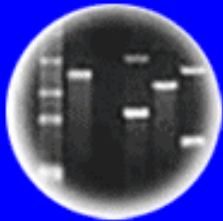


DNA  
Genetic Code of Life



Entire Genetic Code  
of a Bacteria



DNA Fingerprinting



Cloning: Ethical Issues  
and Future Consequences



Plants of Tomorrow

# HC70A & SAS70A Spring 2015 Genetic Engineering in Medicine, Agriculture, and Law

Professors Bob Goldberg. John Harada,  
& Channapatna Prakash  
Lecture 6

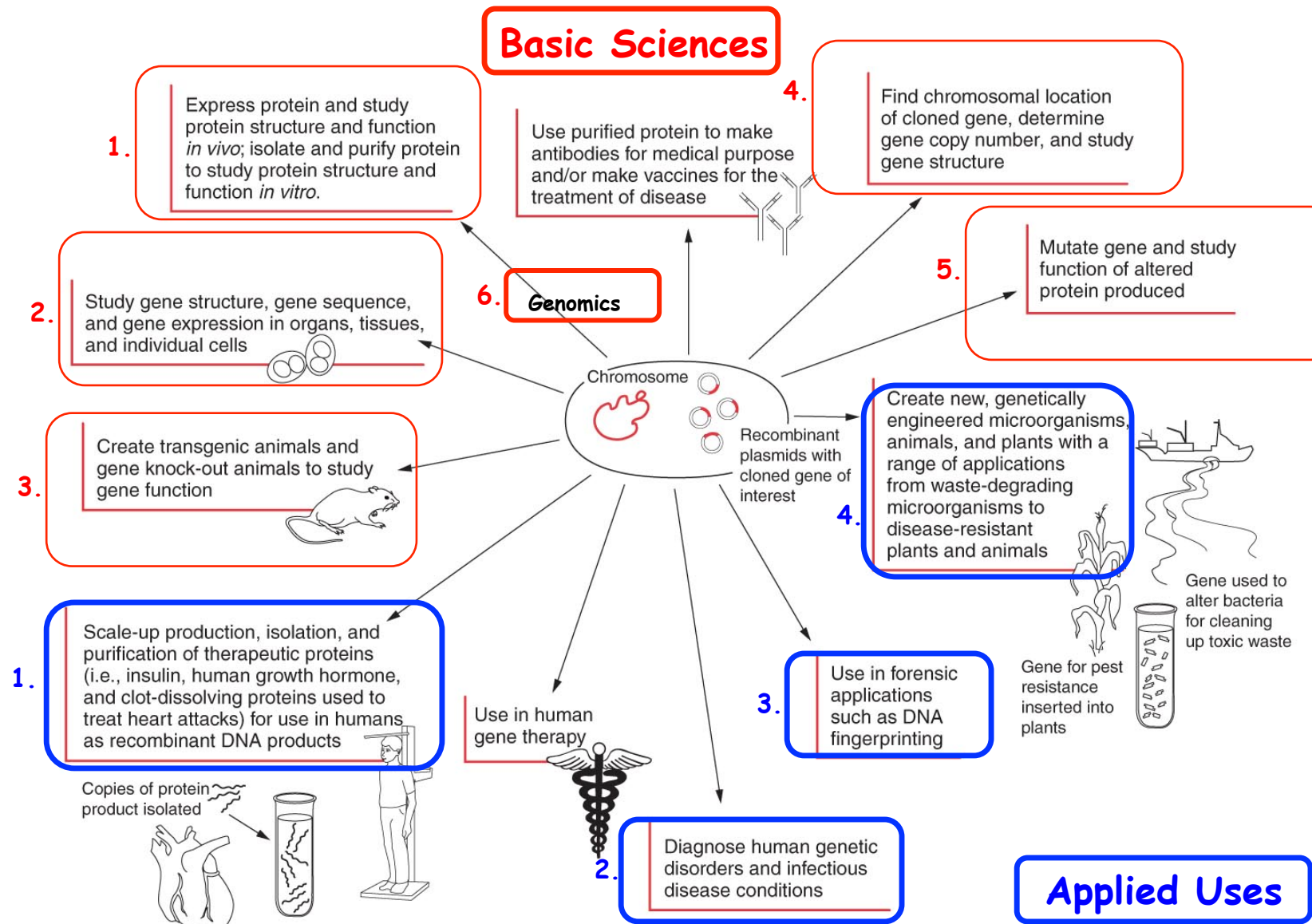
Twenty-First Century Genetic Engineering  
Applications

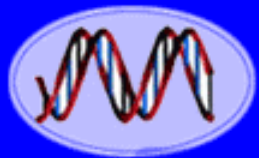
**UCLA**



**UCDAVIS**  
UNIVERSITY OF CALIFORNIA

# There Are Numerous Applications of Genetic Engineering - Many Have Been Discussed in Class To Date

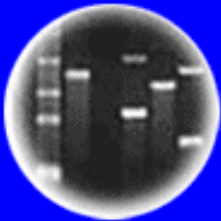




**DNA**  
Genetic Code of Life



Entire Genetic Code  
of a Bacteria



DNA Fingerprinting



Cloning: Ethical Issues  
and Future Consequences



Plants of Tomorrow



# Drugs & Vaccines



# Recombinant Drugs Made In Bacteria And Mammalian Cells To Treat Human & Animal Diseases

**Table 10.1** Some recombinant proteins that have been approved for human use in either the United States or the European Union

Compound	Company	Disorder
Factor VIII	Baxter Healthcare, Genetics Institute, Centeon, Bayer	Hemophilia A
Factor VIIa	Novo Nordisk	Some forms of hemophilia
Factor IX	Genetics Institute	Hemophilia B
Hirudin	Ciba Novartis, Europharm, Hoechst Marion Roussel	Venous thrombosis, heparin-associated thrombocytopenia
Tissue plasminogen activator	Genentech	Acute myocardial infarction
Truncated tissue plasminogen activator	Galenus Mannheim, Boehringer Mannheim/Centocor	Acute myocardial infarction
Insulin	Eli Lilly, Novo Nordisk, Hoechst AG	Diabetes mellitus
Insulin analogues	Eli Lilly, Novo Nordisk, Aventis	Diabetes mellitus
Human growth hormone	Eli Lilly, Genentech, Biotechnology General, Pharmacia, Upjohn, Novo Nordisk, Serono Laboratories	Growth hormone deficiency in children
Human growth hormone analogue	Genentech	Growth hormone deficiency in children
Human growth hormone	Serono Laboratories	AIDS-associated catabolism and wasting
Glucagon	Novo Nordisk	Hypoglycemia
Thyrotrophin- $\alpha$	Genzyme	Thyroid cancer
Follicle-stimulating hormone	Ares-Serono, Organon	Anovulation and superovulation
Erythropoietin	Amgen, Ortho Biotech, Boehringer-Mannheim	Anemia
Platelet-derived growth factor	Ortho-McNeil Pharmaceuticals, Janssen-Cilag	Lower-extremity diabetic neuropathic ulcers
DNase I	Genentech	Cystic fibrosis
$\beta$ -Glucocerebrosidase analogue	Genzyme	Gaucher disease
IFN- $\alpha_{2a}$	Hoffmann-La Roche, Schering-Plough	Hairy cell leukemia, hepatitis B and C
Synthetic type 1 IFN- $\alpha$	Amgen, Yamanouchi Europe	Chronic hepatitis C
IFN- $\alpha_{2b}$	Schering-Plough	Hairy cell leukemia, genital warts, hepatitis B and C
IFN- $\beta_{1b}$ analogues	Schering AG, Berlex Laboratories, Chiron	Multiple sclerosis
IFN- $\beta_{1a}$	Biogen, Ares-Serono	Relapsing multiple sclerosis
IFN- $\gamma_{1b}$	Genentech	Chronic granulomatous disease
IL-2 analogue	Chiron	Renal cell carcinoma
IL-11 analogue	Genetics Institute	Prevention of chemotherapy-induced thrombocytopenia

*Mammalian Cells*  
*Why?*

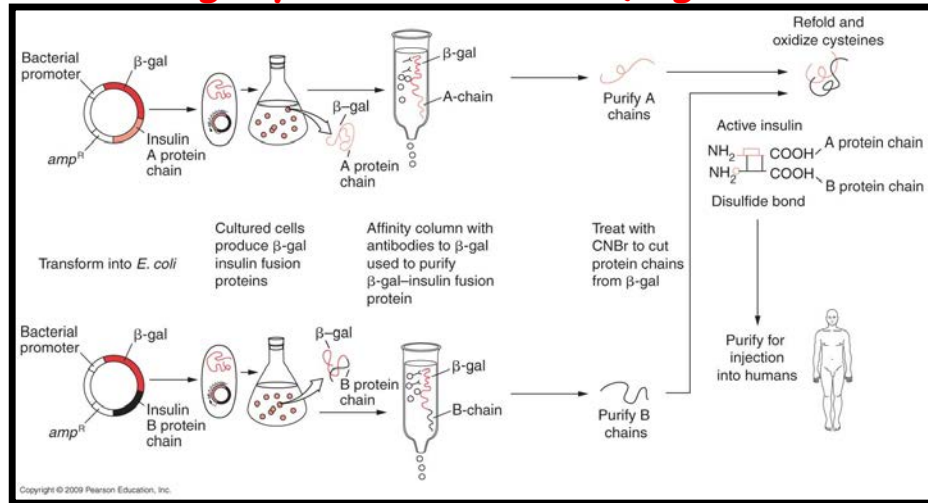
*E. coli*

Abbreviations: IFN, interferon; IL, interleukin.

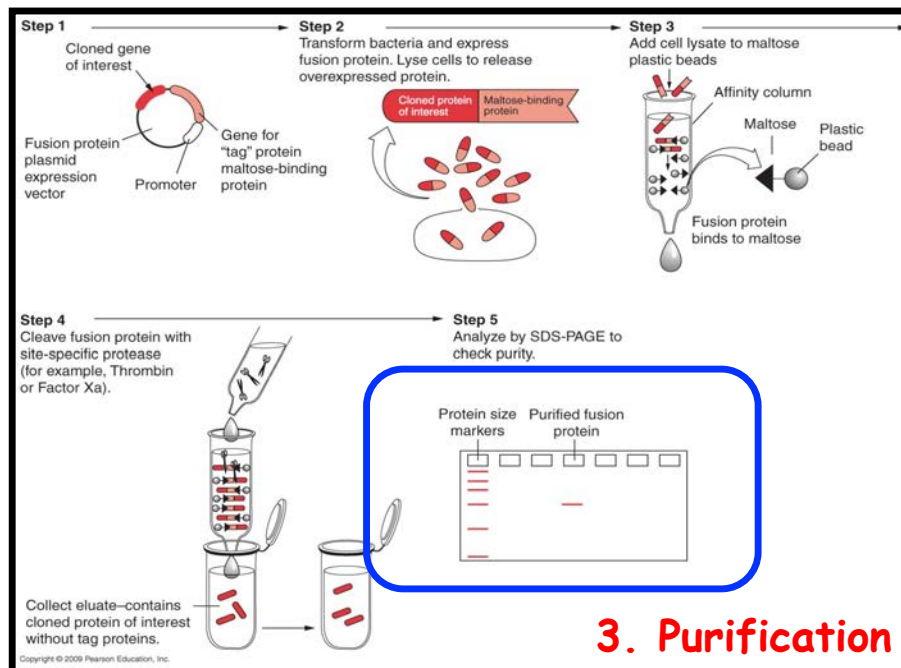
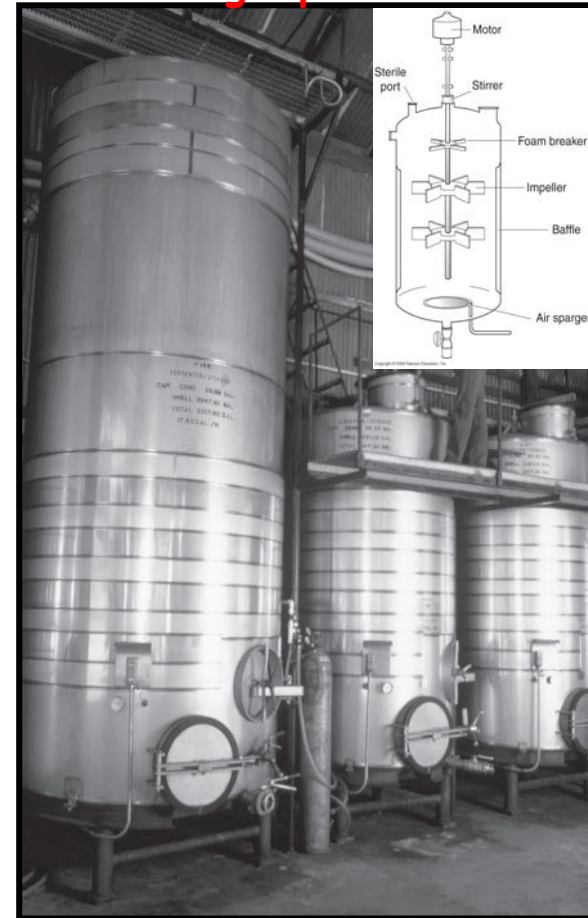


# Manufacturing Recombinant Drugs Requires Industrial-Scale Facilities For Growing Genetically Engineered Cells

## 1. Cloning Synthetic cDNAs (e.g., Insulin cDNA)



## 2. Scaling Up Production



## 3. Purification

## 4. FDA Approval & Sale



# Must Have FDA Approval of Recombinant Drug Production Process, & Drug Safety, and Use

**Table 12.1 PRIMARY FEDERAL REGULATORY AGENCIES IN THE UNITED STATES**

## Regulatory Oversight of Biotechnology Products Agency

U.S. Department of Agriculture

Environmental Protection Agency

Food and Drug Administration

## Product Regulated

Plants, plant pests (including microorganisms), animal vaccines

Microbial/plant pesticides, other toxic substances, microorganisms, animals producing toxic substances

Food, animal feeds, food additives, human and animal drugs, human vaccines, medical devices, transgenic animals, cosmetics

## Major Laws that Empower Federal Agencies to Regulate Biotechnology Law

The Plant Protection Act

The Meat Inspection Act

The Poultry Products Inspection Act

The Eggs Products Inspection Act

The Virus Serum Toxin Act

The Federal Insecticide, Fungicide, and Rodenticide Act

The Toxic Substances Control Act

The Food, Drug, and Cosmetics Act

The Public Health Service Act

The Dietary Supplement Health and Education Act

The National Environmental Protection Act

USDA

USDA

USDA

USDA

USDA

EPA

EPA

FDA, EPA

FDA

FDA

USDA, EPA, FDA



Source: [www.fda.gov](http://www.fda.gov).

Copyright © 2009 Pearson Education, Inc.

# Federal Food, Drug, and Cosmetic Act



Draft Year: 1938 Amendment Years: 1954, 1958

*FDR signed the Food, Drug, and Cosmetic Act on 25 June 1938.*

➤ The FFDCA is a national act

The act requires...

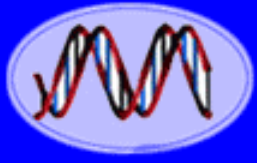
- ✓ Pre-market safety approval of all new drugs
- ✓ Prohibited false therapeutic claims for drugs
- ✓ Drugs labeled with adequate directions for safe use
- ✓ Cosmetics and medical devices under FDA's jurisdiction
- ✓ Drugs be labeled with adequate directions for safe use.

➤ The FDA is responsible for regulations and enforcement



(Left)

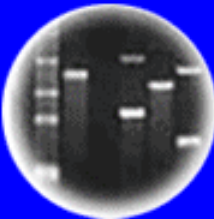
FDA  
employee  
inspecting  
fish for  
pollutants  
or toxins



DNA  
Genetic Code of Life



Entire Genetic Code  
of a Bacteria



DNA Fingerprinting



Cloning: Ethical Issues  
and Future Consequences



Plants of Tomorrow

# How Can the FDA Regulate Drug Approval?

“We the People of the United States, in order to form a more perfect Union, establish justice, insure domestic tranquility, provide for the common defense, promote the General Welfare.....”



# Article I - Section 8.1 of the US Constitution

## The Congress shall have the Power:

[1] “To lay and collect Taxes, Duties, Imposts, and Excises, to pay the Debts and provide for the common Defense and general Welfare of the United States; but all Duties, Imposts, and Excises shall be uniform throughout the United States”

Key Concept: Provide For the General Welfare-Which Can Apply to Almost Everything Dealing With Science, Health, Medicine, Agriculture, and Safety!

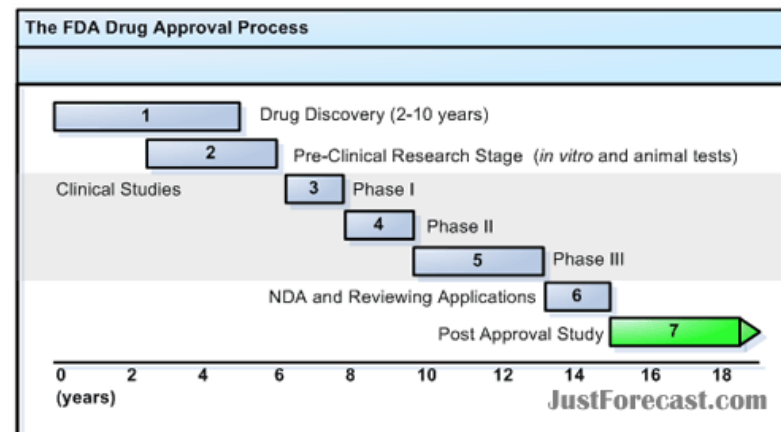
# Drug Testing For FDA Approval is a LONG and COSTLY Process

**TABLE 7.1 FOOD AND DRUG ADMINISTRATION REQUIRED TESTING PHASES FOR DRUG APPROVAL**

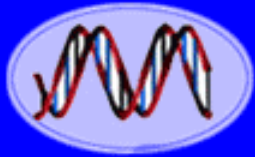
FDA Phase testing involves the use of animals for pre-clinical testing before allowed in humans. If the new drug candidate has proven to be non-toxic and has benefit, then it can be awarded Investigational New Drug (IND) status. If it is successful in the three phases of human testing it can receive a New Drug Application (NDA) and likely approval for marketing. The FDA continues evaluating the NDA for another 2.5 years, resulting a total of about 12 years for a successful drug approval.

	Preclinical Testing		Phase I	Phase II	Phase III		FDA		Phase IV
<b>Years</b>	3.5		1	2	3		2.5	<b>12 total</b>	
<b>Tested on</b>	<b>Animals in the lab</b>		20–80 healthy volunteers	100–300 patient volunteers	1,000–3,000 patient volunteers				
<b>Purpose</b>	Assess safety and biological activity	File IND at FDA	Determine safety and dosage	Evaluate effectiveness and look for side effects	Verify effectiveness, monitor adverse reactions from long-term use	File NDA at FDA	Review process/ approval		Additional testing after approval required by FDA
<b>Success rate</b>	5,000 compounds evaluated			5 enter trials			1 approved		

**\$1 Billion!!!**



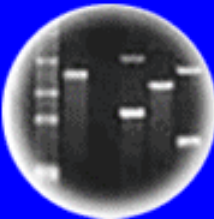
***What If There Was No Patent Protection For Drugs?***



DNA  
Genetic Code of Life



Entire Genetic Code  
of a Bacteria



DNA Fingerprinting



Cloning: Ethical Issues  
and Future Consequences



Plants of Tomorrow

## 1. Article I - Section 8.8

**The Congress shall have the Power:**

[8] “To Promote the Progress of Science and the useful Arts, by securing for limited Times to Authors and Inventors the exclusive Right to their Writings and Discoveries”

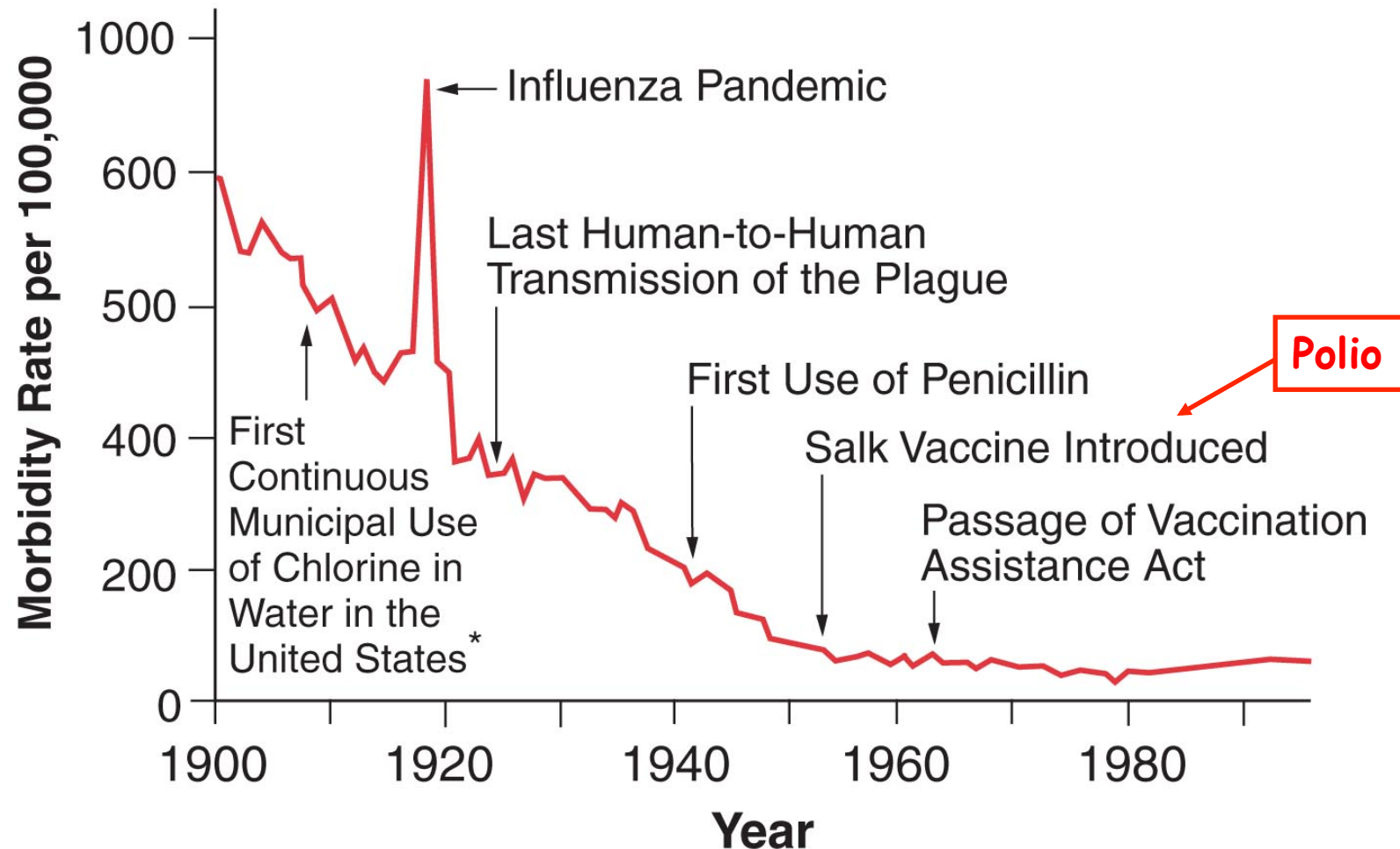
**Keyword:** Inventors not Science.

Wanted to Promote Economic Development &  
Promote a National Economics Policy Grounded  
in Property Rights.  
That is, Entrepreneurship!

