VACCINATION: reversion of P3/sabin (1/790.000)

C Reversion of P3/Sabin



Virus strain	Number of mutations	Number of amino acid changes	Number of mutations required for return to neurovirulence
Sabin strain 1	57	23	>10
Sabin strain 2	23	5	5-6
Sabin strain 3	~6	3	1–2

Limitations to Current Vaccine Production Methods

- Not all pathogens can be grown in culture
- Cell culture is expensive
- Yield and rate of production may be low
- Extensive safety precautions needed
- Inactivation/attenuation must be 100%
- Attenuated strains might revert
- Limited shelf-life and refrigeration requirements
- Not all diseases preventable by vaccines

NEW VACCINE TECHNOLOGY: Impact of rDNA technology approach to vaccine production

- 1. Attenuate organism by specific deletion of virulence gene (bacmid technology)
- Engineer live, non pathogenic carrier (vector)to contain antigenic determinant for disease organism (viral vector technology)
- 1. If non-culturable, clone pathogen's genes, and overexpress a subset of viral proteins to use as vaccines (subunit vaccines, VLPs)

VACCINATION: how to make a vaccine?



VACCINATION: how to make a vaccine?



Recombinant antiviral vaccines

MODERNS VACCINES

- Subunit Vaccines
- Peptide Vaccines
- Genetic Immunization
- Attenuated Vaccines
- Vector Vaccines

MODERNS VACCINES

Subunit Vaccines

- Peptide Vaccines
- Genetic Immunization
- Attenuated Vaccines
- Vector Vaccines

Subunit Vaccines

- Generally whole pathogenic agent used to construct attenuated or inactivated vaccine
- Immune response generally elicited by interaction with proteins on outer surface of pathogen



Subunit vaccines for viruses

В



Subunit Vaccine

• So, is the entire pathogen required?

- No, only outer surface proteins are needed to elicit an immune response
- Vaccines that use components of a pathogen rather than the whole organism are "subunit" vaccines

Subunit Vaccine

Advantages

- Easily to produce
- Using a purified protein ensures that the vaccine is safe and stable
- Inexpensively
- Disadvantages
 - Isolated protein may not have the same conformation as in the pathogen, so may not have the same antigenicity (weakness and short duration of immunity)
 - It may also be necessary to administer the antigen in a specific manner (in a concentrated form, with adjuvants















Negative stained electron micrographs of (A) plasma-derived and (B) yeast-derived hepatitis B surface antigen vaccines

RECOMBIVAX HB® HEPATITIS B VACCINE (RECOMBINANT)

DESCRIPTION

RECOMBIVAX HB* Hepatitis B Vaccine (Recombinant) is a non-infectious subunit viral vaccine derived from hepatitis B surface antigen (HBsAg) produced in yeast cells. A portion of the hepatitis B virus gene, coding for HBsAg, is cloned into yeast, and the vaccine for hepatitis B is produced from cultures of this recombinant yeast strain according to methods developed in the Merck Research Laboratories.

The antigen is harvested and purified from fermentation cultures of a recombinant strain of the yeast *Saccharomyces cerevisiae* containing the gene for the *adw* subtype of HBsAg. The fermentation process involves growth of *Saccharomyces cerevisiae* on a complex fermentation medium which consists of an extract of yeast, soy peptone, dextrose, amino acids and mineral salts. The HBsAg protein is released from the yeast cells by cell disruption and purified by a series of physical and chemical methods. The purified protein is treated in phosphate buffer with formaldehyde and then coprecipitated with alum (potassium aluminum sulfate) to form bulk vaccine adjuvanted with amorphous aluminum hydroxyphosphate sulfate. The vaccine contains no detectable yeast DNA but may contain not more than 1% yeast protein. The vaccine in terms of animal potency (mouse, monkey, and chimpanzee) and protective efficacy (chimpanzee and human).

The vaccine against hepatitis B, prepared from recombinant yeast cultures, is free of association with human blood or blood products.