**PATENTS!!**

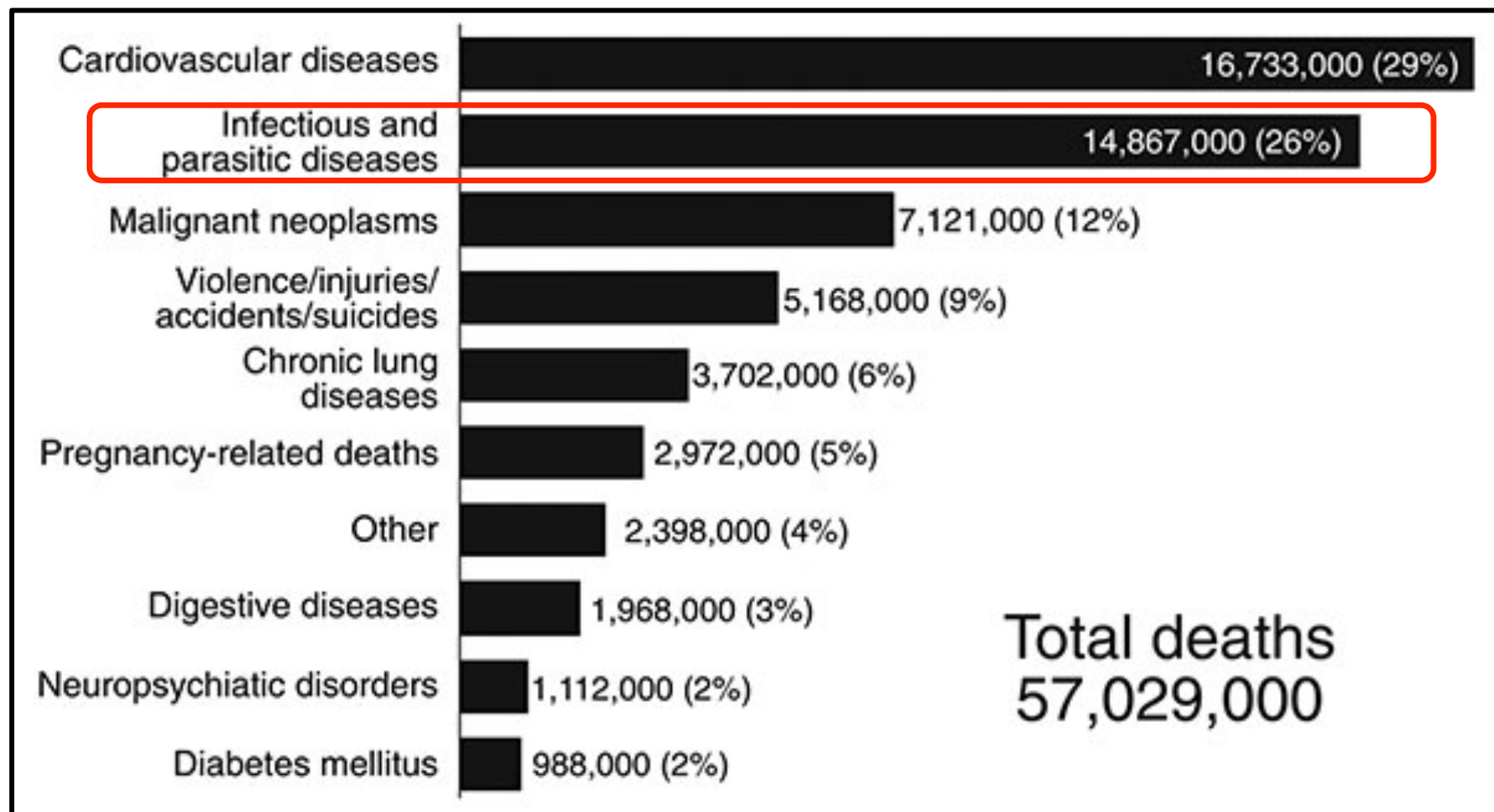
# Over the Past 50 Years **Vaccines and Antibiotics** Have Been Essential in Combating Infectious Diseases

**Crude Death Rate for Infectious Diseases**





## One Fourth of the Annual Deaths Word-Wide Are Caused By Infectious Diseases















# Bacterial Diseases That Are Vaccine Targets

TABLE 28.1 Important Human Bacterial Diseases			
Disease	Pathogen	Vector/Reservoir	Epidemiology
Anthrax	<i>Bacillus anthracis</i>	Animals, including processed skins	Bacterial infection that can be transmitted through contact or ingestion. Rare except in sporadic outbreaks. May be fatal.
Botulism	<i>Clostridium botulinum</i>	Improperly prepared food	Contracted through ingestion or contact with wound. Produces acute toxic poison; can be fatal.
Chlamydia	<i>Chlamydia trachomatis</i>	Humans, STD	Urogenital infections with possible spread to eyes and respiratory tract. Increasingly common over past 20 years.
Cholera	<i>Vibrio cholerae</i>	Human feces, plankton	Causes severe diarrhea that can lead to death by dehydration; 50% peak mortality if untreated. A major killer in times of crowding and poor sanitation; over 100,000 died in Rwanda in 1994 outbreak.
Dental caries	<i>Streptococcus mutans</i> , <i>Streptococcus sobrinus</i>	Humans	A dense collection of these bacteria on the surface of teeth leads to secretion of acids that destroy minerals in tooth enamel; sugar alone will not cause caries.
Diphtheria	<i>Corynebacterium diphtheriae</i>	Humans	Acute inflammation and lesions of respiratory mucous membranes. Spread through respiratory droplets. Vaccine available.
Gonorrhea	<i>Neisseria gonorrhoeae</i>	Humans only	STD, on the increase worldwide. Usually not fatal.
Hansen disease (leprosy)	<i>Mycobacterium leprae</i>	Humans, feral armadillos	Chronic infection of the skin; worldwide incidence about 10–12 million, especially in southeast Asia. Spread through contact with infected individuals.
Lyme disease	<i>Borrelia burgdorferi</i>	Ticks, deer, small rodents	Spread through bite of infected tick. Lesion followed by malaise, fever, fatigue, pain, stiff neck, and headache.
Peptic ulcers	<i>Helicobacter pylori</i>	Humans	Originally thought to be caused by stress or diet, most peptic ulcers now appear to be caused by this bacterium; good news for ulcer sufferers because it can be treated with antibiotics.
Plague	<i>Yersinia pestis</i>	Fleas of wild rodents: rats and squirrels	Killed one-fourth of the population of Europe in the fourteenth century; endemic in wild rodent populations of the western United States today.
Pneumonia	<i>Streptococcus</i> , <i>Mycoplasma</i> , <i>Chlamydia</i> , <i>Haemophilus</i>	Humans	Acute infection of the lungs; often fatal without treatment. Vaccine for streptococcal pneumonia available.
Tuberculosis	<i>Mycobacterium tuberculosis</i>	Humans	An acute bacterial infection of the lungs, lymph, and meninges. Its incidence is on the rise, complicated by the development of new strains of the bacterium that are resistant to antibiotics.
Typhoid fever	<i>Salmonella typhi</i>	Humans	A systemic bacterial disease of worldwide incidence. Fewer than 500 cases a year are reported in the United States. Spread through contaminated water or foods (such as improperly washed fruits and vegetables). Vaccines are available for travelers.
Typhus	<i>Rickettsia typhi</i>	Lice, rat fleas, humans	Historically a major killer in times of crowding and poor sanitation; transmitted from human to human through the bite of infected lice and fleas. Peak untreated mortality rate of 70%.

**DTaP Vaccine – Diphtheria, Tetanus, Pertussis**  
***Corynebacterium diphtheriae*, *Clostridium tetani*, & *Bordetella pertussis***

**All students entering, advancing or transferring into 7th grade need proof of an adolescent whooping cough booster immunization (called "DTap")**

# Viral Diseases That Are Vaccine Targets

TABLE 27.1 Important Human Viral Diseases			
Disease	Pathogen	Genome	Vector/Epidemiology
Chicken pox	Varicella zoster	 Double-stranded DNA	Spread through contact with infected individuals. No cure. Rarely fatal. Vaccine approved in U.S. in early 1995.
Hepatitis B (viral)	Hepadnavirus	 Double-stranded DNA	Highly infectious through contact with infected body fluids. Approximately 1% of U.S. population infected. Vaccine available. No cure. Can be fatal.
Herpes	Herpes simplex virus	 Double-stranded DNA	Blisters; spread primarily through skin-to-skin contact with cold sores/blisters. Very prevalent worldwide. No cure. Exhibits latency—the disease can be dormant for several years.
Mononucleosis	Epstein-Barr virus	 Double-stranded DNA	Spread through contact with infected saliva. May last several weeks; common in young adults. No cure. Rarely fatal.
Smallpox	Variola virus	 Double-stranded DNA	Historically a major killer; the last recorded case of smallpox was in 1977. A worldwide vaccination campaign wiped out the disease completely.
AIDS	HIV	 (+ ) Single-stranded RNA	Acute viral infection of the CNS that can lead to paralysis and is often fatal. Prior to the development of Salk's vaccine in 1954, 60,000 people a year contracted the disease in the U.S. alone.
Yellow fever	Flavivirus	 (+ ) Single-stranded RNA	Spread from individual to individual by mosquito bites; a notable cause of death during the construction of the Panama Canal. If untreated, this disease has a peak mortality rate of 60%.
Ebola	Filoviruses	 (-) Single-stranded RNA	Acute hemorrhagic fever; virus attacks connective tissue, leading to massive hemorrhaging and death. Peak mortality is 50–90% if untreated. Outbreaks confined to local regions of central Africa.
Influenza	Influenza viruses	 (-) Single-stranded RNA (eight segments)	Historically a major killer (20–50 million died during 18 months in 1918–1919); wild Asian ducks, chickens, and pigs are major reservoirs. The ducks are not affected by the flu virus, which shuffles its antigen genes while multiplying within them, leading to new flu strains.
Measles	Paramyxoviruses	 (-) Single-stranded RNA	Extremely contagious through contact with infected individuals. Vaccine available. Usually contracted in childhood, when it is not serious; more dangerous to adults.
SARS	Coronavirus	 (-) Single-stranded RNA	Acute respiratory infection; an emerging disease, can be fatal, especially in the elderly. Commonly infected animals include bats, foxes, skunks, and raccoons. Domestic animals can be infected.
Rabies	Rhabdovirus	 (-) Single-stranded RNA	An acute viral encephalomyelitis transmitted by the bite of an infected animal. Fatal if untreated. Commonly infected animals include bats, foxes, skunks, and raccoons. Domestic animals can be infected.

# Bacterial & Viral Diseases Can Be Prevented Using Vaccines!!!!

## Kids who didn't get whooping cough vaccine a cause of outbreak, scientists say

By Mary MacVean

2:18 PM PDT, September 30, 2013

Children who did not get vaccinated against whooping cough are one of the causes of the 2010 outbreak of the illness, when more cases were reported than in any year since 1947, researchers say. advertisement

Researchers who looked at the geography of the cases suggest that clusters of “nonmedical exemptions” to immunizations were one of several factors in the California outbreak. They reported their findings Monday in the journal Pediatrics.

In California in 2010, there were 9,120 cases of the illness that’s also called [pertussis](#) – one-third of all the U.S. cases. Los Angeles had 1,000 of those cases. Whooping cough is a respiratory ailment marked by bouts of coughing that are accompanied by a noise that can frighten parents – hence the name.

**DTaP Vaccine – Diphtheria, Tetanus, Pertussis**  
*Corynebacterium diphtheriae, Clostridium tetani, & Bordetella pertussis*

*All students entering, advancing or transferring into 7th grade need proof of an adolescent whooping cough booster immunization (called “Tdap”)*

*How Can There Be a Law REQUIRING Vaccinations?*



## Edward Jenner Using Cowpox to Vaccinate a Child Against Smallpox



Vaccine From Vacca or Cow

~1797



**In 1776 George Washington Lost 1,000 Men to Battle  
And 100,000 Men to Smallpox!**

**Washington Had His Army Innoculated With a Small Amount of Fluid From a  
Smallpox Victim and the Smallpox Rate Went Down**

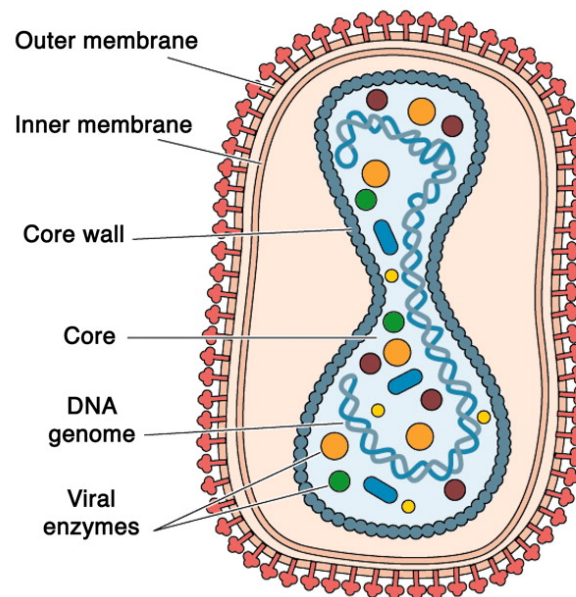
**Smallpox Was One of the First Biological Warfare Agents-Having Been  
Used For Centuries**

**Responsible For 300-500 MILLION Deaths in the 20th Century**



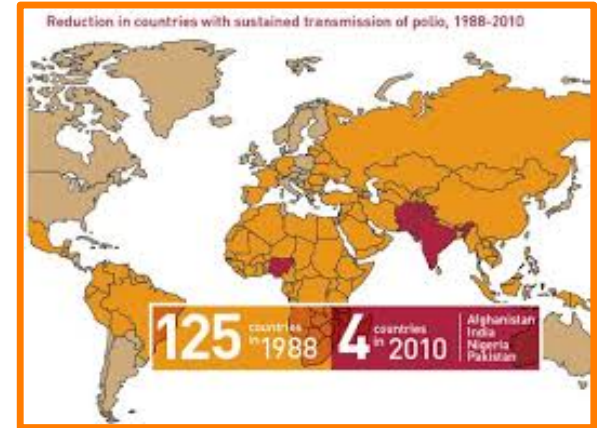
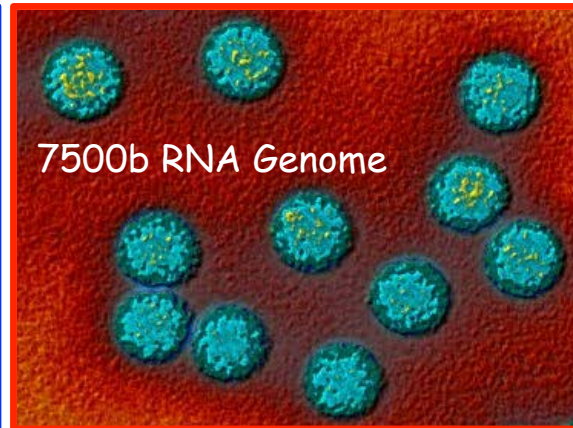
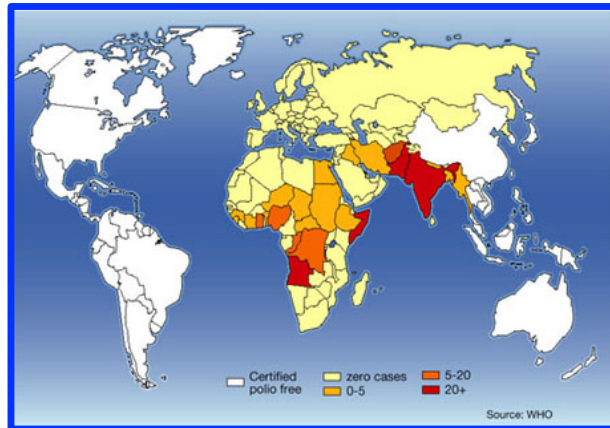
# Smallpox is the Only Human Infectious Disease That Has Been Eradicated Globally

186,000 bp dsDNA Genome

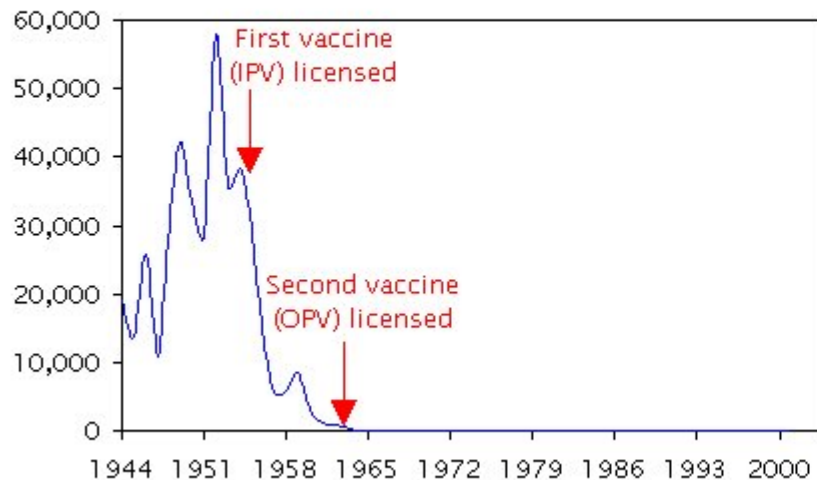


*The Last Reported Case of Smallpox in US was in Was Reported in 1949 & in World in 1977 - Smallpox Vaccinations Are No Longer Given. Smallpox Virus Destroyed in 1980! What About Monkeypox?*

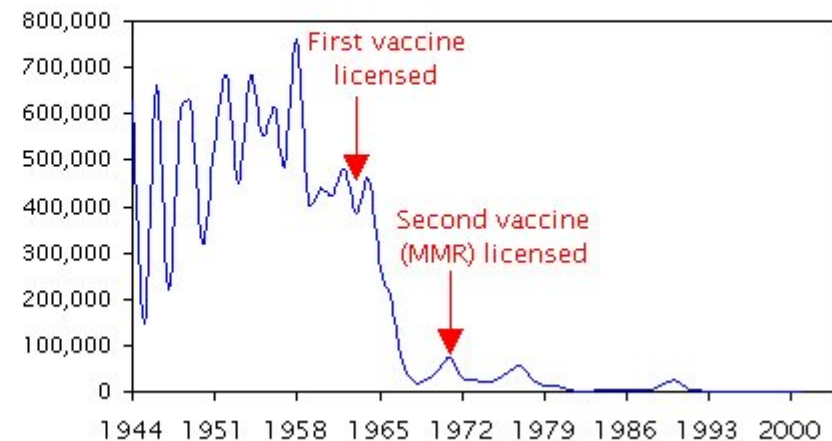
# What About Polio?



## U.S. Polio Cases 1944-2001



## U.S. Measles Cases 1944-2001





# California Vaccination Requirements

## GUIDE TO IMMUNIZATIONS REQUIRED FOR SCHOOL ENTRY

### Grades K-12



**INSTRUCTIONS** Use this guide as a quick reference to help you determine whether children seeking admission to your school meet California's school immunization requirements. For the actual laws, see Health and Safety Code, Division 105, Part 2, Chapter 1, Sections 120325-120380; California Code of Regulations, Title 17, Division 1, Chapter 4, Subchapter 8, Sections 6000-6075. If you have any questions, call the Immunization Coordinator at your local health department.

**IMMUNIZATION REQUIREMENTS** To enter into public and private elementary and secondary schools (grades kindergarten through 12, including transitional kindergarten), children under age 18 years must have immunizations.

VACCINE	REQUIRED DOSES
Polio	<b>4 doses at any age, but...</b> 3 doses meet requirement for ages 4–6 years if at least one was given on or after the 4 <sup>th</sup> birthday <sup>1</sup> ; 3 doses meet requirement for ages 7–17 years if at least one was given on or after the 2 <sup>nd</sup> birthday. <sup>1</sup>
Diphtheria, Tetanus, and Pertussis	<p><b>Age 6 years and under:</b> DTP, DTaP or any combination of DTP or DTaP with DT (diphtheria and tetanus) <b>5 doses at any age, but...</b> 4 doses meet requirements for ages 4–6 years if at least one was on or after the 4<sup>th</sup> birthday.<sup>1</sup></p> <p><b>Age 7 years and older:</b> Tdap, Td, or DTP, DTaP or any combination of these <b>4 doses at any age, but...</b> 3 doses meet requirement for ages 7–17 years if at least one was on or after the 2<sup>nd</sup> birthday.<sup>1</sup> If last dose was given before the 2<sup>nd</sup> birthday, one more (Tdap) dose is required.</p>
Measles, Mumps, Rubella (MMR)	<p><b>Age 4-6 years (kindergarten and above): 2 doses<sup>2</sup></b> both on or after 1<sup>st</sup> birthday.<sup>1</sup></p> <p><b>7<sup>th</sup> grade: 2 doses<sup>2</sup></b> both on or after 1<sup>st</sup> birthday.<sup>1</sup></p> <p><b>Age 7-17 years and not entering or advancing into 7<sup>th</sup> grade: 1 dose</b> on or after 1<sup>st</sup> birthday.<sup>1</sup></p>
Hepatitis B <sup>3</sup>	<b>Age 4-6 years (kindergarten and above): 3 doses.</b>
Varicella	<b>1 dose<sup>4, 6</sup></b>
Tdap Booster (Tetanus, reduced diphtheria, and pertussis)	<b>7<sup>th</sup> grade: 1 dose</b> on or after 7 <sup>th</sup> birthday. <sup>5, 7</sup>

# California Vaccination Requirements

## Quick Facts and Resources for California Residents

 Medical

 Religious

 Philosophical

**Quick Fact:** In 2012 the personal belief exemption became more restrictive in that there is an additional requirement of a signature from a health care practitioner to obtain the exemption. Effective Jan. 1, 2014 parents, guardians and emancipated minors must now obtain this additional signature when filing with the governing authority the necessary documents that state which vaccinations have not been given on the basis that they are contrary to his or her beliefs. When the law was amended for this change, Governor Brown also issued an [executive order](#) directing the health department to include a separate religious exemption on the new exemption form.

In 2011 a new law was passed which allows minors 12 years old and older to consent to vaccines for sexually transmitted diseases without the knowledge or consent of their parents. This includes HPV vaccine.

# California's measles outbreak is over, but vaccine fight continues

The state epidemiologist, Dr. Gil Chavez, said immunization rates in some schools are at 50% or lower, creating an ideal environment for the virus to spread quickly. A [study](#) published in JAMA Pediatrics last month calculated that the measles virus that caused the outbreak spread in areas where vaccination rates were likely between 50% and 86%.

The outbreak prompted two state lawmakers, Sens. Pan and Ben Allen (D-Santa Monica) to push for closing a loophole in state law that gives parents the right to refuse state-required vaccinations due to their personal beliefs while still sending their children to public and private schools.

“  
**If we had higher levels of immunity in the community, this outbreak would not have happened.**

”

- Dr. Gil Chavez, state epidemiologist

 [SHARE THIS QUOTE](#)

Early on, the bill appeared to have momentum, winning approval of the Senate Health Committee after Gov. Jerry Brown signaled he was open to considering an elimination of all but medical waivers to vaccines.

But SB 277 [stalled](#) this week in the Senate Education Committee, where members demanded changes after hundreds of parents lined up to say they would pull their kids out of school if the bill passed. A vote is



# Senate Bill 277 Introduced to End California's Vaccine Exemption Loophole

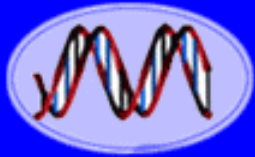
February 19, 2015

**SACRAMENTO** –Dr. Richard Pan, a pediatrician and Senator representing Sacramento, Senator Ben Allen, the former Board President of the Santa Monica-Malibu Unified School District and Assemblymember Lorena Gonzalaz who represents San Diego have introduced Senate Bill 277 to repeal the personal belief exemption that currently allows parents to opt their child out of vaccines in our schools.

If this legislation is passed, California will join thirty-two other states that don't allow parents to opt out of vaccination requirements using a personal belief exemption.



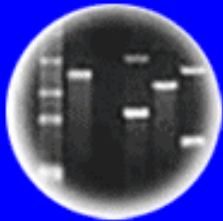




DNA  
Genetic Code of Life



Entire Genetic Code  
of a Bacteria



DNA Fingerprinting



Cloning: Ethical Issues  
and Future Consequences



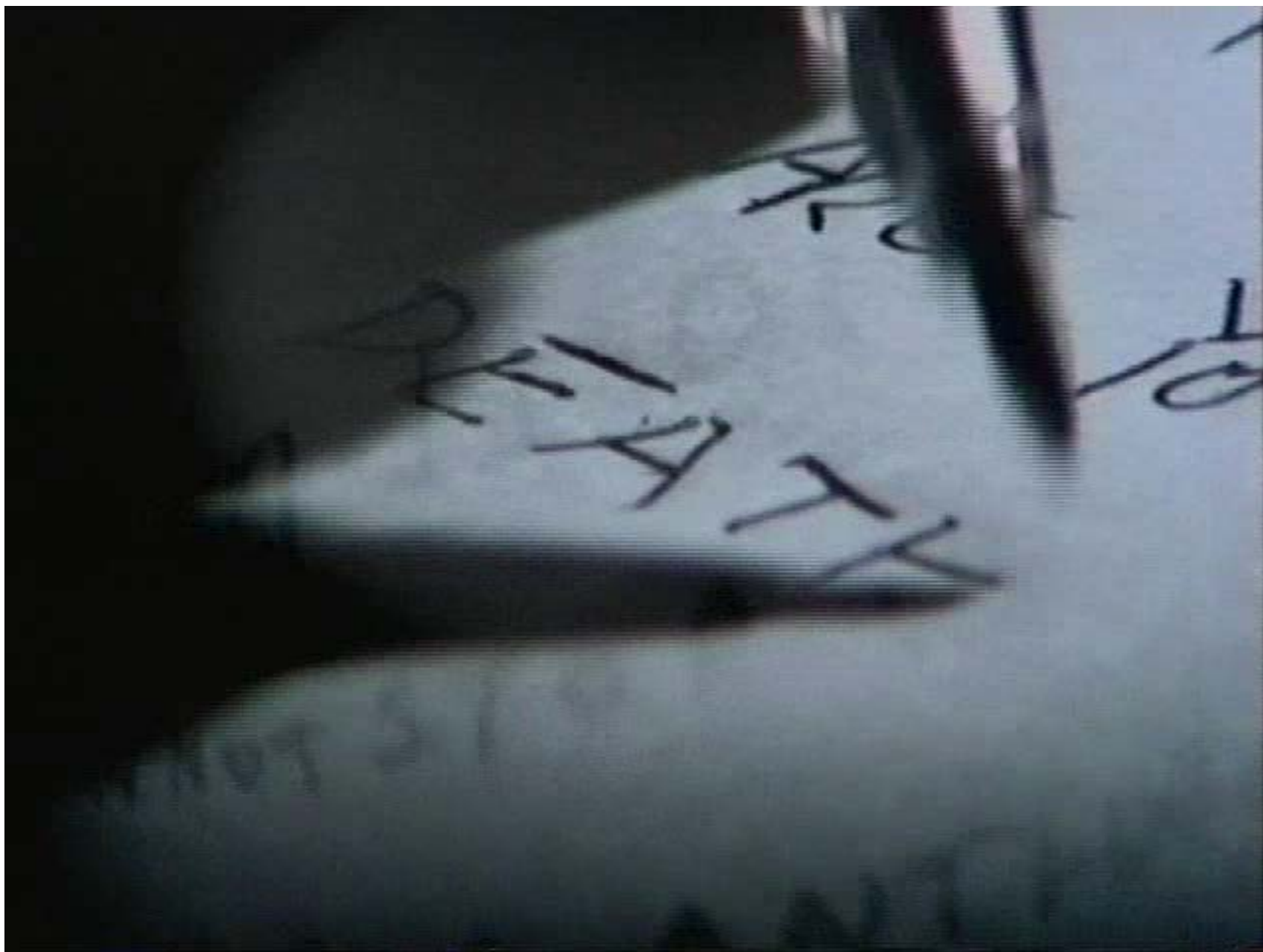
Plants of Tomorrow

Should Parents Be Allowed To  
Exempt Their Children From  
Required Vaccinations?

a. yes

b. no

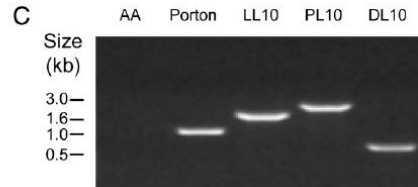
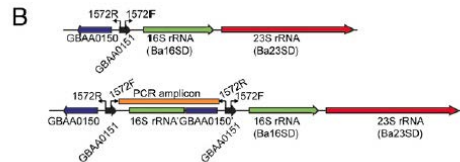
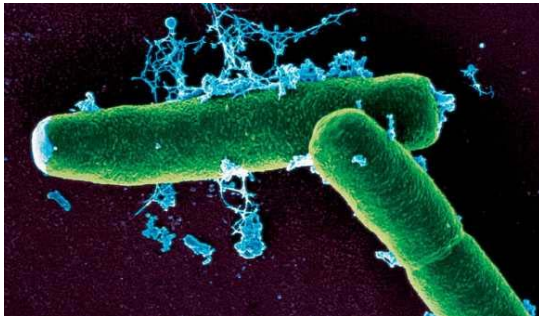




October 18, 2001

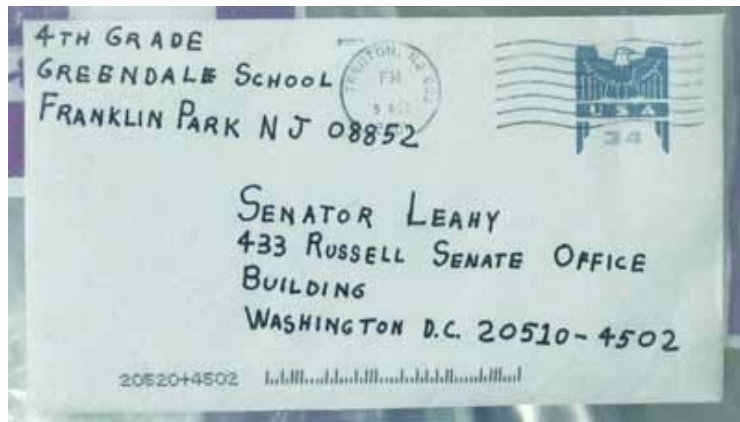
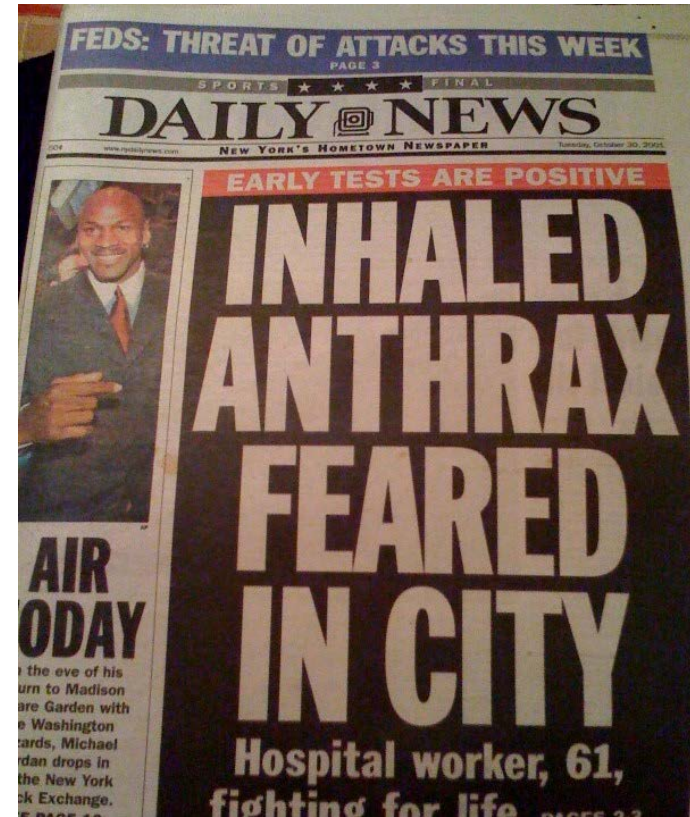
A NATION CHALLENGED: THE ANTHRAX THREAT

## A NATION CHALLENGED: THE ANTHRAX THREAT; TESTS SHOW ANTHRAX EXPOSURE IN AT LEAST 30 CAPITAL WORKERS



DNA  
Fingerprints  
Were Used  
To Find the  
Source of the  
Strain - *Rare  
Ames Strain*

Where Did  
the Anthrax  
Strain Come  
From?





HART  
SENATE OFFICE BUILDING

MEMBERS  
AND STAFF  
AT ALL TIMES  
VISITORS  
FROM  
9:00A.M.











# Using Genetic Engineering To Produce Vaccines Can Play a Big Role in Combating Bioweapons

**TABLE 5.5 POTENTIAL BIOLOGICAL WEAPONS**

Agent	Disease Threat and Common Symptoms
<i>Brucella</i> (bacteria)	Different strains of <i>Brucella</i> infect livestock such as cattle and goats. They can cause brucellosis in animals and humans. Prolonged fever and lethargy are common symptoms. The disease can be mild or life-threatening.
<i>Bacillus anthracis</i> (bacterium)	Anthrax. Skin form (cutaneous) produces skin-surface lesions that are generally treatable. Inhalation anthrax initially produces flu-like symptoms leading to pulmonary pneumonia, which is usually fatal.
<i>Clostridium botulinum</i> (bacterium)	Botulism. Caused by ingestion of food contaminated with <i>C. botulinum</i> or its toxins. Varying degrees of paralysis of the muscular system created by botulinum toxins are typical. Respiratory paralysis and cardiac arrest often cause death.
Ebola virus or Marburg virus	Both are highly virulent viruses that cause hemorrhagic fever. Symptoms include severe fever, muscle/joint pain, and bleeding disorders.
<i>Francisella tularensis</i> (bacterium)	Tularemia. Lung inflammation can cause respiratory failure, shock, and death.
Influenza viruses (a large, highly contagious group)	Influenza (flu). Severity and outcome depend largely on the strain of the virus.
<i>Rickettsia</i> (several bacteria strains)	Different strains cause diseases such as Rocky Mountain spotted fever and typhus.
Variola virus	Smallpox. Chills, high fever, backache, headache, and skin lesions.
<i>Yersinia pestis</i> (bacterium)	Bubonic plague. High fever, headache, painful swelling of lymph nodes, shock, circulatory collapse, organ failure, and death within days after infection in a majority of cases.



# Omics Find Chinks in Ebola Armor for Vaccine and Drug Development

The guiding principle is global information accessibility.

## Pint-sized DNA sequencer impresses first users

Portable device offers on-the-spot data to fight disease, catalogue species and more.

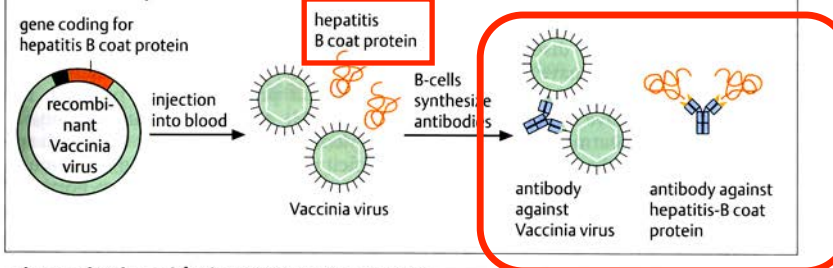


# Using Genetic Engineering To Make Vaccines

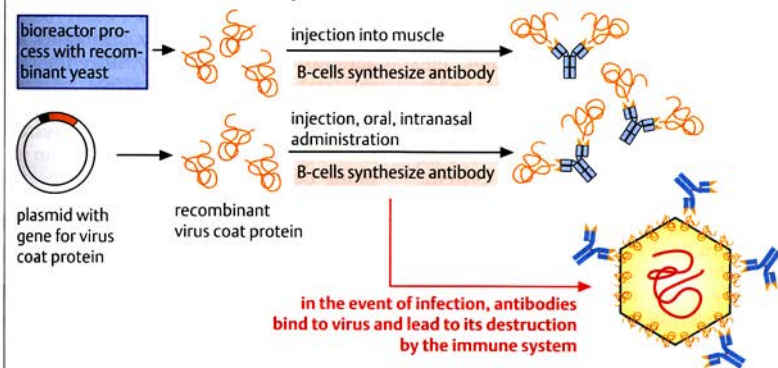
## Recombinant vaccines (selection)

		antigen	status
viruses	hepatitis B	surface antigens	registered
	<i>Herpes simplex</i> type 2	surface antigens	clinical studies
	rabies vaccine	surface antigens	not registered
	yellow fever virus	surface antigens	preclinical studies
	AIDS virus	surface antigens	clinical studies
bacteria	<i>Streptococcus pneumoniae</i>	polysaccharide conjugate	registered
	<i>Clostridium tetani</i>	tetanus toxin	not registered
	<i>Mycobacterium tuberculosis</i>	surface antigens	clinical studies
parasites	<i>Plasmodium falciparum</i>	(malaria)	clinical studies
	<i>Trypanosoma</i> sp.	(sleeping sickness)	clinical studies
	<i>Schistosoma mansoni</i>	(bilharziosis)	clinical studies

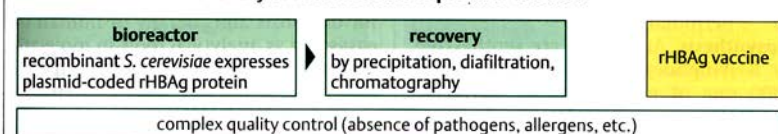
## Vaccination by recombinant Vaccinia virus



## Immunization with virus coat protein or DNA



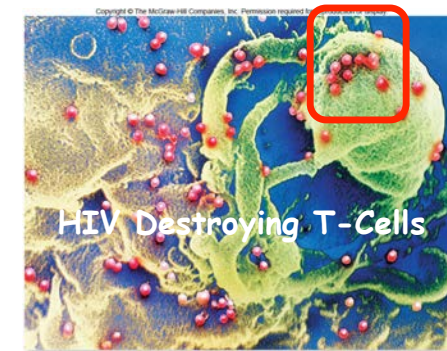
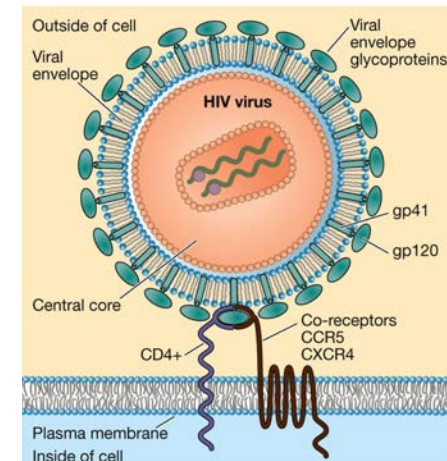
## Fermentation and recovery of recombinant hepatitis B vaccine



Clone Pathogenic Antigen Gene in *E. Coli* or Other Host (e.g., Yeast, Virus) And Synthesize Large Amounts of Antigen

# Recombinant Vaccines Are Being Developed To Combat Many Pathogens

Pathogenic agent	Disease(s)
<b>Viruses</b>	
Varicella-zoster virus	Chicken pox
Cytomegalovirus	Infection in infants and immunocompromised patients
Dengue virus	Hemorrhagic fever
Hepatitis A virus	High fever, liver damage
Hepatitis B virus	Long-term liver damage
Herpes simplex virus type 2	Genital ulcers
Influenza A and B viruses	Acute respiratory disease
Japanese encephalitis virus	Encephalitis
Parainfluenza virus	Inflammation of the upper respiratory tract
Rabies virus	Encephalitis
Respiratory syncytial virus	Upper and lower respiratory tract lesions
Rotavirus	Acute infantile gastroenteritis
Yellow fever virus	Lesions of heart, kidney, and liver
Human immunodeficiency virus	AIDS
<b>Bacteria</b>	
<i>Vibrio cholerae</i>	Cholera
<i>E. coli</i> enterotoxin strains	Diarrheal disease
<i>Neisseria gonorrhoeae</i>	Gonorrhea
<i>Haemophilus influenzae</i>	Meningitis, septicemic conditions
<i>Mycobacterium leprae</i>	Leprosy
<i>Neisseria meningitidis</i>	Meningitis
<i>Bordetella pertussis</i>	Whooping cough
<i>Shigella</i> strains	Dysentery
<i>Streptococcus</i> group A	Scarlet fever, rheumatic fever, throat infection
<i>Streptococcus</i> group B	Sepsis, urogenital tract infection
<i>Streptococcus pneumoniae</i>	Pneumonia, meningitis
<i>Clostridium tetani</i>	Tetanus
<i>Mycobacterium tuberculosis</i>	Tuberculosis
<i>Salmonella typhi</i>	Typhoid fever
<b>Parasites</b>	
<i>Onchocerca volvulus</i>	River blindness
<i>Leishmania</i> spp.	Internal and external lesions
<i>Plasmodium</i> spp.	Malaria
<i>Schistosoma mansoni</i>	Schistosomiasis
<i>Trypanosoma</i> spp.	Sleeping sickness
<i>Wuchereria bancrofti</i>	Filariasis



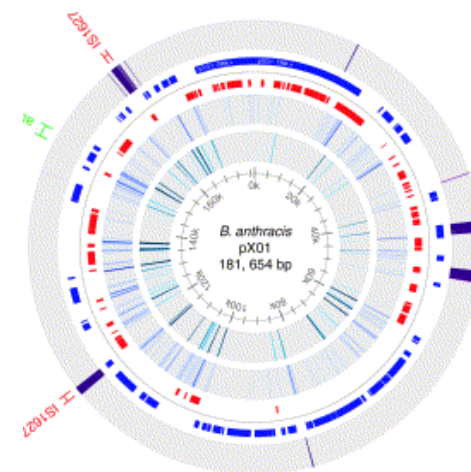
**But a Vaccine To The AIDS Virus Remains Elusive!!**



# Genomics Can Provide Valuable Information About Pathogen Protein Targets For Vaccine Production

**TABLE 5.4** EXAMPLES OF MEDICALLY IMPORTANT VIRAL GENOMES THAT HAVE BEEN SEQUENCED

Virus	Human Disease or Illness	Year Sequenced
Ebola virus	Ebola hemorrhagic fever	1993
Hepatitis A virus	Hepatitis A	1987
Hepatitis B virus	Hepatitis B	1984
Hepatitis C virus	Hepatitis C	1990
Herpes simplex virus, type I	Cold sores	1988
Human immunodeficiency virus (HIV-1)	Acquired immunodeficiency syndrome (AIDS)	1985
Human papillomavirus	Cervical cancer	1985
Human poliovirus	Poliomyelitis	1981
Human rhinovirus	Common cold	1984
Influenza A virus		
• Subtype H5N1 (Avian flu)	Severe flu	2007
• Subtype H5N1 (Swine flu)	Severe flu	2009
Severe acute respiratory coronavirus (SARS-CoV)	Severe acute respiratory syndrome (SARS)	2003
Variola virus	Smallpox	1992



**TABLE 5.3** SELECTED MICROBIAL GENOMES

Bacterium	Human Disease Condition (megabases, mB)	Approximate Genome Size	Approximate Number of Genes
<i>Bacillus anthracis</i>	Anthrax	5.23	5,000
<i>Borrelia burgdorferi</i>	Lyme disease	1.44	853
<i>Chlamydia trachomatis</i>	Eye infections, genitourinary tract infections (e.g., pelvic inflammatory disease)	1.04	896
<i>Escherichia coli</i> O157:H7	Severe food-borne illness (diarrhea)	4.10	5,283
<i>Haemophilus influenzae</i>	Serious infections in children (eye, throat, and ear infections, meningitis)	1.83	1,746
<i>Helicobacter pylori</i>	Stomach (gastric) ulcers	1.66	1,590
<i>Listeria monocytogenes</i>	Listeriosis (serious food-borne illness)	2.94	2,853
<i>Mycobacterium tuberculosis</i>	Tuberculosis	4.41	3,974
<i>Neisseria meningitidis</i> (MC58) infections	Meningitis and blood	2.27	2,158
<i>Pseudomonas aeruginosa</i>	Pneumonia, chronic lung infections	6.30	5,570
<i>Rickettsia prowazekii</i>	Typhus	1.11	834
<i>Rickettsia conorii</i>	Mediterranean spotted fever	1.30	1,374
<i>Streptococcus pneumoniae</i>	Acute (short-term) respiratory infection	2.16	2,236
<i>Yersinia pestis</i>	Plague	4.65	4,012
<i>Vibrio cholerae</i>	Cholera (diarrheal disease)	4.00	3,885



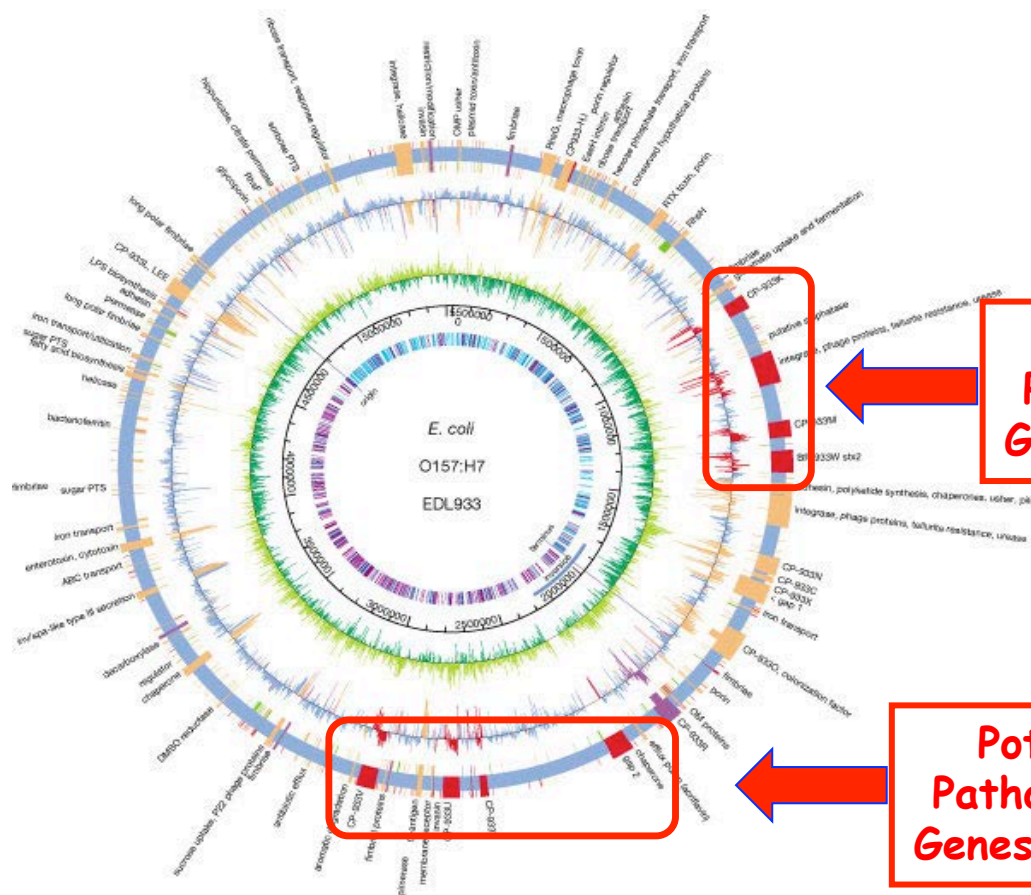
# Comparison of Pathogenic and Non-Pathogenic *E. coli* Genomes



*E. coli* O157:H7 was first recognized as a foodborne pathogen in 1982 during an investigation into an outbreak of hemorrhagic colitis (bloody diarrhea) associated with consumption of contaminated hamburgers (Riley, et al., 1983). The following year, Shiga toxin (Stx), produced by the then little-known *E. coli* O157:H7, was identified as the real culprit.



**75,000 Illnesses  
& 650 Deaths  
Due To *E. coli*  
H0157  
&  
72 Million  
Illnesses &  
5,000 Deaths  
Due to All  
Foodborne  
Diseases  
In US!!!  
A BIG PROBLEM**



**Potential  
Pathogenicity  
Genes/Proteins**

*Nature* **409**, 529-533 (25 January 2001)  
November 2000

# Understanding Pathogen Genes and Infection Process Leads to New Drugs

## For Example - A New Anthrax Drug Just Released

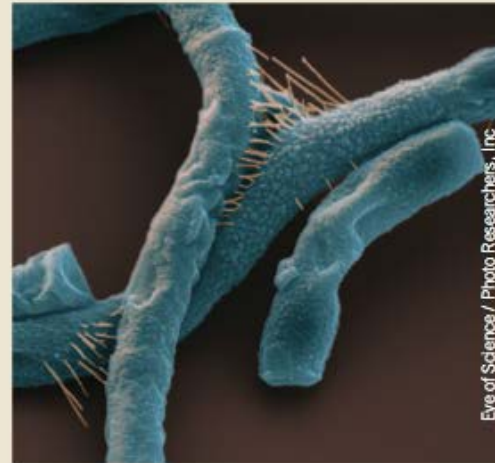
### Anthrax drug first antibacterial mAb to win approval

Officials of the US Food and Drug Administration (FDA) in mid-December approved ABthrax, or raxibacumab, for use in patients with inhalational anthrax. This approval is the first under the agency's 'animal rule', established for evaluating the efficacy of products that would be unethical or impossible to test (except for safety) in humans. ABthrax is a human monoclonal antibody (mAb), licensed for use as an adjunct to conventional antibiotics such as Cipro (ciprofloxacin).