PRESCRIBING INFORMATION



ENGERIX-B[®] [Hepatitis B Vaccine (Recombinant)]

DESCRIPTION

ENGERIX-B [Hepatitis B Vaccine (Recombinant)] is a noninfectious recombinant DNA hepatitis B vaccine developed and manufactured by GlaxoSmithKline Biologicals. It contains purified surface antigen of the virus obtained by culturing genetically engineered *Saccharomyces cerevisiae* cells, which carry the surface antigen gene of the hepatitis B virus. The surface antigen expressed in *Saccharomyces cerevisiae* cells is purified by several physicochemical steps and formulated as a suspension of the antigen adsorbed on aluminum hydroxide. The procedures used to manufacture ENGERIX-B result in a product that contains no more than 5% yeast protein. No substances of human origin are used in its manufacture.

ENGERIX-B is supplied as a sterile suspension for intramuscular administration. The vaccine is ready for use without reconstitution; it must be shaken before administration since a fine white deposit with a clear colorless supernatant may form on storage.

7994332

Virus-like particles (VLPs) vaccines: HPV



Virus-like particles (VLPs) vaccines: HPV





Virus-like particles (VLPs)vaccines: HPV







VLP



Virus-like particles (VLPs)vaccines: HPV

HPV VLPs vaccine development: *timeline*



The failure of a subunit Vaccine - HSV

- Herpes Simplex Viruses (HSV-1,2)
 - sexually transmitted disease (HSV-2)
 - cancer associated agent (HSV-2)
 - encephalitis and severe eye infections

Subunit vaccine would be best

 inactivated or attenuated virus would have
 to be 100% or risk infecting patient

Subunit Vaccine - Example: HSV Target Antigenic Protein

Herpes Simplex Virus

 What is the antigenic target?

HSV viral envelope glycoprotein D
gD interacts with entry receptors (HVEM, nectin-1, 3-O-sulphated HS)

Structure and function of HSV gD glycoprotein



3) endo-domain: aa 340-369
Functions of HSV gD: receptor recognition

The three natural gD receptors:

 herpesvirus entry mediator (HVME): tumor necrosis factor receptor family; in T-lymphocytes or lymphoyd organ. HVEM binding-site: aa 1-32 (contact residues between aa 7-15 and 24-32).

2. nectin 1:

intercellular adhesion molecules family; in sensory neurons, muco-epithelia or epithelia cells. nectin-1 binding-site: critical aa residues (aa 34, 38, 215 and aa 222-223).

 O-sulphated HS (heparan sulfate): modified heparan sulfate by enzymes in neuronal and endothelial cells, corneal fibroblasts.

HSV Glycoprotein D



HSV glycoprotein D vaccine

- Isolate gene
- Clone into mammalian exp. system
- Purify expressed protein

 Membrane proteins typically difficult to purify - aggregate

– Mutate so not membrane bound

A Mammalian expression vector

Comments for pcDNA3.1/V5-His A

5503 nucleotides

CMV promoter: bases 209-863 T7 promoter/priming site: bases 863-882 Multiple cloning site: bases 902-999 V5 epitope: bases 1000-1041 Polyhistidine tag: bases 1051-1068 pcDNA3.1/BGH reverse priming site: bases 1091-1108 BGH polyadenylation signal: bases 1090-1304 f1 origin of replication: bases 1357-1780 SV40 promoter and origin: bases 1845-2170 Neomycin resistance gene: bases 2206-3000 SV40 polyadenylation signal: bases 3019-3257 ColE1 origin: bases 3689-4362 Ampicillin resistance gene: bases 4507-5367

> * After the Xho I site, there is a unique BstE II site, but no Xba I or Apa I sites in version C.

** There is a unique Sac II site between the Apa I site and the Sfu I site in version B only.



Modified HSV glycoprotein D



Development of Subunit Vaccine against HSV



HSV glycoprotein D vaccine

- HSV Subunit vaccine production
 - Transfect modified glycoprotein gene into mammalian cells
 - -Overexpress
 - -Purify
- Modified HSV glycoprotein D was effective against both HSV-1 and HSV-2 in animal lab tests

HSV glycoprotein D vaccine

Table 4: Summary of human trials of prophylactic herpes simplex virus (HSV) vaccines

	Vaccine	Manufacturer				
Live-attenuated/replication disabled						
	R7020	Pasteur Merieux ² *				
	ICP10DPK	AuRx Inc.32				
Killed/viral component						
	Skinner ⁷⁻⁹					
	Cappel et al.14					
	HSV-2 GS	Merck, Sharpe & Dohme ¹⁶				
	Biocine	Chiron ⁶⁰ *				
	gD2/gB2/ MF59	Chiron ³⁵				
	Simplirix (gD2/MPL-alum)	GlaxoSmithKline ³				
*Trial referred to within this reference.						