This is also the first time FDA approved a mAb for an antibacterial indication, according to Steven Projan, a senior vice president at MedImmune in Gaithersburg, Maryland. "This should signal a new era in pathogen-specific drugs for the prevention and/or treatment of bacterial infections for bacteria like *Staphylococcus aureus* and *Pseudomonas aeruginosa*, where there are already monoclonal antibodies in clinical trials," he says.

ABthrax interferes with the binding of a key antigen of *Bacillus anthracis*, the bacterial pathogen responsible for anthrax—a potentially deadly infection, particularly when it involves the lungs and becomes systemic. Spores of this pathogen can be used as a bioterror agent—in 2001, spores deliberately distributed through the US Postal Service led to 5 deaths amid 17 cases of anthrax—or in biological warfare. The mAb was developed by Rockville, Maryland-based Human Genome Sciences, a biotech company that GlaxoSmithKline of London acquired last August (*Nat. Biotechnol.* **30**, 815, 2012).

Under a contract from 2005, FDA allowed the US Department of Health and Human Services to purchase and stockpile ABthrax under Project BioShield and within its Biomedical Advanced Research and Development Authority (BARDA). Until full approval came in 2012, however, the mAb was subject to FDA emergency use authority (EUA), according to Amesh Adalja, senior associate at the Center for Biosecurity, a nonprofit organization of the University of Pittsburgh Medical School, in Baltimore. "FDA approval [of ABthrax] makes it easier for physicians to use the product," he says.



*Bacillus anthracis* is the causative agent of anthrax, which affects both humans and animals.

# Synthetic Biology Can Be Used to Rapidly Synthesize Vaccines

## VACCINES

**Synthetic Generation of Influenza Vaccine Viruses for Rapid Response to Pandemics**

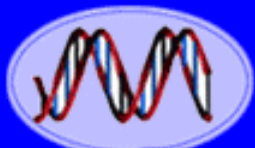
**Synthetic Biologists Engineer A Custom Flu Vaccine In A Week**

A synthetic biology method proves its chops.

**Synthetic Biology Could Speed Flu Vaccine Production**

Advanced genetic engineering is already changing vaccine development and could make inroads into other branches of medicine.

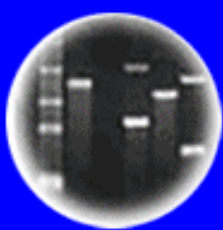




DNA  
Genetic Code of Life



Entire Genetic Code  
of a Bacteria



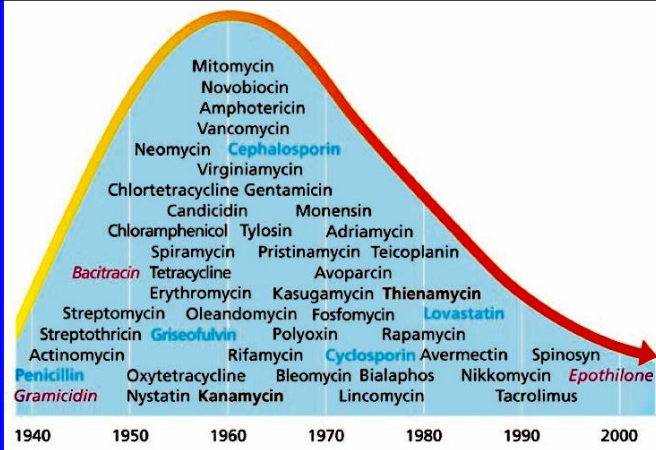
DNA Fingerprinting



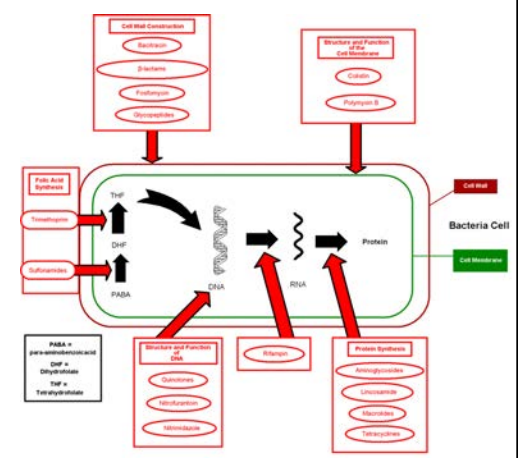
Cloning: Ethical Issues  
and Future Consequences



Plants of Tomorrow



# Antibiotics



# Antibiotic Resistance Is Also A Major Problem in Combating Pathogens

## RISING RESISTANCE

Griffith's & Avery's Bacteria

**MANY ANTIBIOTICS** are no longer effective against certain strains of bacteria, as these examples—collected from different hospitals in the late 1990s—show. One strain of *Staphylococcus aureus* found in Korea, for instance, is 98 percent resistant to penicillin (*top left*); another, found in the U.S., is 32 percent resistant to methicillin (*bottom left*). All these strains are not resistant to vancomycin, for now.

*STAPHYLOCOCCUS AUREUS*  
VS. PENICILLIN



*ENTEROCOCCUS FAECIUM*  
VS. CIPROFLOXACIN (CIPRO)



*STREPTOCOCCUS PNEUMONIAE*  
VS. TETRACYCLINE



*STAPHYLOCOCCUS AUREUS*  
VS. METHICILLIN



*ENTEROCOCCUS FAECIUM*  
VS. AMPICILLIN



*STREPTOCOCCUS PNEUMONIAE*  
VS. PENICILLIN



**Methicillin Resistant  
*Staphylococcus aureus*  
MRSA!!**

IMMUNIZE FOR A HEALTHY FUTURE



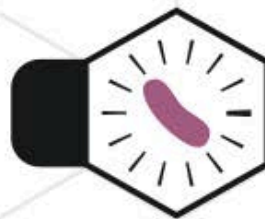
## Media centre

# **WHO's first global report on antibiotic resistance reveals serious, worldwide threat to public health**

New WHO report provides the most comprehensive picture of antibiotic resistance to date, with data from 114 countries



# Antibiotic Resistance Is Also A Major Problem in Combating Pathogens



## How Antibiotic Resistance Happens

**1.**

Lots of germs.  
A few are drug resistant.



**2.**

Antibiotics kill  
bacteria causing the illness,  
as well as good bacteria  
protecting the body from  
infection.



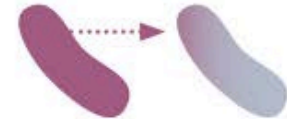
**3.**

The drug-resistant  
bacteria are now allowed to  
grow and take over.

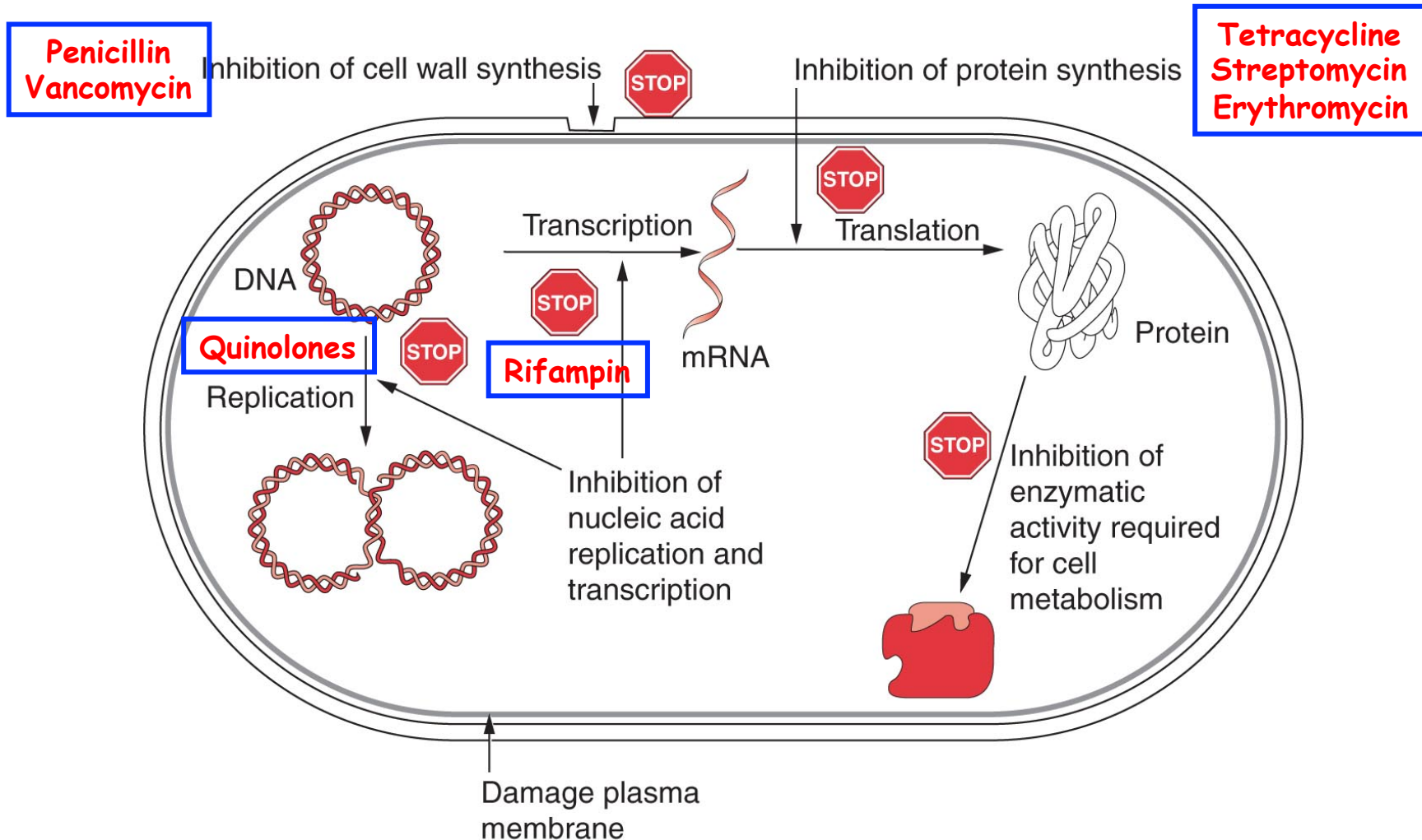


**4.**

Some bacteria give  
their drug-resistance to  
other bacteria, causing  
more problems.

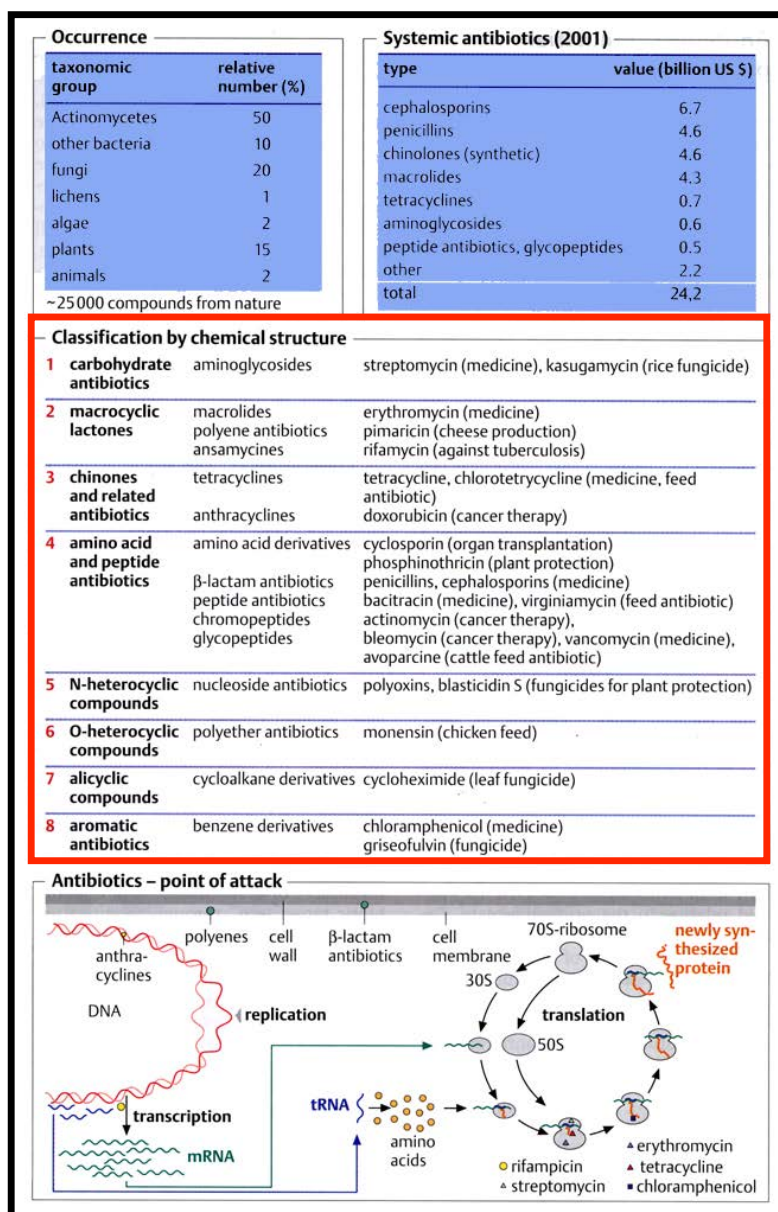


# A Review.....How Do Antibiotics Kill Bacterial Cells?

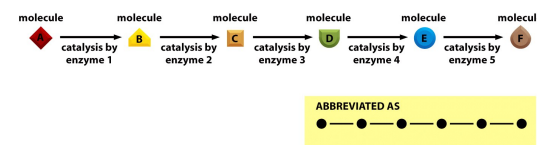


**By Inhibiting Basic Microbial Cell Processes**

# Genetic Engineering Can Be Used To Make Better/More Effective Antibiotics



By Modifying Pathways Leading to Antibiotics In Bacterial Cells. But Need To Know Genes/Proteins in Pathway & By Finding Their Targets In Pathogens As Well







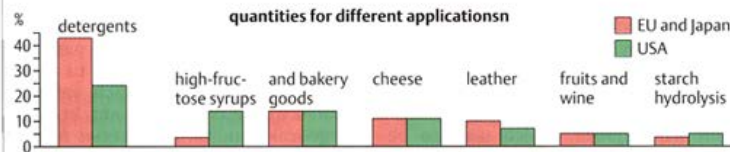
## Industrial & Food Products



# Bacteria & Other Microbes Are the Source Of Many Different Products

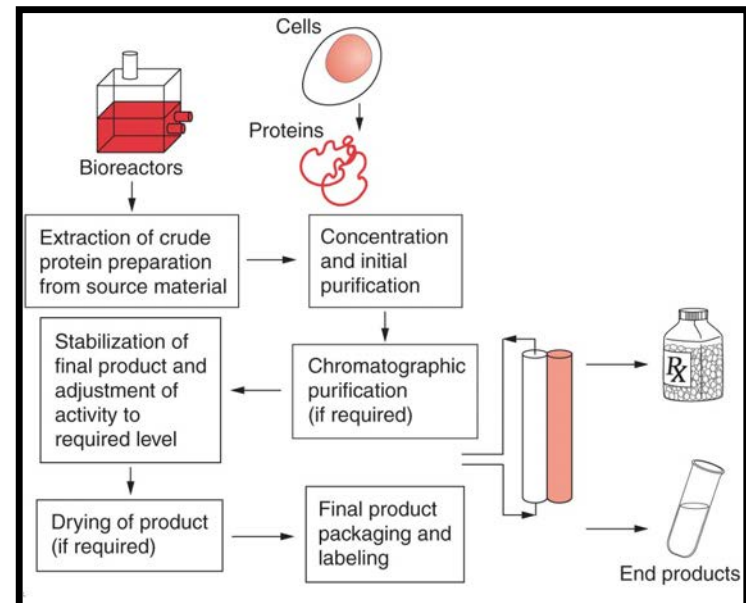
Enzymes as additives in industry

application	enzyme type	organisms (examples)	market size (% of total)	economic advantage
detergents	proteases, cellulases, lipases	<i>Bacillus licheniformis</i> <i>Aspergillus nidulans</i> <i>Trichoderma reesei</i>	40	1
starch hydrolysis	$\alpha$ -amylase	<i>Bacillus amyloliquefaciens</i>	5	3, 4
glucose isomerization	glucose isomerase	<i>Streptomyces venezuelae</i>	7	1, 3
beer brewing	amylase	<i>Bacillus subtilis</i>	3	3, 4
fruit processing, wine	cellulases, hemicellulases, pectinases	<i>Aspergillus niger</i>	5	3, 4, 5, 6
flour, bakery goods	$\alpha$ -amylase, proteases	<i>Aspergillus oryzae</i>	8	1, 3
cheese manufacture, aroma	proteases, chymosin, lipases	animal rennin, <i>Rhizomucor miehei</i> , <i>Saccharomyces cerevisiae</i>	12	2
silage and animal feed	phytases	<i>Aspergillus niger</i>	8	3
paper and textiles	$\alpha$ -amylase, lipase	<i>Bacillus</i> , <i>Humicola</i>	2	4
leather treatment	proteases	<i>Aspergillus oryzae</i>	10	1, 7

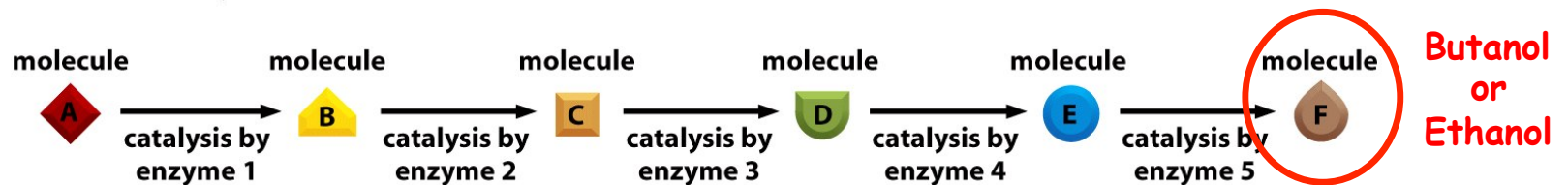
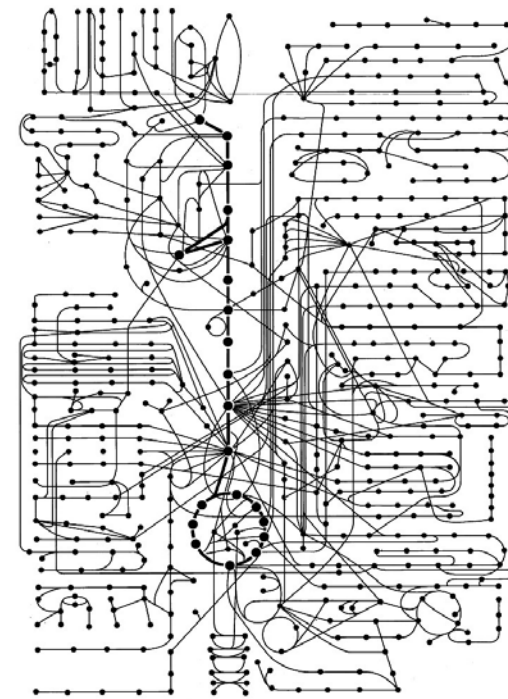
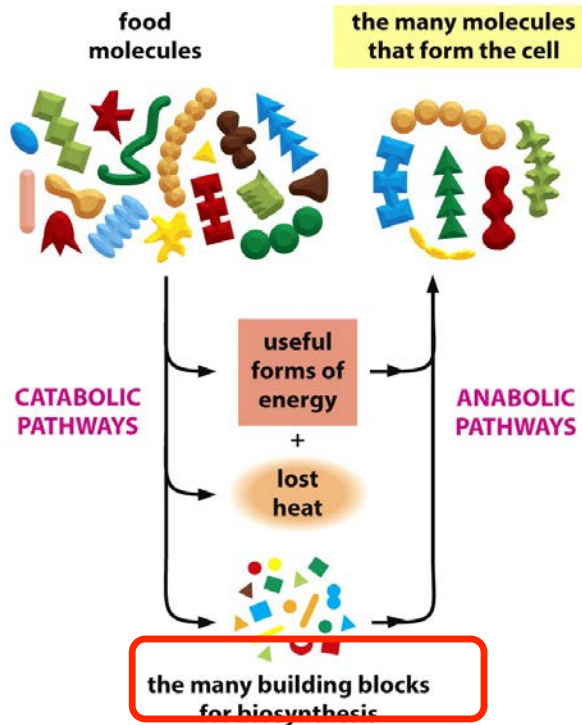


process/application	enzyme cost per unit quantity (US \$)	important goals in application technology
starch liquefaction	ca. \$ 2 per t starch	1 higher product quality
glucose from starch	\$ 3.5 per t starch	2 improved taste
isomerization of glucose	\$ 6 per t starch	3 better yields
HFS in USA	\$ 6–7 per t starch	4 reduced process costs
ethanol	\$ 1 per t starch	5 better filtration
beer	\$ 0.1 per 100L	6 better conservation
bakery goods USA	\$ 0.1 per 100 kg flour	7 improved working conditions, reduced environmental load
bakery goods EU	\$ 0.1–0.5 per 100 kg flour	
fruit juice	\$ 0.1–0.5 per 100L juice	
wine	\$ 0.1–0.5 per 100L wine	
stabilization of fruit		
lemonade by glucose oxidase	\$ 0.3–0.8 per 1000L	
cheese manufacture	\$ 0.05 per 100L milk	
detergents	\$ 0.05 per kg detergent	
leather tanning	\$ 1.2–3 per t skin	

Specific Proteins and/or Metabolic Pathways Can Be Improved and/or Manipulated By Recombinant DNA!



# Metabolites Are Produced By Cellular Pathways That Use Specific Enzymes and Genes To Synthesize Specific Small Molecules



ABBREVIATED AS





# Engineering *E.coli* Pathways To Make BioFuel

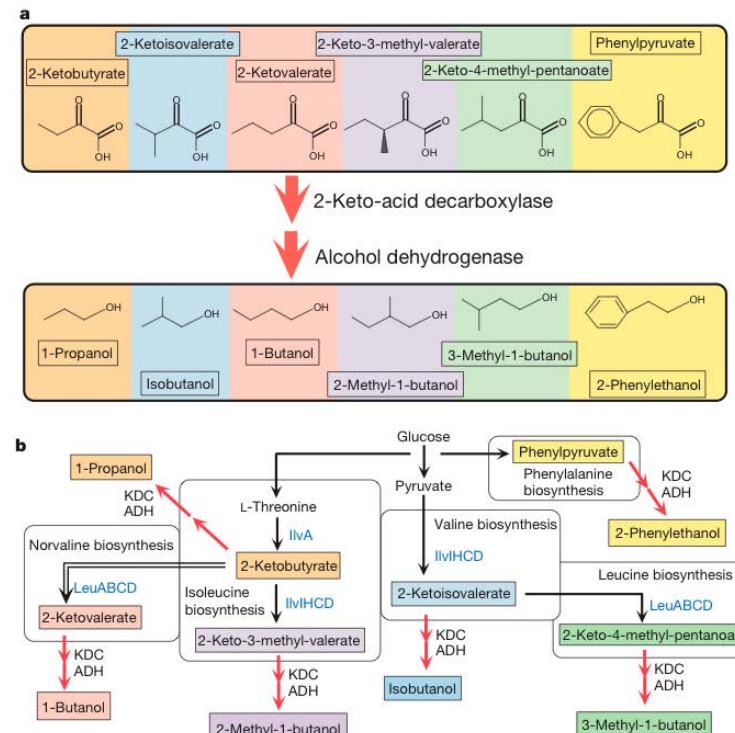
nature

Vol 451 | 3 January 2008 | doi:10.1038/nature06450

## LETTERS

### Non-fermentative pathways for synthesis of branched-chain higher alcohols as biofuels

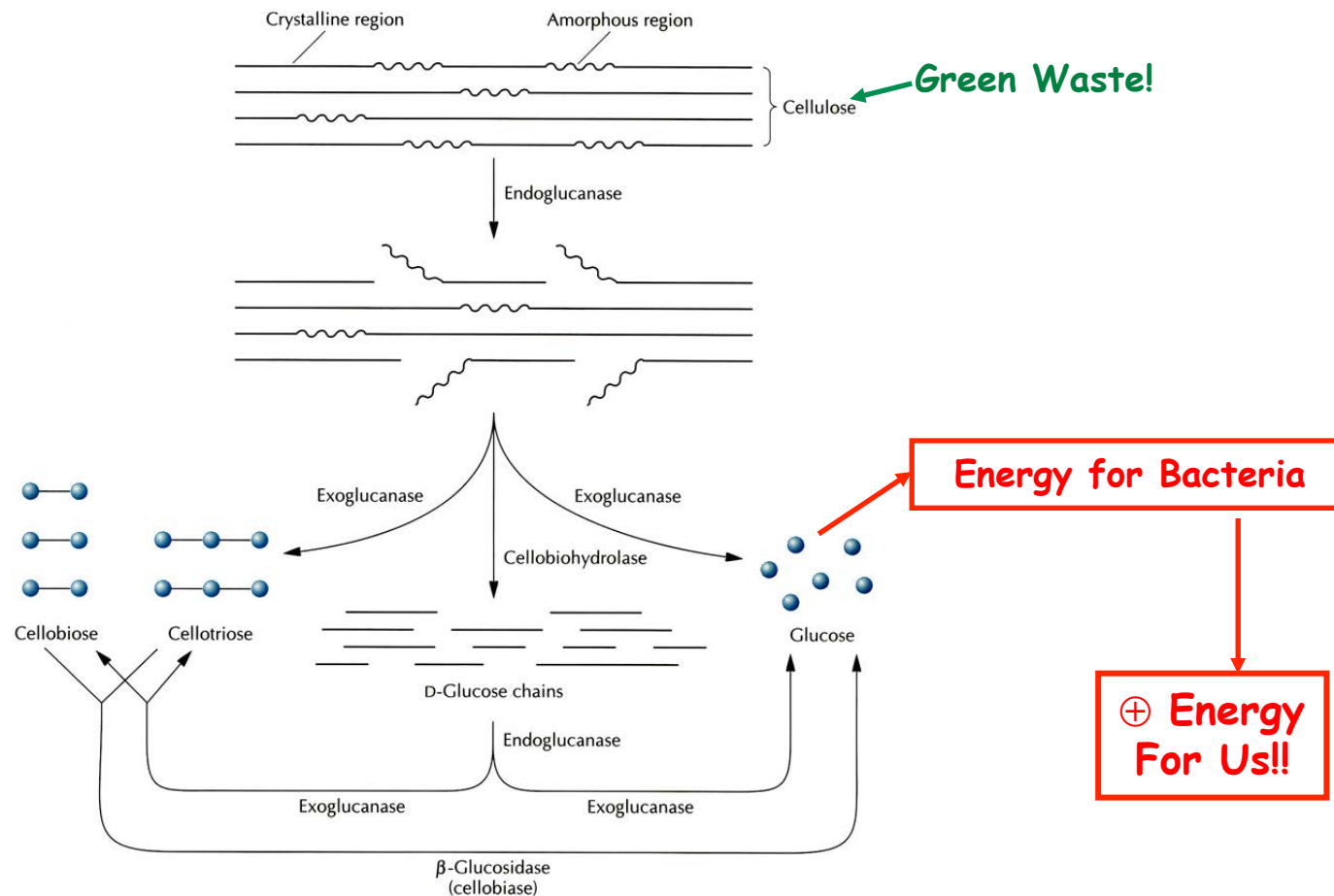
Shota Atsumi<sup>1</sup>, Taizo Hanai<sup>1</sup> & James C. Liao<sup>1,2</sup>



**Figure 1 | Production of higher alcohols through the synthetic non-fermentative pathways. a,** Various 2-keto acid precursors lead to corresponding alcohols through 2-ketoacid decarboxylase and alcohol dehydrogenase. **b,** The synthetic networks for the non-fermentative alcohol

production in engineered *E. coli*. Red arrows represent the 2-keto acid decarboxylation and reduction pathway. Blue enzyme names represent amino acid biosynthesis pathways. The double lines represent a side pathway leading to norvaline and 1-butanol biosynthesis.

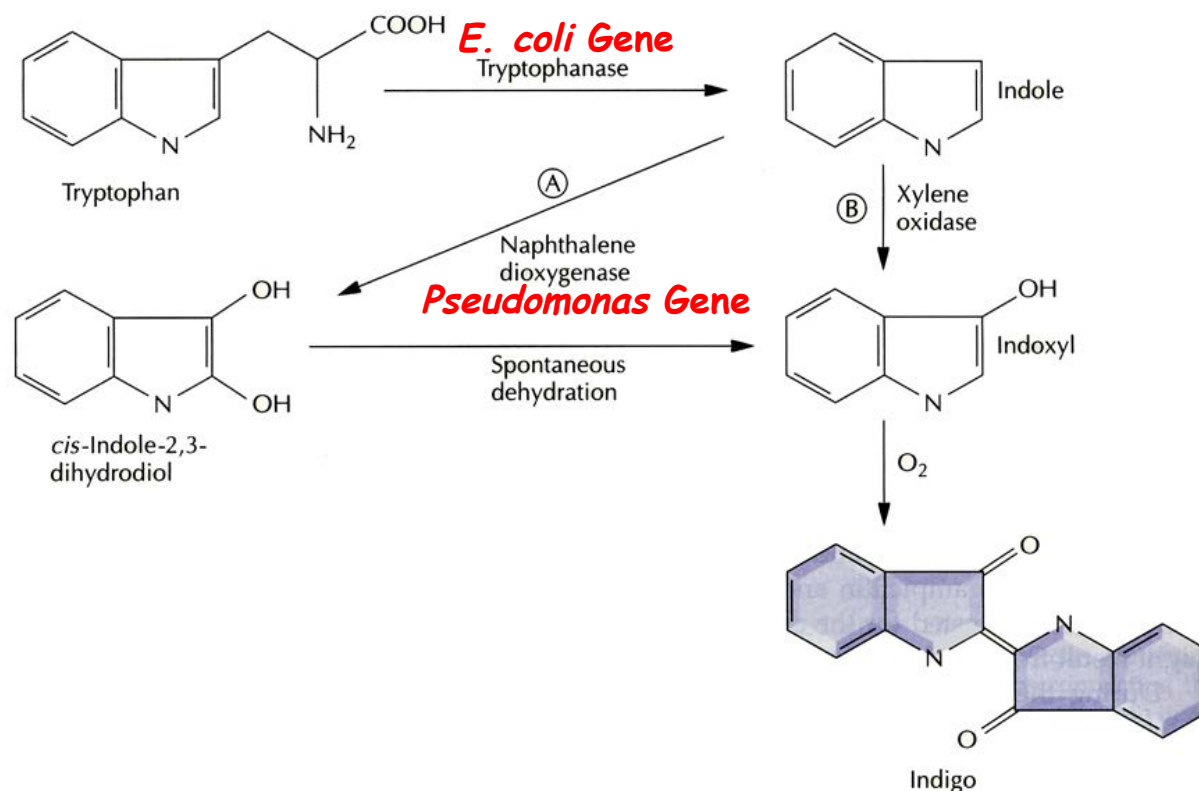
# Bacteria Can Be Engineered To Degrade Biomass Waste-Containing Cellulose (e.g., paper)



**Figure 13.27** Enzymatic biodegradation of cellulose. Cellulose hydrolysis begins with the cleavage of  $\beta$ -1,4-linkages within the accessible amorphous regions of the cellulose chains by endoglucanase(s). This reaction is followed by the removal of oligosaccharides from the reducing ends of the partially cleaved cellulose chains by exoglucanase(s) and cellobiohydrolase(s). The degradation of cellulose is completed when the cellobiose and cellotriose are converted to glucose by  $\beta$ -glucosidase.

**Agriculture, Timber Processing, Human Activities: e.g., Plants Left Over From Harvests, Animal Manure With Grasses, Municipal Water Paper, Cotton Leftovers, Hay, Etc.**

# Engineering *E. coli* To Synthesize Indigo- The Major Blue Dye For Jeans & Other Clothes & Uses



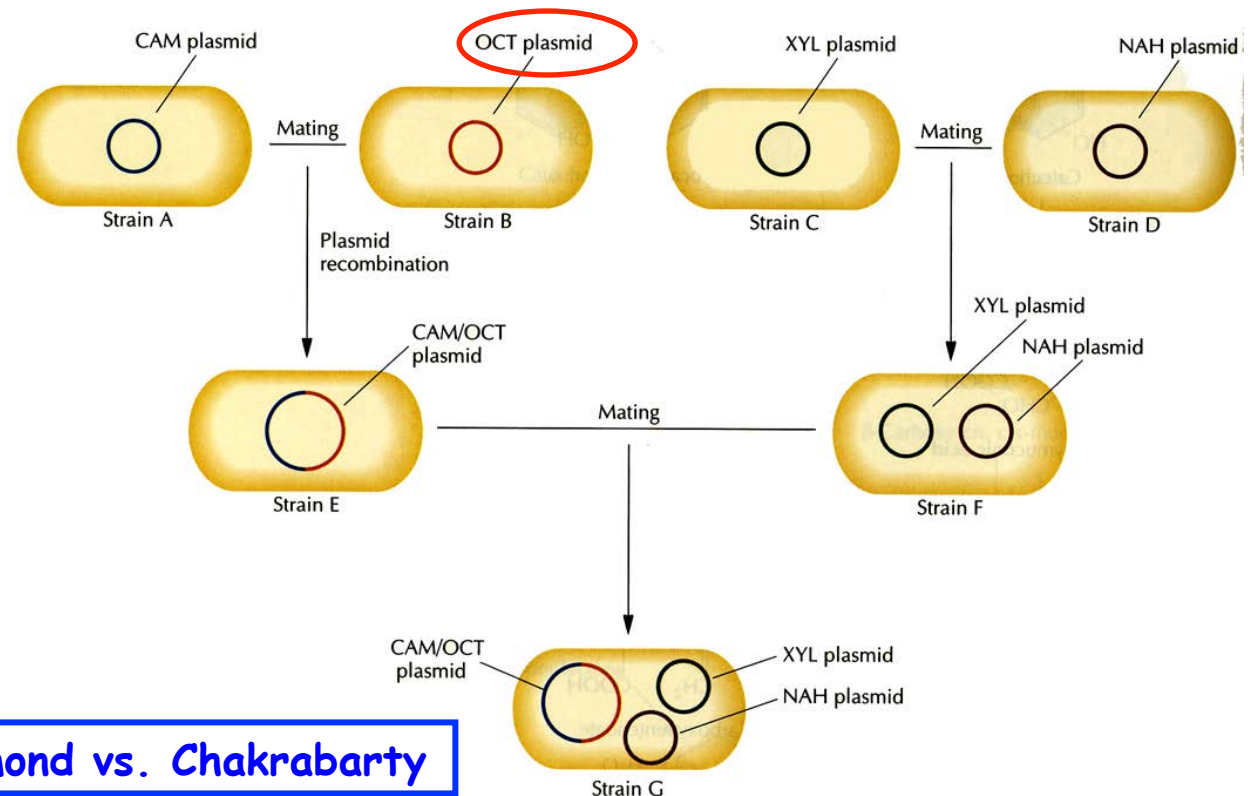
**Figure 12.8** Indigo biosynthesis from tryptophan in genetically engineered *E. coli*. Tryptophanase is an *E. coli* enzyme. In pathway A, the naphthalene dioxygenase is derived from the NAH plasmid; in pathway B, the xylene oxidase is from the TOL plasmid. *E. coli* transformants that synthesize indigo contain either pathway A or B but not both pathways.

**\$200M/Year Industry**  
**Indigo Previously Obtained From Plants!**



# Bacteria Can Be Engineered To Degrade Several Different “Toxic” Compounds

**Pseudomonas**



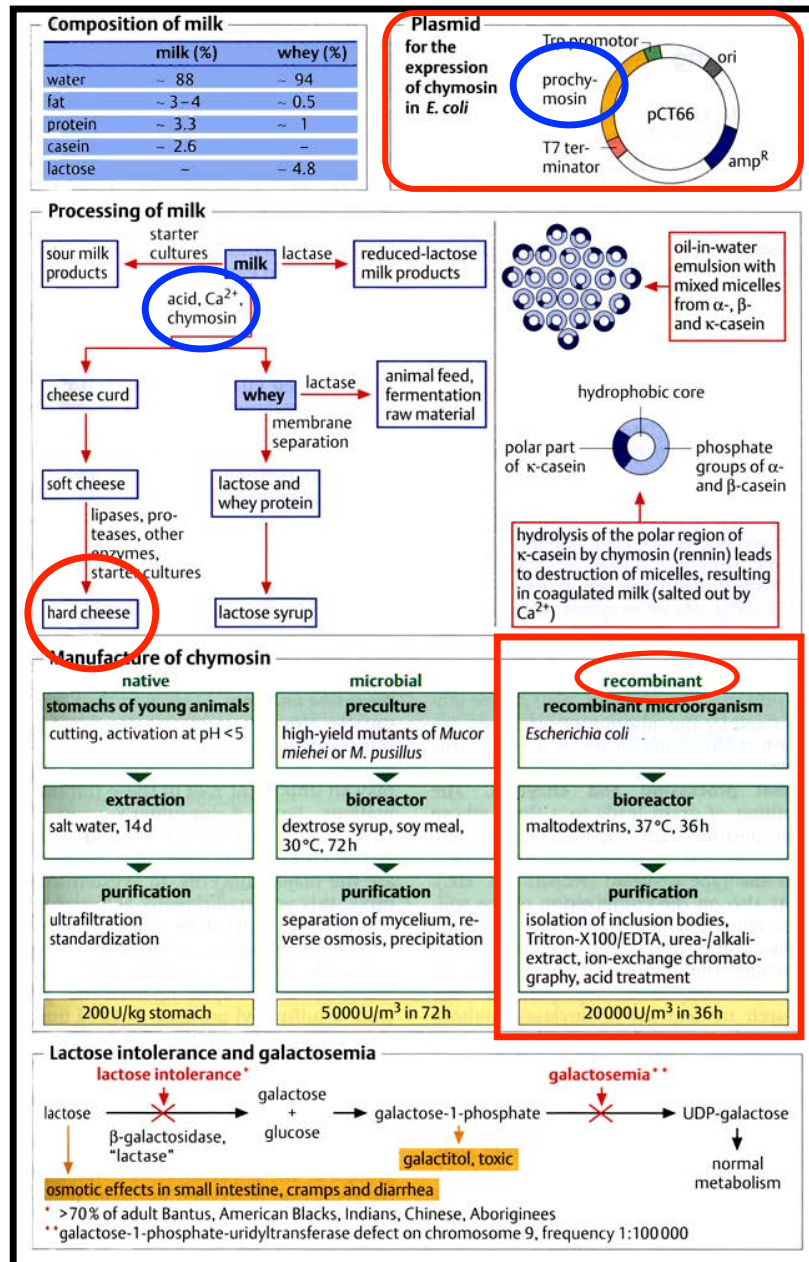
**A Landmark Decision- Diamond vs. Chakrabarty**

**Chakrabarty US Patent 4,259,444 1981  
Genetically Engineered Microorganisms  
Are “Inventions”**

**Life Can Be Patented !**

*Figure 13.5* Schematic representation of the development of a bacterial strain that can degrade camphor, octane, xylene, and naphthalene. Strain A, which contains a CAM (camphor-degrading) plasmid, is mated with strain B, which carries an OCT (octane-degrading) plasmid. Following plasmid transfer and homologous recombination between the two plasmids, strain E carries a CAM and OCT biodegradative fusion plasmid. Strain C, which contains a XYL (xylene-degrading) plasmid, is mated with strain D, which contains a NAH (naphthalene-degrading) plasmid, to form strain F, which carries both of these plasmids. Finally, strains E and F are mated to yield strain G, which carries the CAM/OCT fusion plasmid, the XYL plasmid, and the NAH plasmid.

# Recombinant Chymosin Is Used To Make Cheese



**Chymosin (Rennin)  
Acts On Milk  
Proteins To  
Coagulate Milk →  
Cheese**

**Is Cheese A GMO?**



# Chymosin In Cheese Making

1. ~80-90% of Cheeses Are Made With Recombinant Chymosin (a Protease)
2. Approved For Use In Cheese Making By FDA - 1992
3. Not Different From Non-Recombinant Chymosin-  
∴ GRAS- Generally Regarded As Safe & No Labeling Needed — Because Not An Additive & Not Different From Non-Recombinant Chymosin!!

Is Cheese Made Using Recombinant Chymosin a GMO?

Industry Adds Claim That Recombinant Chymosin is “Kosher” & “Vegetarian”



Why No Fuss?

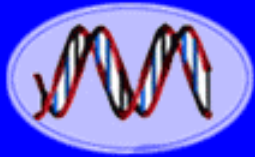




## FDA Approval of Cheese Made With Recombinant Chymosin

**Extraordinary precautions** were taken before chymosin, made by recombinant DNA technology, was marketed. Regulators ensured that no toxins of any kind had been introduced and that no live recombinant organisms were present. Indeed, the product contained nothing but pure chymosin. Cheese made with it is completely indistinguishable from that produced with animal rennet. In any case, chymosin itself is degraded during cheese making and none is left in the finished product. **Today, in North America, over 80 percent of all cheese is made using chymosin produced by recombinant DNA technology.** Cheese makers no longer have to worry about a shortage of calf stomachs and turophiles can satisfy their critical tastebuds. Thanks to biotechnology they can "say cheese" and smile.

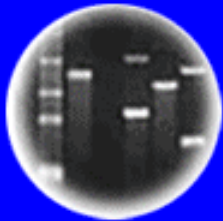




DNA  
Genetic Code of Life



Entire Genetic Code  
of a Bacteria



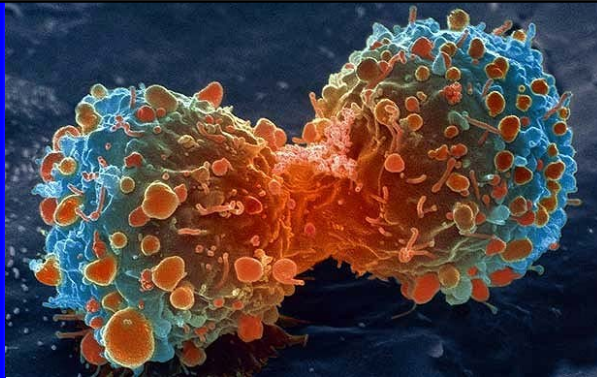
DNA Fingerprinting



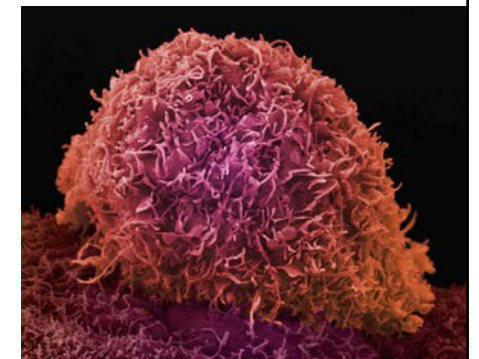
Cloning: Ethical Issues  
and Future Consequences



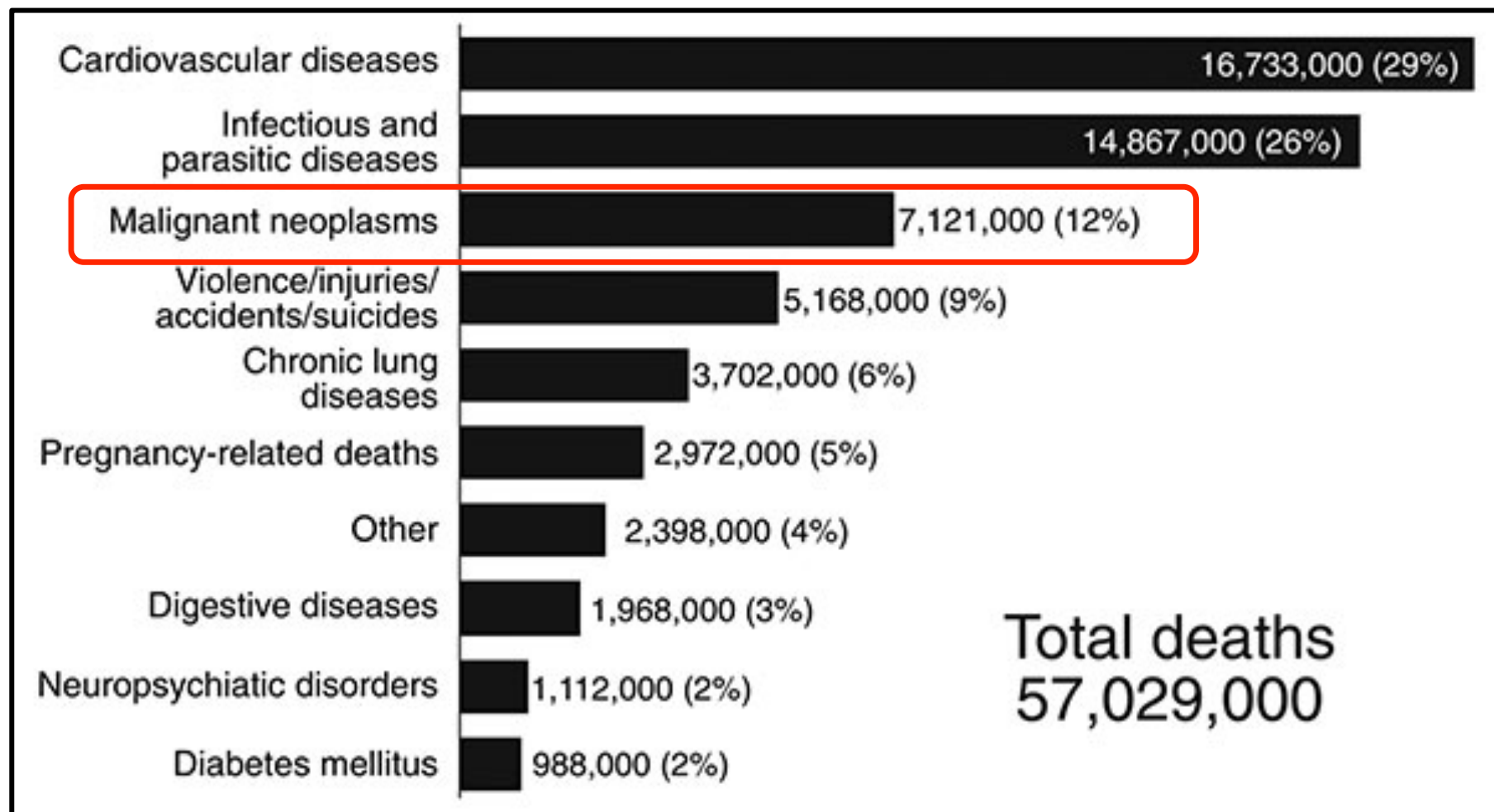
Plants of Tomorrow



# Cancer Treatment & Discovery



## Over 10% of Annual Deaths Word-Wide Are Caused By Various Types of Cancer





# Frequency of Different Cancer Types

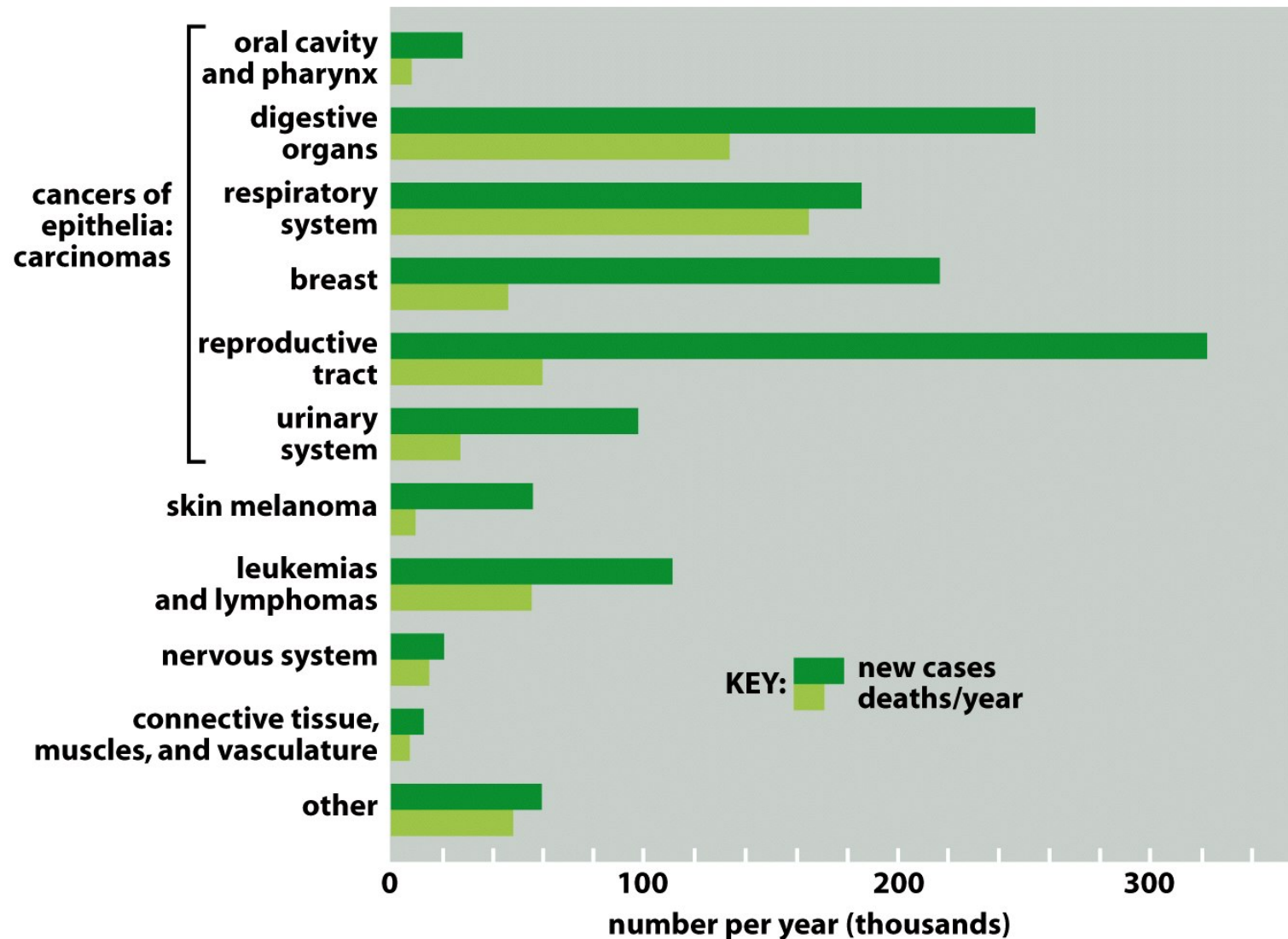


Figure 20-2 *Molecular Biology of the Cell* (© Garland Science 2008)