Efficacy Results of a Trial of a Herpes Simplex Vaccine

Robert B. Belshe, M.D., Peter A. Leone, M.D., David I. Bernstein, M.D., Anna Wald, M.D., Myron J. Levin, M.D., Jack T. Stapleton, M.D., Iris Gorfinkel, M.D., Rhoda L. Ashley Morrow, Ph.D., Marian G. Ewell, Sc.D., Abbie Stokes-Riner, Ph.D., Gary Dubin, M.D., Thomas C. Heineman, M.D., Ph.D., Joann M. Schulte, D.O., and Carolyn D. Deal, Ph.D., for the Herpevac Trial for Women

ABSTRACT

BACKGROUND

Two previous studies of a herpes simplex virus type 2 (HSV-2) subunit vaccine containing glycoprotein D in HSV-discordant couples revealed 73% and 74% efficacy against genital disease in women who were negative for both HSV type 1 (HSV-1) and HSV-2 antibodies. Efficacy was not observed in men or HSV-1 seropositive women.

METHODS

We conducted a randomized, double-blind efficacy field trial involving 8323 women 18 to 30 years of age who were negative for antibodies to HSV-1 and HSV-2. At months 0, 1, and 6, some subjects received the investigational vaccine, consisting of 20 μ g of glycoprotein D from HSV-2 with alum and 3-0-deacylated monophosphoryl lipid A as an adjuvant; control subjects received the hepatitis A vaccine, at a dose of 720 enzyme-linked immunosorbent assay (ELISA) units. The primary end point was occurrence of genital herpes disease due to either HSV-1 or HSV-2 from month 2 (1 month after dose 2) through month 20.

RESULTS

The HSV vaccine was associated with an increased risk of local reactions as compared with the control vaccine, and it elicited ELISA and neutralizing antibodies to HSV-2. Overall, the vaccine was not efficacious; vaccine efficacy was 20% (95% confidence interval [CI], -29 to 50) against genital herpes disease. However, efficacy against HSV-1 genital disease was 58% (95% CI, 12 to 80). Vaccine efficacy against HSV-1 infection (with or without disease) was 35% (95% CI, 13 to 52), but efficacy against HSV-2 infection was not observed (~8%; 95% CI, -59 to 26).

CONCLUSIONS

In a study population that was representative of the general population of HSV-1and HSV-2-seronegative women, the investigational vaccine was effective in preventing HSV-1 genital disease and infection but not in preventing HSV-2 disease or infection. (Funded by the National Institute of Allergy and Infectious Diseases and GlaxoSmithKline; ClinicalTrials.gov number, NCT00057330.)



MODERNS VACCINES

- Subunit Vaccines
- Peptide Vaccines
- Genetic Immunization
- Attenuated Vaccines
- Vector Vaccines

Peptide Vaccines

- Subunit vaccine uses entire protein

 Contains several antigenic determinants (B and T cells epitopes)
- Peptide Vaccine
 - vaccine from a specific domain of an antigenic protein
 - single epitope or antigenic determinant (immunogenic epitope)

Generalized membrane-bound protein with external epitopes



Peptide Vaccines

• Advantages of peptide vaccines:

- can be made by chemical synthesis and eliminate the need of expensive process of protein purification (remove host DNA,LPS).

- they are less expensive, purer and more stable than protein-containing subunit vaccines.

-in addition, using only a part of the antigenic protein there will be no unwanted immunological reactions.