# The Frequency of Cancer Increases With Age

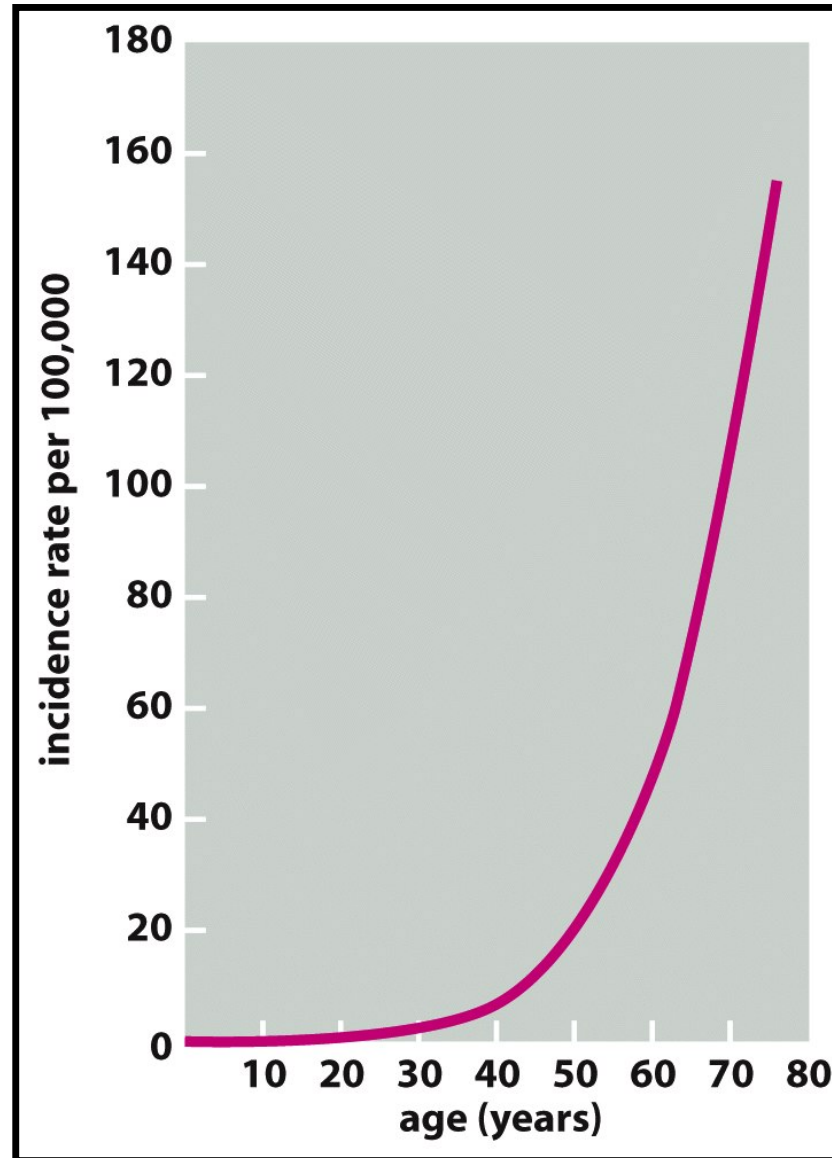
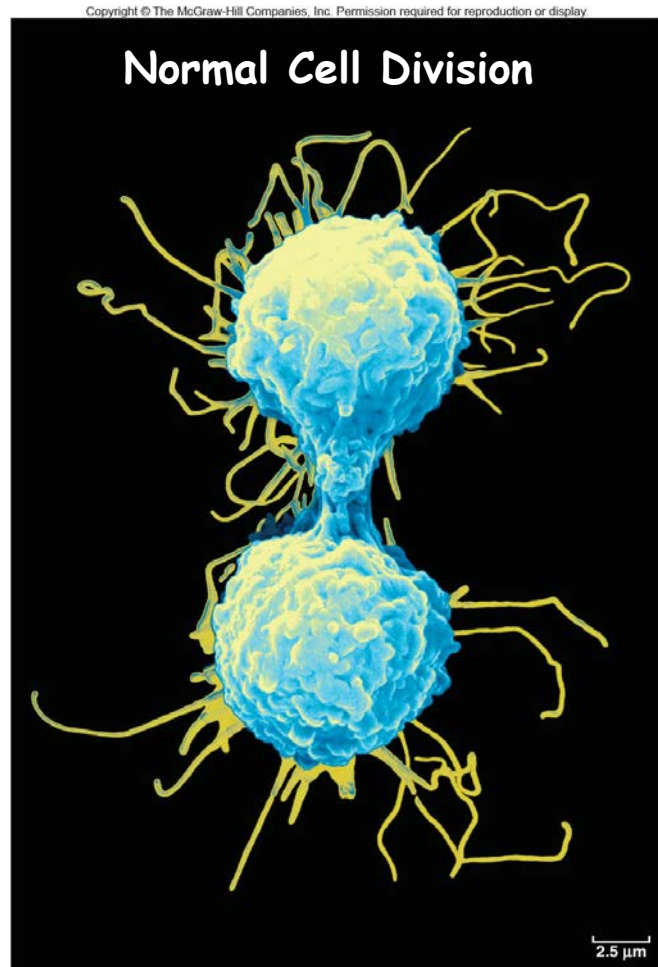
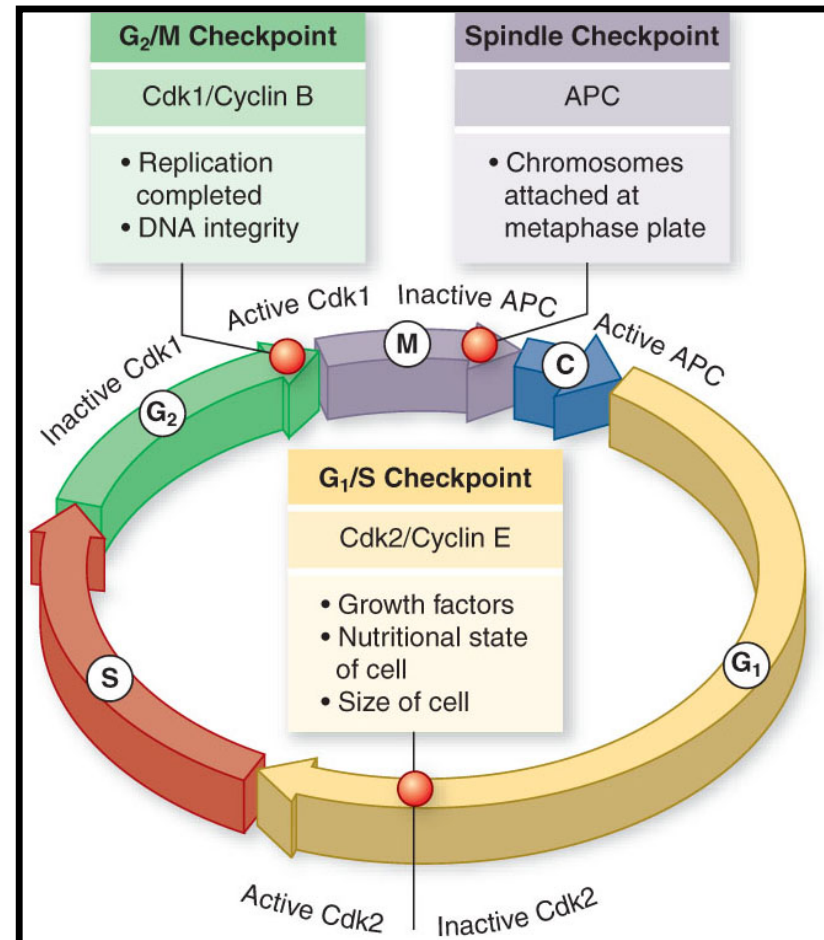


Figure 20-7 *Molecular Biology of the Cell* (© Garland Science 2008)

# Vaccines Can Also Be Made To Treat Cancer - Checking Abnormal Cell Division



## Normal Cell Cycle





# Check Points Controlling Cell Division

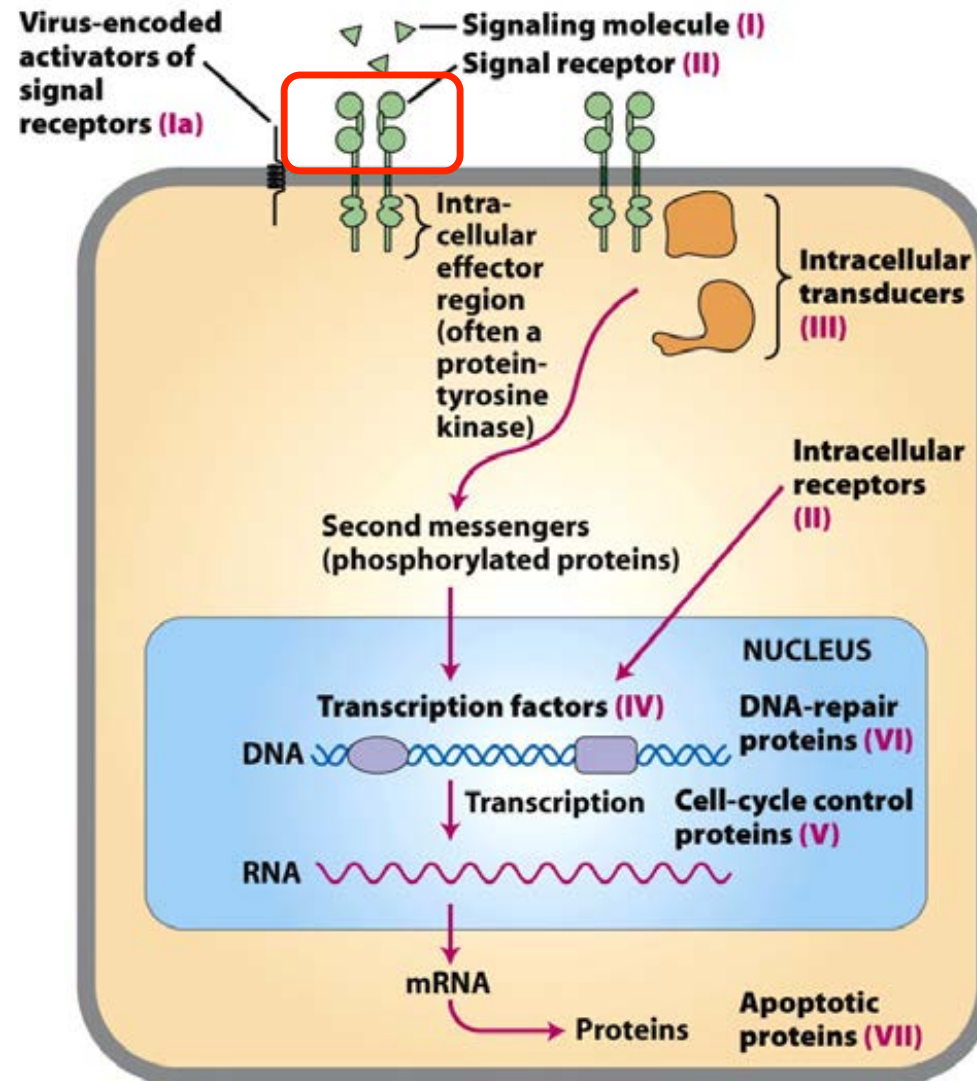


Figure 25-11  
*Molecular Cell Biology, Sixth Edition*  
© 2008 W. H. Freeman and Company

# Mutations in Check Point Genes/Proteins Lead To Cancer

- Cancer is a "Gene Mutation" Disease

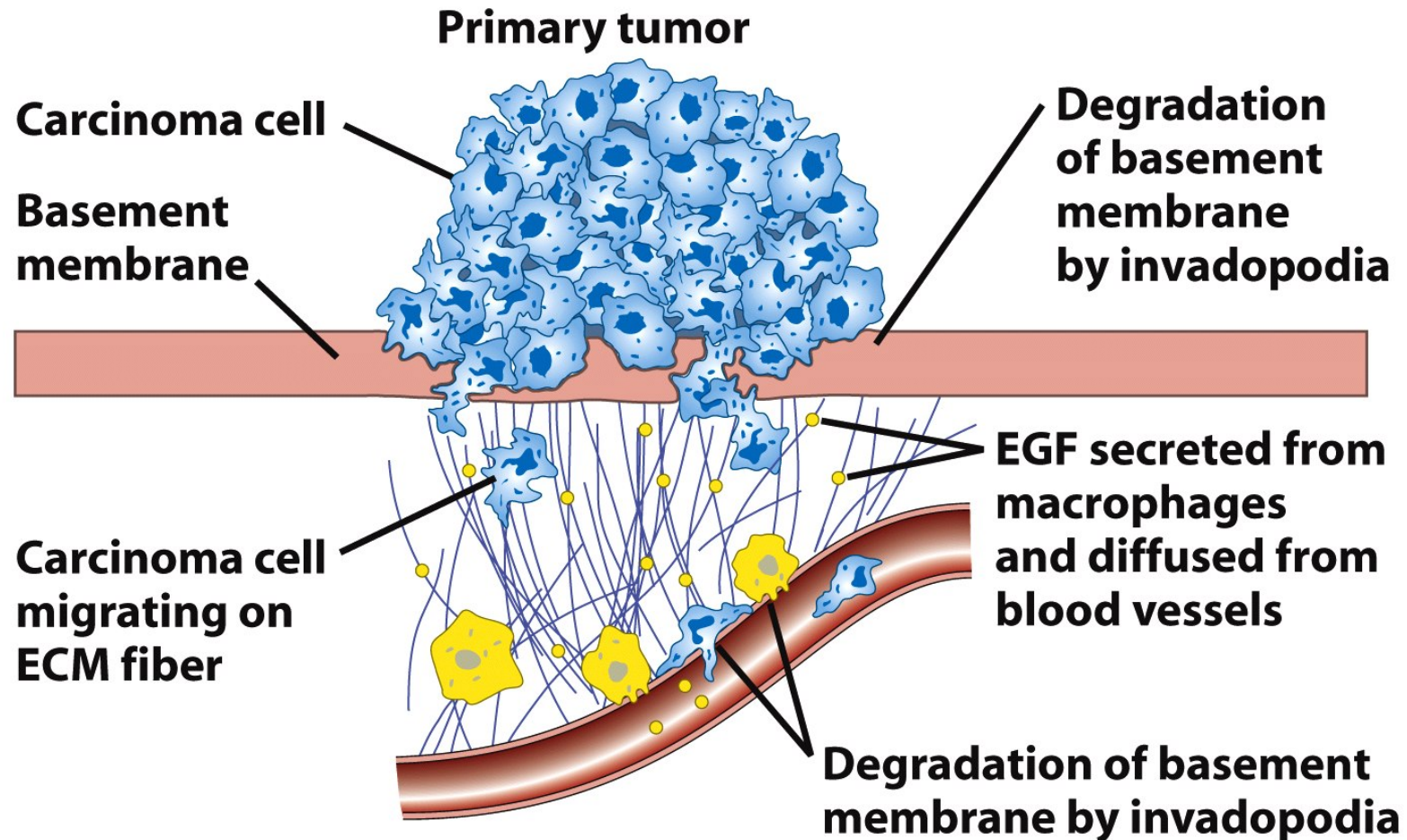
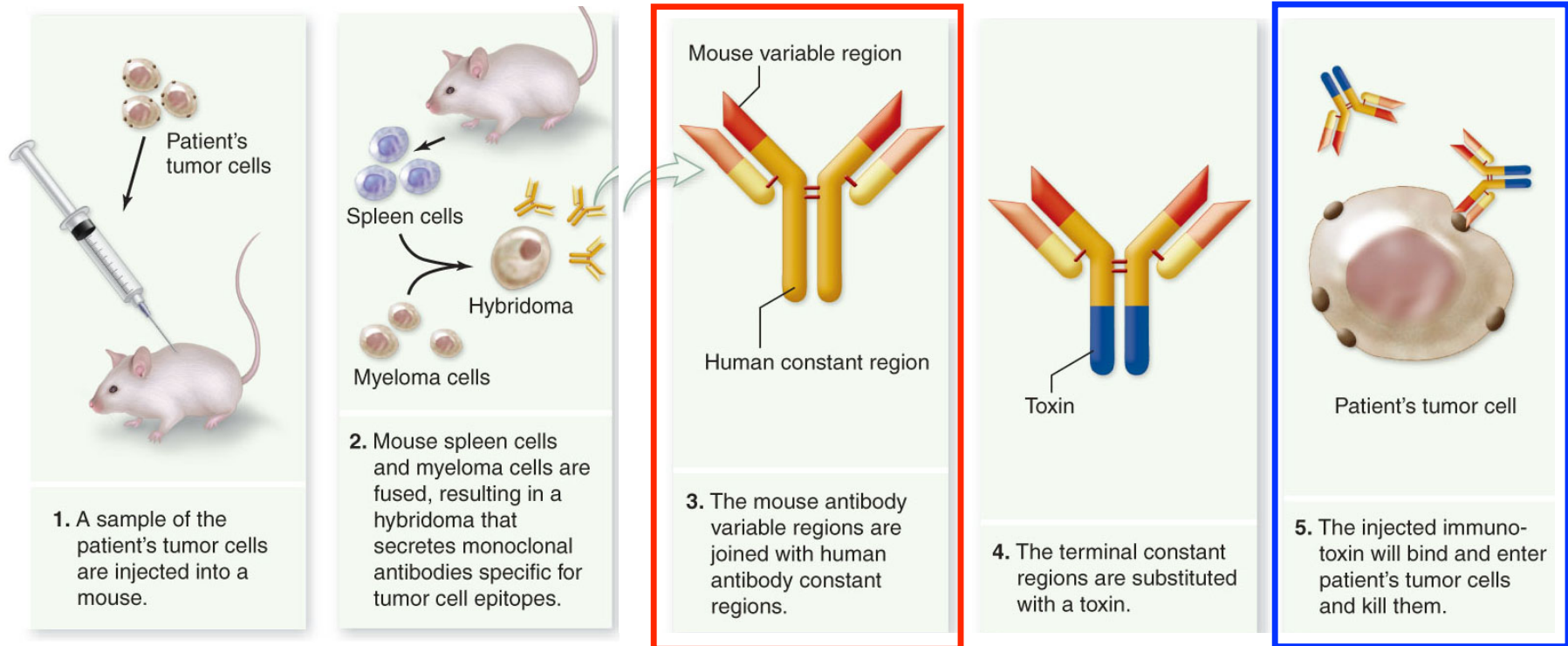
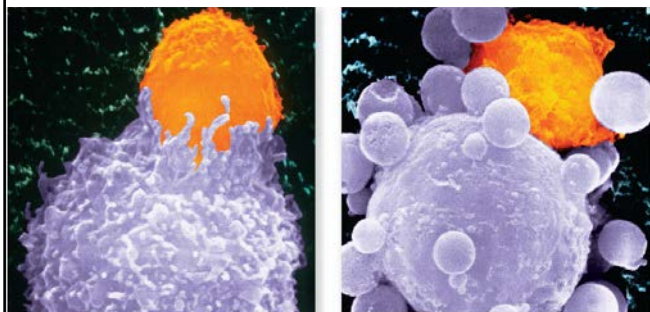


Figure 25-3a  
*Molecular Cell Biology, Sixth Edition*  
© 2008 W. H. Freeman and Company

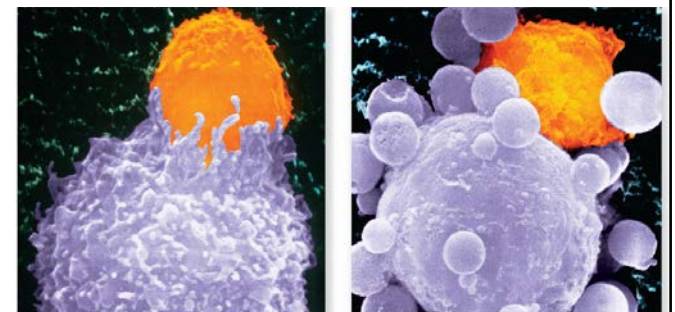
# Recombinant Vaccines Are Being Developed To Fight Cancer