Peptide Vaccines

Peptides need to be linked to another molecule to prevent rapid degradation

Keyhole limpet hemocyanin

- an inert carrier protein
- from a marine gastropos mollusk
- Hepatitis B core protein (HBcAg)
 - highly immunogenic carrier protein
 - self assembles into small particles

Keyhole Limpet (Diodora aspera)



Structure of a peptide vaccine



Limitations of Peptide Vaccines

- Epitope must consist of a contiguous stretch of amino acids
- Not all peptides are effective in eliciting an immune response (may need 2 or more)
- Peptide must have the same conformation as in pathogen
- Selection of immune escape mutants is highly probable (may need 2 or more)
- Amount of peptide required to elicit an immune response may be 1000X more than for inactivated pathogen

Peptide Vaccine Effectiveness Foot and Mouth Disease Virus (FMDV)









FMDV peptide vaccines





Peptide Vaccine Effectiveness - Example FMVD

Foot and Mouth Disease peptide vaccines

FMDV particles	1
VP1-HBcAg particles VP1=142-160 aa	1/10
VP1-β-gal fusion VP1=137-162 aa	1/350
VP1 142-160 aa	1/5000

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GENETIC IMMUNIZATION

- Novel variation of recombinant vaccine strategy
 - First reported in 1992
 - Instead of injecting protein antigen to elicit immune response, inject gene encoding antigen
- Also called "DNA Vaccine"

- Gene directs production of encoded protein antigen directly into tissues of the vaccinee
- A relative low dose of immunizing protein (ng range) seems to be sufficient to produce long-lasting immune response
- Antigen activates an immune response
 - Antibody production
 - Cell mediated response

- The method of inoculation appears to dictate the type of immune response:
 - Direct Injection into muscle of an aqueous DNA solution (DNA travels to spleen Th1 response)
 - Biolistic method DNA coated metal beads (keratinocytes express ag, Langerhans cells move to the draining lymph node – Th2 response)

- Advantages of genetic immunization
 - Cultivation of pathogen not required
 - No chance of reversion
 - No adjuvants are necessary to stimulate an immune response
 - Inexpensive no need to purify protein
 - Inexpensive purified DNA is stable
 - One plasmid could encode several antigens





- Disadvantage
 - Don't know fate of DNA
 - Transient will repeated treatments be needed?
 - Integration into genome mutations?

Representative results of DNA vaccine trials

Virus	Proteins	Induction of antibody	Induction of CTL response	Protection against challenge
Bovine herpesvirus	gD	+	ND	+ (cattle)
Hepatitis B virus	Surface and core antigens	+ (chimpanzees); ND (humans)	+ (chimpanzees)	+ (chimpanzees)
Hepatitis C virus	Nucleocapsid	+	+	+ (mice)
Herpes simplex virus type 1	gD, gB	+	+	+ (mice)
HIV type 1	Env, Gag, Rev	+	+	+ (rhesus macaques)
Influenza virus	HA, M1, Np	+	+	+ (chickens, mice)
Lymphocytic choriomeningitis virus	NP	+	+	+ (mice)
Rabies virus	Glycoprotein, NP	+	+	+ (cynomolgus monkeys)
Respiratory syncytial virus	Glycoprotein	+	+	+ (mice)

"Data from A. Reyes-Sandoval and H. C. Ertl, Curr. Mol. Med. 1:217-243, 2001, with permission. ND, not detected.

Genetic Immunization/ DNA Vaccine Example Influenza Virus

Inject mice with plasmid Gene influenza virus nucleocapsid NP coding sequences Promoter Cytomegalovirus (CMV) promoter **Injection site** quadriceps muscles of both rear legs

Genetic Immunization of Mice Against Influenza Virus

- Couldn't detect expression of protein
- Could detect Abs to flu virus nucleoprotein
- Injected mice were protected from Influenza virus (Iv)

Survival of DNA-immunized Mice



The immunostimulatory sequences of plasmid vector DNA have adjuvant effects:

•Bacterial DNA has intrinsic immunostimulatory activity in mammalian cells through TLR9 stimulation.

•DNA motifis containing unmethylated CpG dinucleotides flanked by two 5' purines and by two 3' pyrimidines (5'-GACGTC-3')induce synthesis of IL6, IFN-γ, IL12, TNF.