**Genetic Engineering Step**

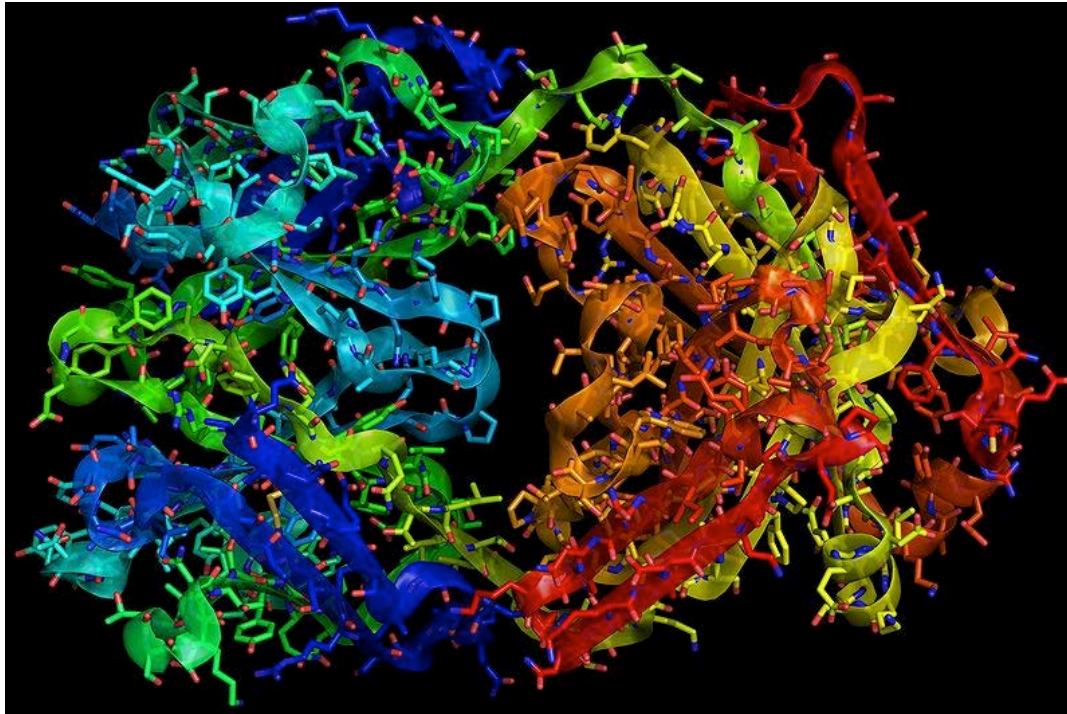


**Cancer Cell Being  
Destroyed By T-Cell  
Containing Cancer-  
Cell-Specific Antibody**





# Using Herceptin® to Treat Breast Cancer



Trastuzumab® or Herceptin®

Dr. Dennis Slamon, UCLA Jonsson Cancer Treatment Center

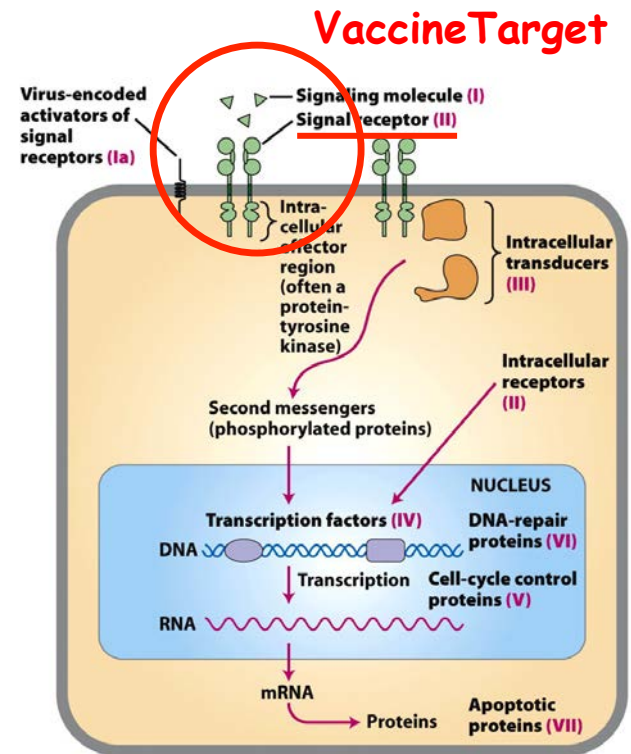
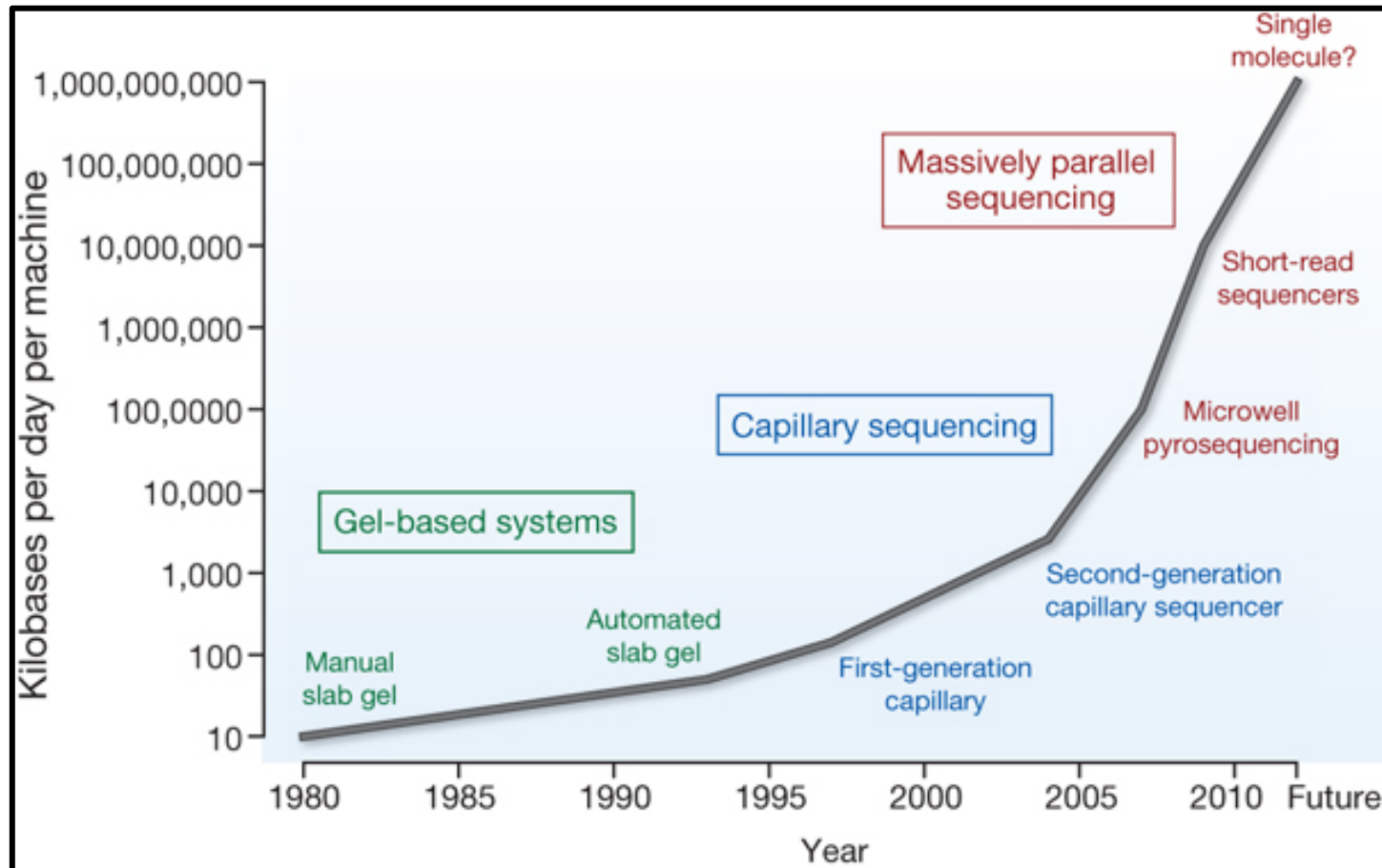


Figure 25-11  
Molecular Cell Biology, Sixth Edition  
© 2008 W. H. Freeman and Company



# The Cancer Genome Project Is Mapping Tumor-Specific Genes To Find Drug Targets



*Sequencing Costs Have Dropped Exponentially Allowing the Genome Sequence of Specific Tumors - As They Progress - To Be Sequenced In Order To Identify The Mutated Genes Causing the Cancer*



# **A small-cell lung cancer genome with complex signatures of tobacco exposure**

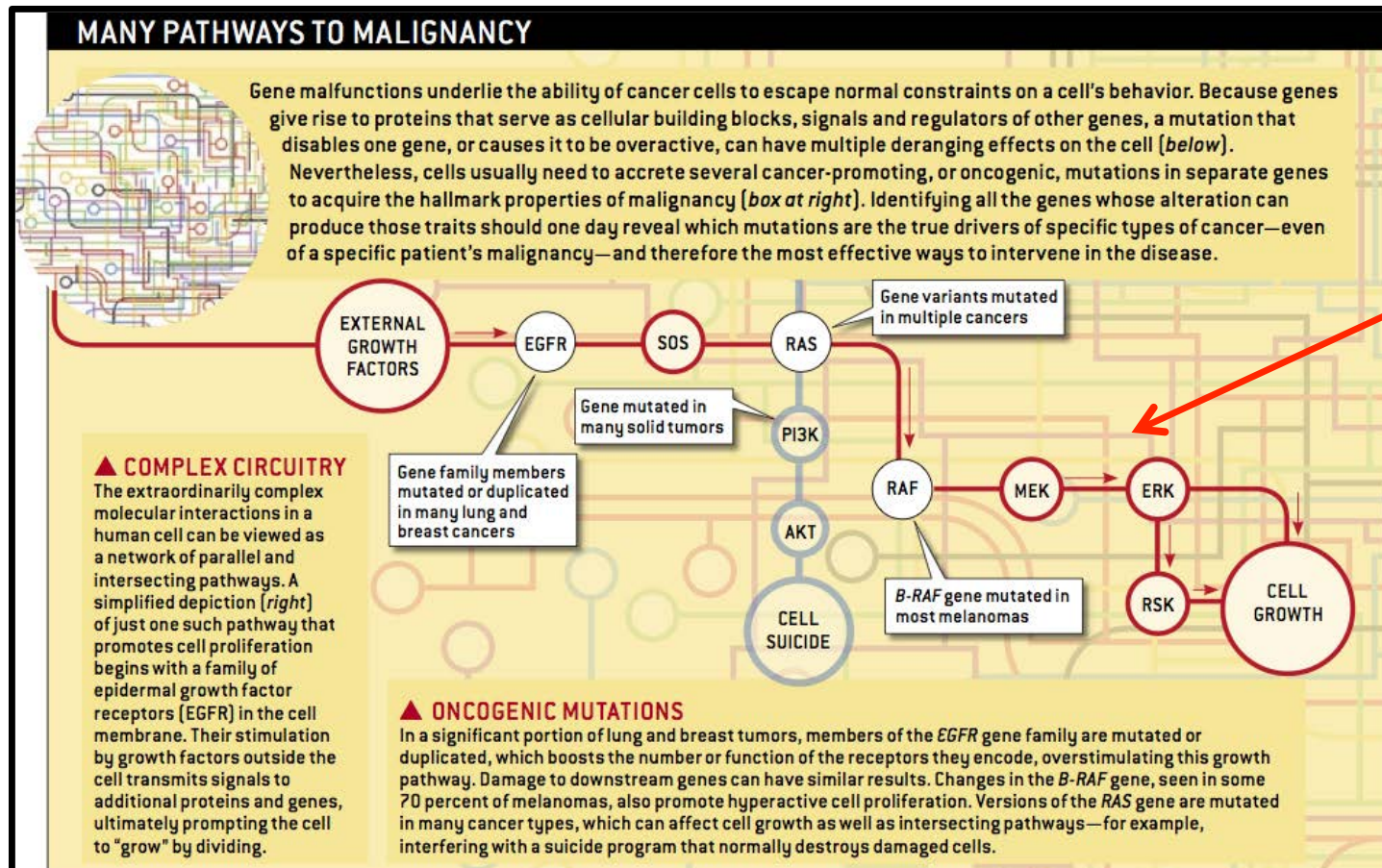
Erin D. Pleasance<sup>1</sup>, Philip J. Stephens<sup>1</sup>, Sarah O'Meara<sup>1,2</sup>, David J. McBride<sup>1</sup>, Alison Meynert<sup>3</sup>, David Jones<sup>1</sup>, Meng-Lay Lin<sup>1</sup>, David Beare<sup>1</sup>, King Wai Lau<sup>1</sup>, Chris Greenman<sup>1</sup>, Ignacio Varela<sup>1</sup>, Serena Nik-Zainal<sup>1</sup>, Helen R. Davies<sup>1</sup>, Gonzalo R. Ordoñez<sup>1</sup>, Laura J. Mudie<sup>1</sup>, Calli Latimer<sup>1</sup>, Sarah Edkins<sup>1</sup>, Lucy Stebbings<sup>1</sup>, Lina Chen<sup>1</sup>, Mingming Jia<sup>1</sup>, Catherine Leroy<sup>1</sup>, John Marshall<sup>1</sup>, Andrew Menzies<sup>1</sup>, Adam Butler<sup>1</sup>, Jon W. Teague<sup>1</sup>, Jonathon Mangion<sup>2</sup>, Yongming A. Sun<sup>4</sup>, Stephen F. McLaughlin<sup>5</sup>, Heather E. Peckham<sup>5</sup>, Eric F. Tsung<sup>5</sup>, Gina L. Costa<sup>5</sup>, Clarence C. Lee<sup>5</sup>, John D. Minna<sup>6</sup>, Adi Gazdar<sup>6</sup>, Ewan Birney<sup>3</sup>, Michael D. Rhodes<sup>4</sup>, Kevin J. McKernan<sup>5</sup>, Michael R. Stratton<sup>1,7</sup>, P. Andrew Futreal<sup>1</sup> & Peter J. Campbell<sup>1,8</sup>

Cancer is driven by mutation. Worldwide, tobacco smoking is the principal lifestyle exposure that causes cancer, exerting carcinogenicity through >60 chemicals that bind and mutate DNA. Using massively parallel sequencing technology, we sequenced a small-cell lung cancer cell line, NCI-H209, to explore the mutational burden associated with tobacco smoking. A total of 22,910 somatic substitutions were identified, including 134 in coding exons. Multiple mutation signatures testify to the cocktail of carcinogens in tobacco smoke and their proclivities for particular bases and surrounding sequence context. Effects of transcription-coupled repair and a second, more general, expression-linked repair pathway were evident. We

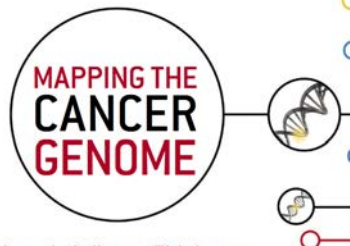
# **A comprehensive catalogue of somatic mutations from a human cancer genome**



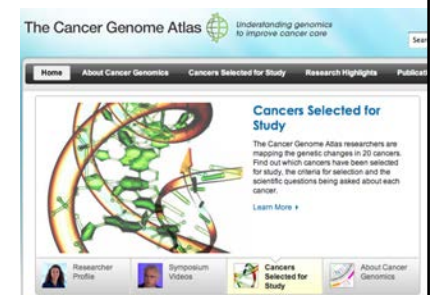
# The Cancer Genome Project Is Mapping Tumor-Specific Genes To Find Drug Targets

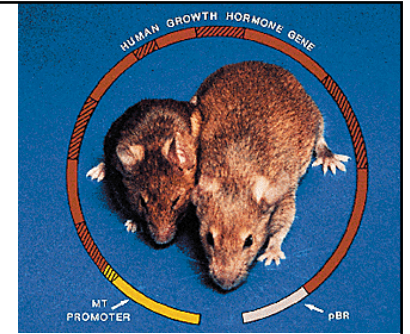


Possible  
Drug  
Targets



Pinpointing the genes involved in cancer will help chart a new course across the complex landscape of human malignancies



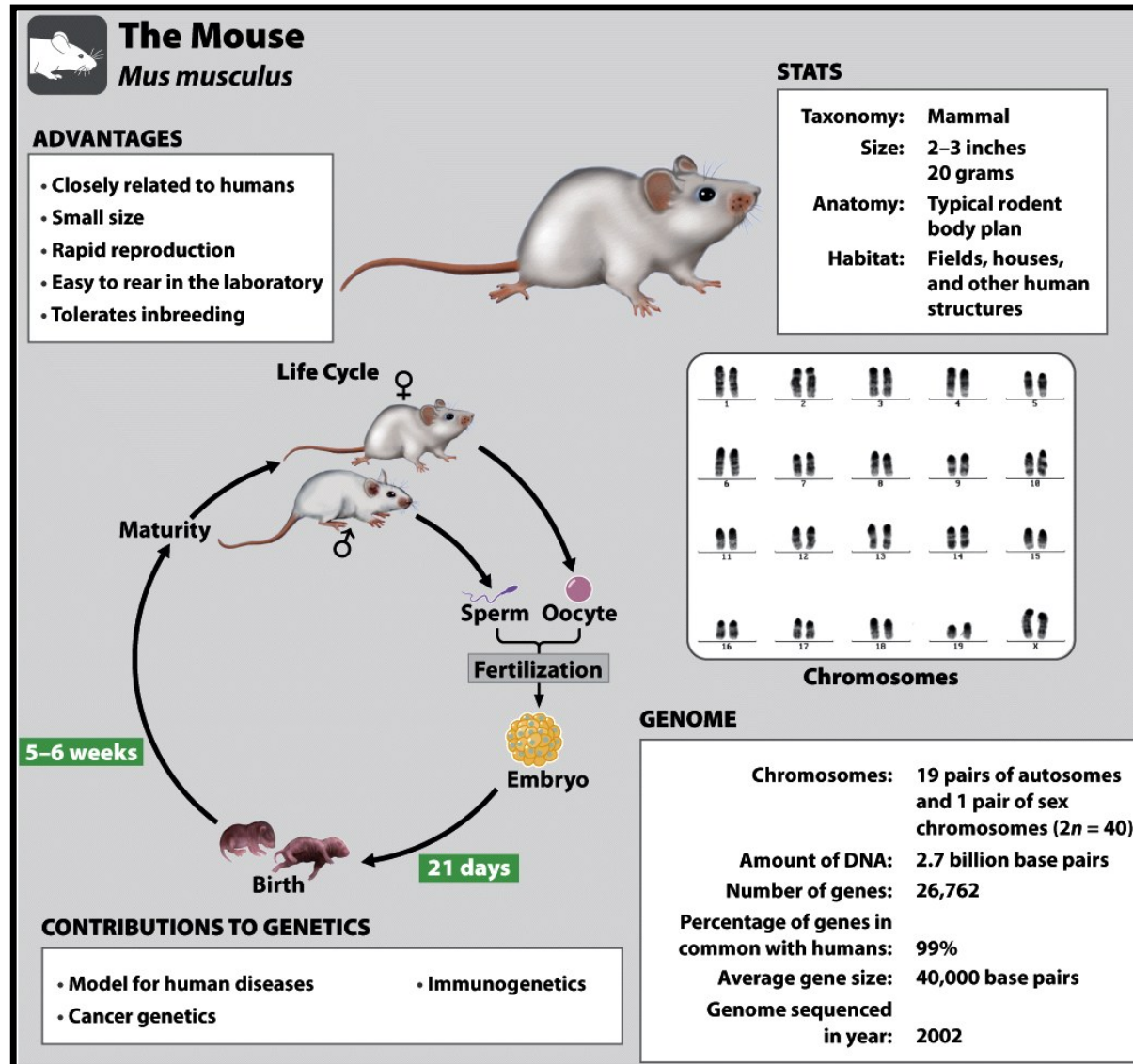


## Animals (and Plants) Can Also Be Engineered For Applied Purposes





# Using the Mouse as a Model to Determine the Function of Human Genes



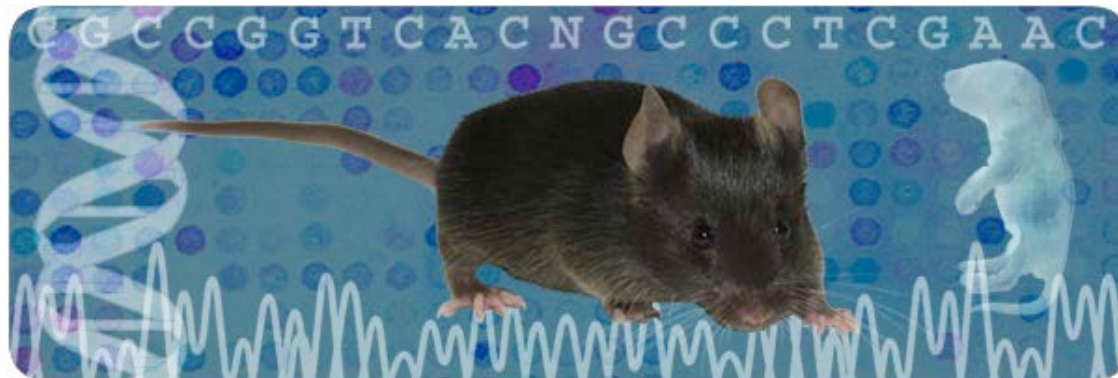
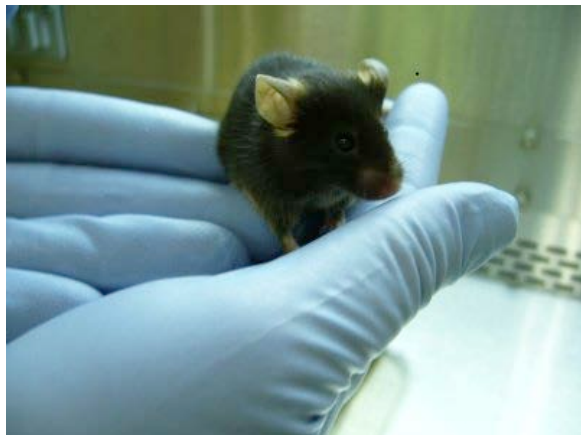
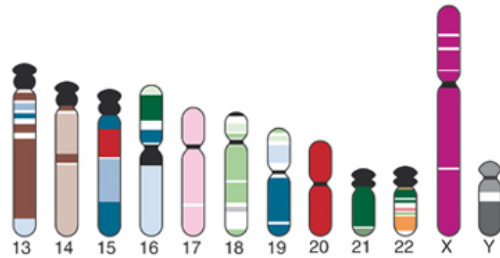
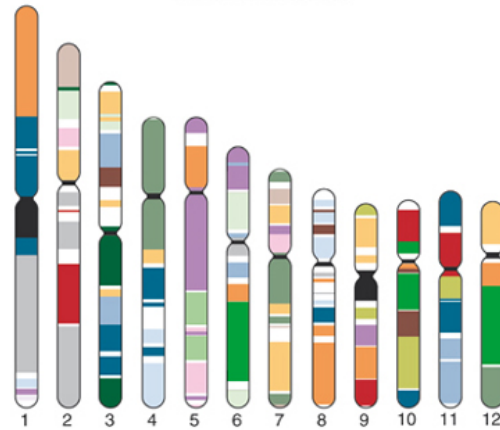


# Human and Mouse Genomes 99% Similar

## ∴ Can Study Human Genes Using Mouse as a Model

Copyright © The McGraw-Hill Companies, Inc. Permission required for reproduction or display.

Human chromosomes



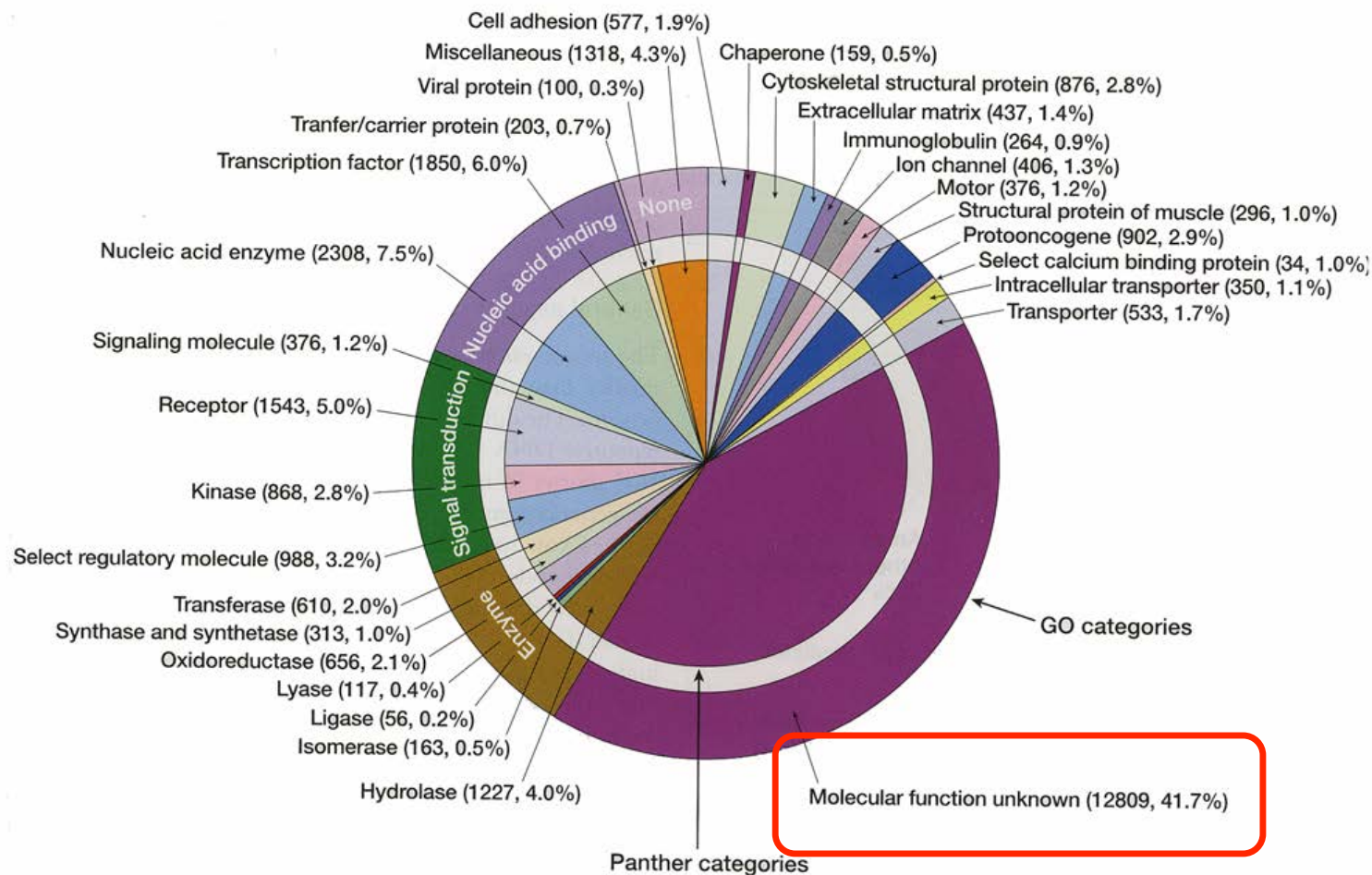
# How Many Human Disease Genes Have Been Identified?