•CpG DNA as a PAMP such as LPS, PGN, LTA dsRNA

MODERNS VACCINES

- Subunit Vaccines
- Peptide Vaccines
- Genetic Immunization
- Attenuated Vaccines
- Vector Vaccines

ATTENUATED VACCINES: a modern approach

- Attenuated vaccines
 - generally more effective vaccines
 - BUT must not have virulent forms present
- rDNA techniques permit genetic manipulation for generating attenuated vaccines

rDNA Technology Based Approach for Modifying Organisms to Generate Attenuated Vaccines

- A non-pathogenic organism engineered to carry and express antigenic determinants from a target pathogenic agent (vector vaccines)
- A pathogenic organism engineered such that the virulence genes have been modified or deleted (Bacmid technology)

Benefits of Attenuated Vaccines generated by rDNA technology

- Only a selected antigenic determinant from the pathogen is put into the non-pathogen, so there is no chance of vaccine causing disease
- Deletion of virulence genes greatly decreases the likelihood of reversion back to the virulent form
- Whole pathogen much more immunogenic than subunit or peptide vaccines (concentrated forms, in the presence of PAMPs)
Isolate pathogenic virus

Mutate

gene

Construction of attenuated Viruses by using recombinant **DNA** tecnology



Estimated Global Rotavirus Deaths in 2008



*Tate JE, Burton AH, Boschi-Pinto C, Duncan Steele D, et. al. 2008 estimate of worldwide rotavirus-associated mortality in children younger than 5 years before the introduction of universal rotavirus vaccination programmes: a systematic review and meta-analysis. *The Lancet*. Published online October 25, 2011.

Rotaviruses are the leading cause of severe diarrhea among infants and young children. Each year an estimated 453,000 children die from diarrhoeal disease caused by Rotavirus, most of whom live in developing countries, and another two million are hospitalised.

Rotavirus pathogenesis

Reoviruses				
Rotavirus				
Virus	Disease	Epidemiology		
Orthoreovirus	Mild upper respiratory tract disease, gastroenteritis, biliary atresia	Transmission • Fecal-oral route	Distribution of virus • Ubiquitous (type A) • Less common in summer	
Orbivirus/coltivirus	Colorado tick fever: febrile disease, headache, myalgia (zoonosis)	At risk • Rotavirus type A Infants <24 months of age (gastroenteritis, dehydration)	Vaccines or antiviral drugs • None	
Rotavirus	Gastroenteritis	Older children (mild diarrhea) Undernourished persons in underdeveloped countries (diarrhea, dehydration, death)		
		 Rotavirus type B Infants, older children, adults in China (severe gastroenteritis) 		

Disease mechanisms

Transmitted by fecal-oral route

nsP4 is a viral enterotoxin that causes diarrhea

Disease is serious in infants <24 months old, asymptomatic in adults

Large quantities of virions released in diarrhea

Immunity to infection depends on IgA in gut lumen

Rotavirus is highly contagious and resistant and, regardless of water quality and available sanitation nearly every child in the world is at risk of infection.



The mechanisms of Rotavirus pathogenesis and immunity



Nature Reviews | Microbiology

Schematic representation of a rotavirus virion.



Rotavirus vaccine

- Rotaviruses contain 11 segments of double-stranded RNA.
- Rotaviruses are classified into seven groups (A to G) on the basis of their distinct antigenic and genetic properties. Human infection has been reported with group A, B, and C rotaviruses. Of these, group A rotavirus is the most important, being a significant cause of severe gastroenteritis in children worldwide.
- The two outer capsid proteins, VP4 and VP7, allow classification of rotavirus into P and G genotypes, respectively. In rotavirus, at least 15 G genotypes have been recognized by neutralization assay and 27 P genotypes have been identified by hybridization or sequence analysis. Of these, four rotavirus G-P combinations, i.e., G1P[8], G2P[4], G3P[8], G4P[8], and G9P[8], are the most common globally and are therefore the targets for current vaccine development strategies.
- Since effective antirotavirus drugs have not been developed, a rotavirus vaccine would be very useful.

Rotavirus reassortant to generate oral live virus vaccine





a) **Rotarix** is an attenuated human rotavirus vaccine made of a tissue-culture-adapted human P1A[8]G1, VP6 subgroup II and NSP4 geno-group B strain.

b) **RotaTeq** is a bovine (WC3)-human reassortant vaccin. RotaTeq is a live, oral pentavalent vaccine that contains five rotaviruses produced by reassortment. The rotavirus A parent strains of the reassortants were isolated from human and bovine hosts. Four reassortant rotaviruses express one of the outer capsid, VP7, proteins (serotypes G1, G2, G3, or G4) from the human rotavirus parent strain and the attachment protein VP4 (type P7[5]) from the bovine rotavirus parent strain. The fifth reassortant virus expresses the attachment protein VP4, (type P1[8]), from the human rotavirus parent strain and the outer capsid protein VP7 (serotype G6) from the bovine rotavirus parent strain.



RotaTeq is a bovine (WC3)-human reassortant vaccin. RotaTeq is a live, oral pentavalent vaccine that contains five rotaviruses produced by reassortment. Four reassortant rotaviruses express one of the outer capsid, VP7, proteins (serotypes G1, G2, G3, or G4) from the human rotavirus parent strain and the attachment protein VP4 (type P7[5]) from the bovine rotavirus parent strain. The fifth reassortant virus expresses the attachment protein VP4, (type P1[8]), from the human rotavirus parent strain and the outer capsid protein VP7 (serotype G6) from the bovine rotavirus parent strain.



RotaTeq[®] [Rotavirus Vaccine, Live, Oral, Pentavalent]

	Human Rotavirus Parent Strains	Bovine Rotavirus Parent	Reassortant Outer Surface Protein Composition	
Name of	and Outer Surface Protein	Strain and Outer Surface	(Human Rotavirus Component	Minimum Dose Levels
Reassortant	Compositions	Protein Composition	in Bold)	(10 ⁶ infectious units)
G1	WI79 – G1, P1[8]		G1 , P7[5]	2.2
G2	SC2 – G2, P2[6]		G2 , P7[5]	2.8
G3	WI78 – G3, P1[8]	WC3 - G6, P7[5]	G3 , P7[5]	2.2
G4	BrB – G4, P2[6]		G4 , P7[5]	2.0
P1[8]	WI79 – G1, P1[8]		G6, P1[8]	2.3

DESCRIPTION

RotaTeq^{*} is a live, oral pentavalent vaccine that contains 5 live reassortant rotaviruses. The rotavirus parent strains of the reassortants were isolated from human and bovine hosts. Four reassortant rotaviruses express one of the outer capsid proteins (G1, G2, G3, or G4) from the human rotavirus parent strain and the attachment protein (P7) from the bovine rotavirus parent strain. The fifth reassortant virus expresses the attachment protein, P1A (genotype P[8]), hereafter referred to as P1[8], from the human rotavirus parent strain and the outer capsid protein G6 from the bovine rotavirus parent strain (see Table 1).

Overview of the development of a vaccine against Rotavirus (Rotateq)

Overview of development of RotaTeq



As of September 2012, 41 Countries Have Introduced Rotavirus Vaccines



World Health Organization, 2012 PATH, 2012

MODERNS VACCINES

- Subunit Vaccines
- Peptide Vaccines
- Genetic Immunization
- Attenuated Vaccines
- Vector Vaccines, see. Viral vectors

Some antiviral vaccines currently under development

Virus family	Virus species	Most promising strategies
Arenavirus	Junin	Live attenuated
	Lassa fever	Recombinant vaccinia virus
Calicivirus	Norwalk	Recombinant capsid "ghosts"
Filovirus	Ebola/Marburg	DNA
		Recombinant vesicular stomatitis virus
Flavivirus	Hepatitis C	Recombinant envelope glycoprotein, peptide, polypeptide
	Dengue	Tetravalent attenuated, DNA
Hepadnavirus	Hepatitis B	DNA, recombinant surface antigen including pre-S1/S2 regions; curative hepatitis B vaccine: DNA with CpG motifs
Herpesvirus	Epstein-Barr	Whole inactivated, subunit (gp350)
	Herpes simplex 1, 2	Recombinant polypeptide, vectored
Orthomyxovirus	Influenza A	Vectored, DNA, peptide
Paramyxovirus	Measles	Vectored, DNA
	Parainfluenza	Live attenuated
	Respiratory syncytial	Subunit G and fusion protein, DNA, vectored, peptide
Parvovirus	Parvovirus B19	Empty capsid with VP1 protein
Picornavirus	Enterovirus 71	Whole inactivated
Reovirus	Rotavirus	Live reassortants of simian or human origin
Retrovirus	HIV-1, 2	Recombinant polypetide, DNA, vectored, chimeric
Unclassified	Hepatitis E	Recombinant protein ORF2