The screenshot shows the OMIM website interface. At the top, there is a navigation bar with links to All Databases, PubMed, Nucleotide, Protein, Genome, Structure, PMC, and OMIM. The OMIM logo is prominently displayed, along with the text "Online Mendelian Inheritance in Man" and the Johns Hopkins University logo. A search bar is located at the top left, with a dropdown menu set to "OMIM" and a "for" field. Below the search bar, there are buttons for "Limits", "Preview/Index", "History", "Clipboard", and "Details". On the left side, there is a sidebar with links to Entrez, OMIM, Search OMIM, Search Gene Map, Search Morbid Map, Help, OMIM Help, How to Link, FAQ, Numbering System, Symbols, and How to Print. The main content area contains a list of search instructions: "Enter one or more search terms.", "Use Limits to restrict your search by search field, chromosome, and other criteria.", "Use Index to browse terms found in OMIM records.", and "Use History to retrieve records from previous searches, or to combine searches." Below this, there is a section titled "OMIM® - Online Mendelian Inheritance in Man®" with a welcome message: "Welcome to OMIM®, Online Mendelian Inheritance in Man®. OMIM is a comprehensive, authoritative, and timely compendium of human genes and genetic phenotypes. The full-text, referenced overviews in OMIM contain information on all known mendelian disorders and over 12,000 genes. OMIM focuses on the relationship between phenotype and genotype. It is updated daily, and the entries contain copious links to other genetics resources."

**There are ~25,000 Genes in The Human Genome**

1. ~4034 Genes Correlate With a Disease Phenotype
2. The Molecular Basis of These Genetic Diseases Are Known (e.g., Sickle Cell Anemia, Hemophilia A)
3. ~1717 Disease Genes - Molecular Basis Unknown

# The Human Genome Contains ~25,000 Different Genes

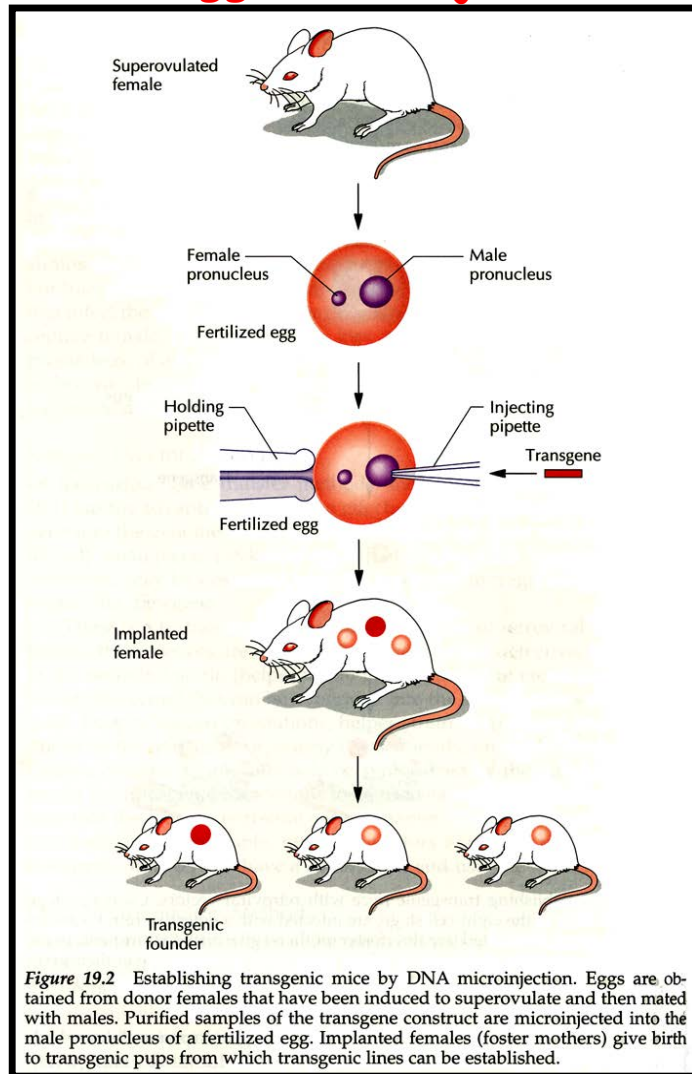


**Do Not Know Functions of  
Most Human Genes!**  
**How Find?**

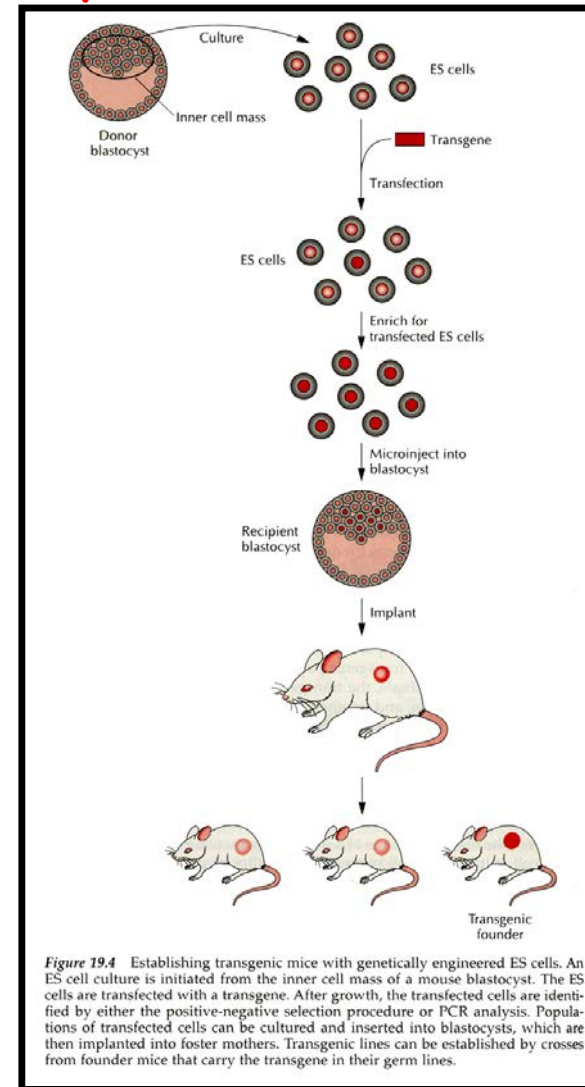


# Genetic Engineering Mice and Other Mammals

## 1. Egg DNA Injection



## 2. Embryo Stem Cell DNA Transformation



**Both Methods Alter the Germ Line (i.e., Genes are Inherited)**

# Mouse Genes Related to Those in the Human Genome Can be “Knocked Out” In Order to Determine Their Functions

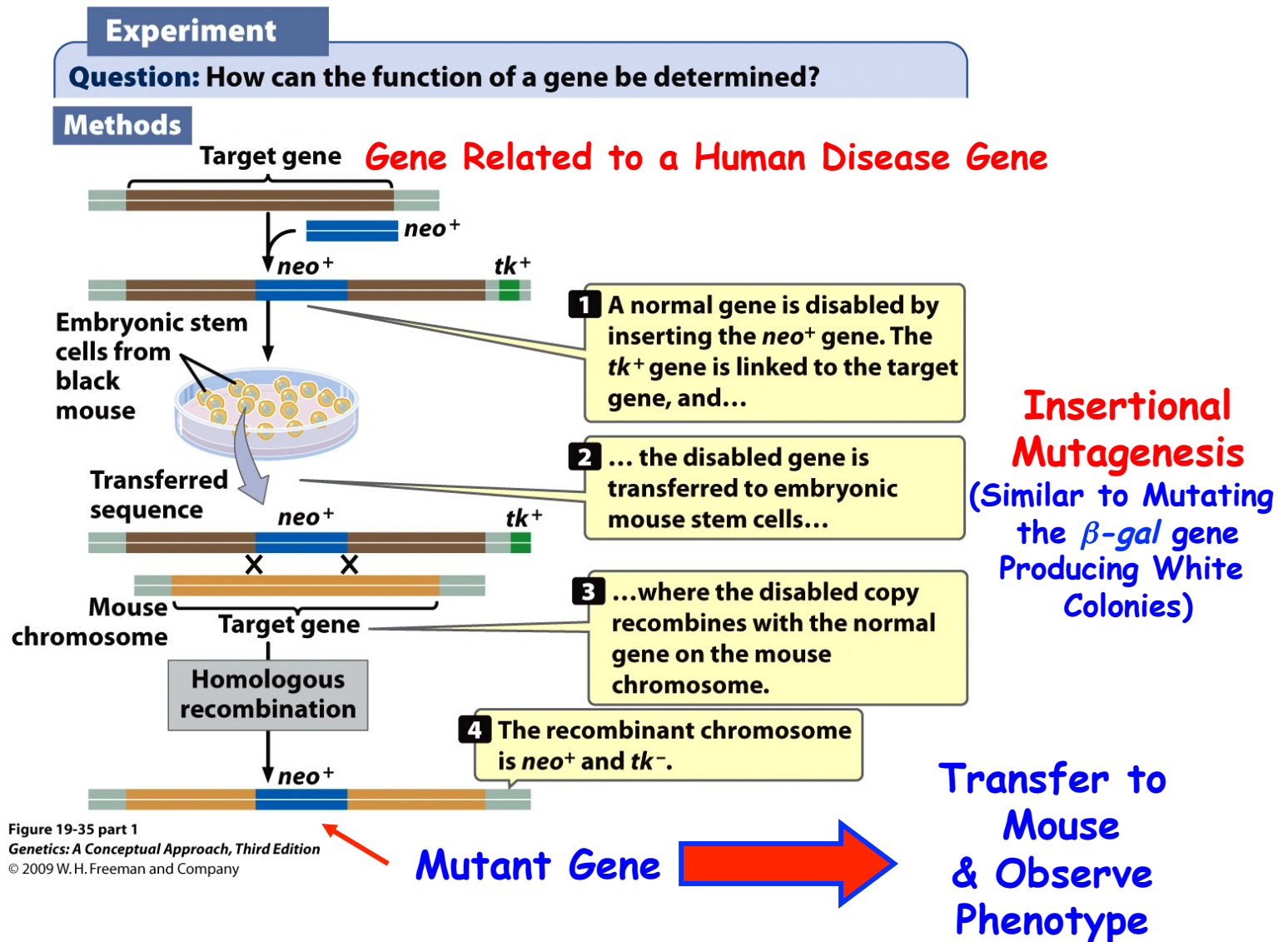
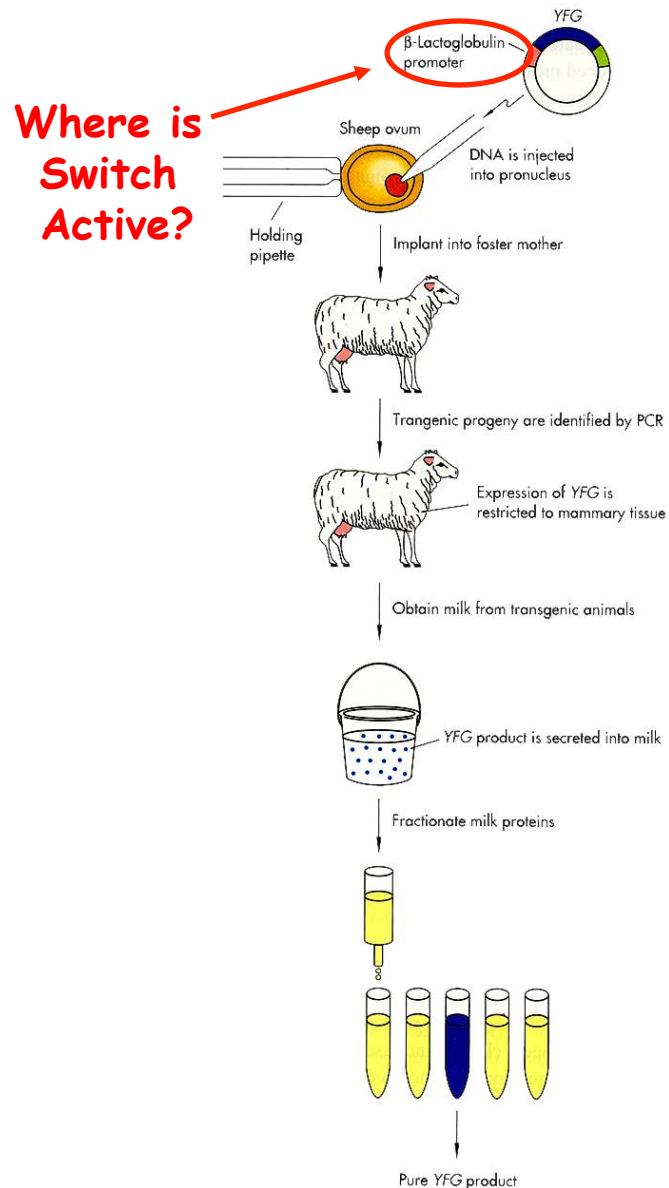


Figure 19-35 part 1  
Genetics: A Conceptual Approach, Third Edition  
© 2009 W. H. Freeman and Company

# Animals Can Also be Used as Factories to Produce Large Amounts of Human Proteins

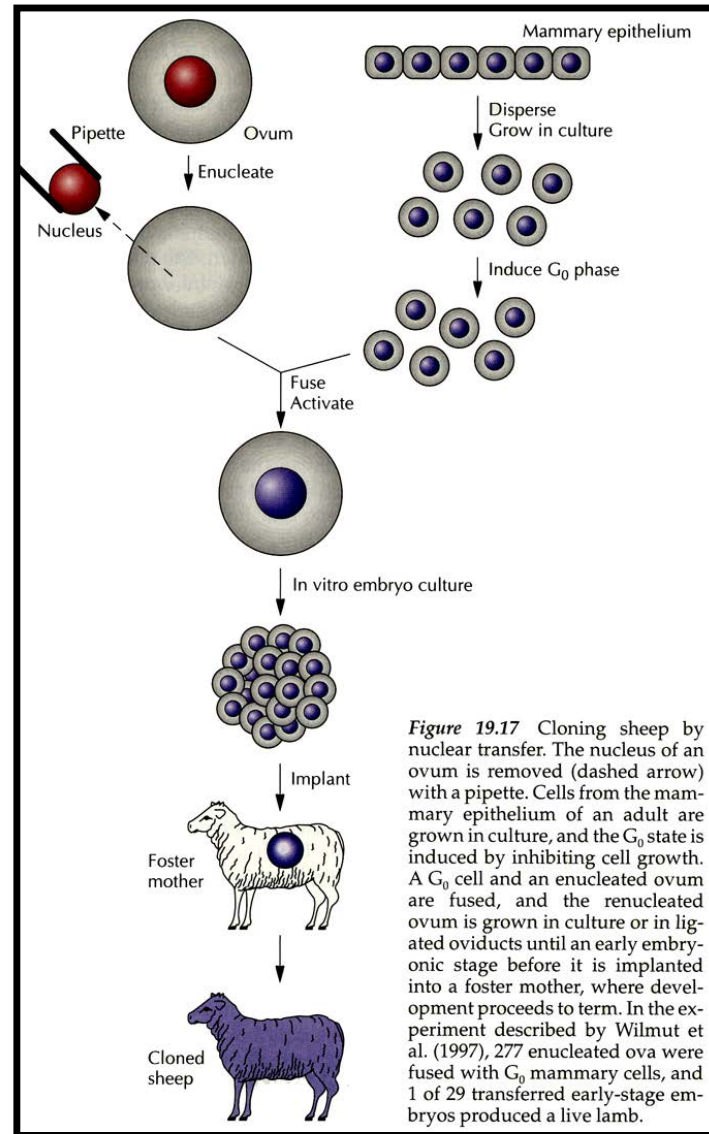


## Advantages of Molecular Pharming

1. Many human proteins need to be modified after translation to be active. Only eukaryotic cells can do this.
2. Bacteria need big fermentors + elaborate protein purification schemes--Farm animals can be used for this purpose w/o special processing/machinery.
3. Proteins stable, can be made in large amounts, and purified easily



# Genetically Engineered Drug-Producing Mammals Can Also Be Cloned



**Somatic Cells  
Can Also Be  
Genetically  
Engineered and  
Then Inserted  
Into Egg**

February 7, 2009

## F.D.A. Approves Drug From Gene-Altered Goats

Examined Data From Seven Generations of Genetically Engineered Goats

### New Drug From Genetically Engineered Goat

FDA OKs ATryn, 1st Drug Made in Milk of a Genetically Engineered Animal

By [Miranda Hitti](#)  
WebMD Health News

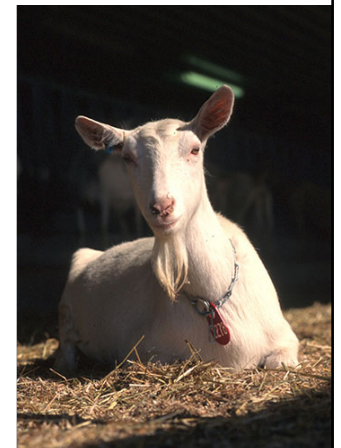
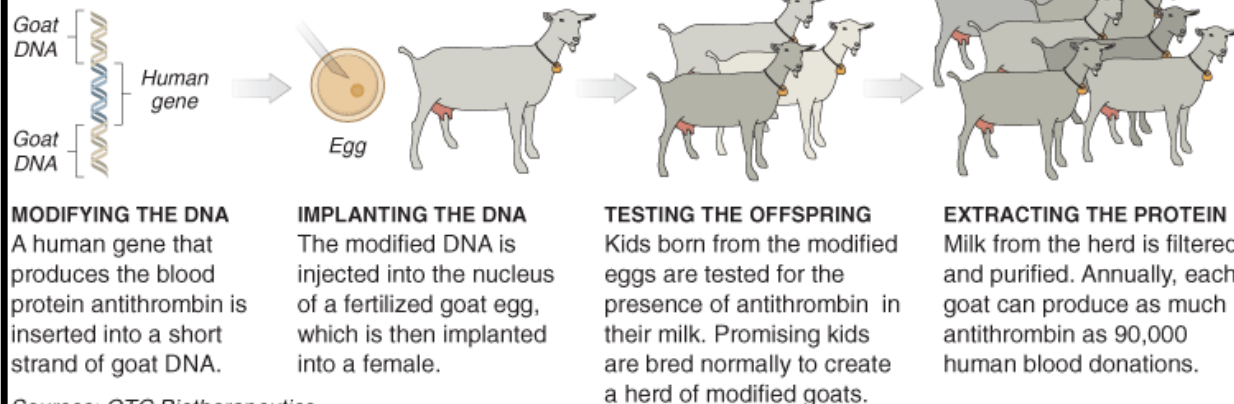
Feb. 6, 2009 -- The FDA today approved ATryn, the first drug made in genetically engineered animals.

#### Issues

Food Supply?  
Containment?  
Animal Health?  
Effective Drug?

#### Bioengineering on the Farm

The Food and Drug Administration has approved the first drug produced in the milk of genetically engineered animals.



# Making Recombinant Human Proteins in Animals

**Table 19.3** Some exogenous proteins that have been expressed in the mammary glands of transgenic animals

Antithrombin III  
Calcitonin  
Erythropoietin  
Factor IX  
Factor VIII  
Fibrinogen  
Glucagon-like peptide  
Granulocyte colony-stimulating factor  
Growth hormone  
Hemoglobin  
Human serum albumin  
Insulin  
Insulin-like growth factor 1  
Interleukin 2  
Lactoferrin  
Lysozyme  
Monoclonal antibodies  
Nerve growth factor  $\beta$   
Protein C  
Superoxide dismutase  
Tissue plasminogen activator  
 $\alpha$ 1-Antitrypsin  
 $\alpha$ -Glucosidase  
 $\alpha$ -Lactalbumin

## Advantages over Bacteria?

**Table 19.2** Milk production and estimated recombinant protein yields from organisms used for the expression of transgenes in mammary glands

Organism	Annual milk yield (liters)	Estimated recombinant protein per female (kg/yr)
Rabbit	5	0.02
Pig	300	1.5
Sheep	500	2.5
Goat	900	4
Cow	10,000	60 !!!



# And Don't Forget Plants!

## First plant-made biologic approved



Carrot cell bioreactors

The US Food and Drug Administration in May approved Eleyso (taliglucerase alfa), an enzyme produced in genetically engineered carrot cells, for treating type 1 Gaucher's disease. This is the first plant-made drug approved

by the regulators, and for Israeli company Protalix BioTherapeutics of Carmiel, it is the first product made in their ProCellEx protein expression system to reach the market. The plant cell platform produces recombinant proteins with a glycan and amino acid structure similar to naturally produced human counterparts. Some 10,000 patients worldwide have Gaucher's, a rare genetic disorder in which individuals fail to produce the enzyme glucocerebrosidase.

## Drug-making plant blooms

*Approval of a 'biologic' manufactured in plant cells may pave the way for similar products.*

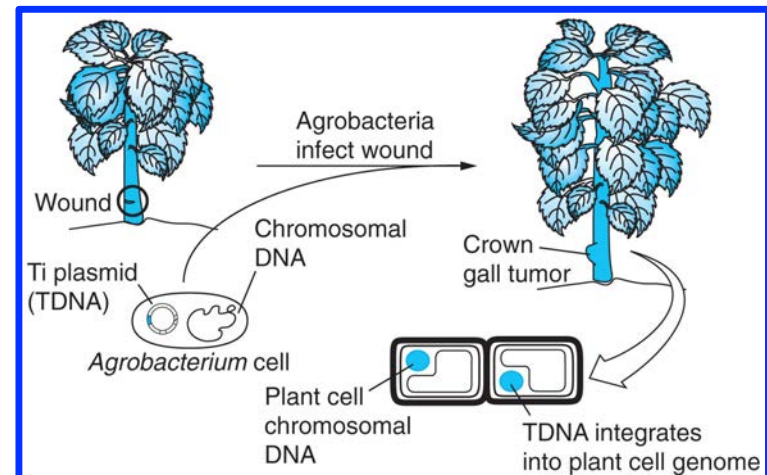
### PLANTS IN THE PIPELINE

Manufacturers have begun or completed phase II clinical trials on a handful of biologics made in plants, and hope to follow Eleyso to market.

Drug	Condition	Company	Platform
Locteron (interferon- $\alpha$ )	Hepatitis C	Biolex Therapeutics	Duckweed
H5N1 vaccine	Influenza	Medicago	Tobacco
VEN100	Antibiotic-associated diarrhoea	Ventria Bioscience	Rice
CaroRx	Dental caries	Planet Biotechnology	Tobacco

**Eleyso® Made in Engineered Carrot Cells To Treat Gaucher's Disease - A Lysosomal Storage Disease That Prevents Molecules From Being Degraded and Disposed of Properly in Cells - 100x Prevalence in Ashkenazi Jews. Gene on Chromosome 1, and Encodes a Glucocerebrosidase.**

**Advantages of Plants?**



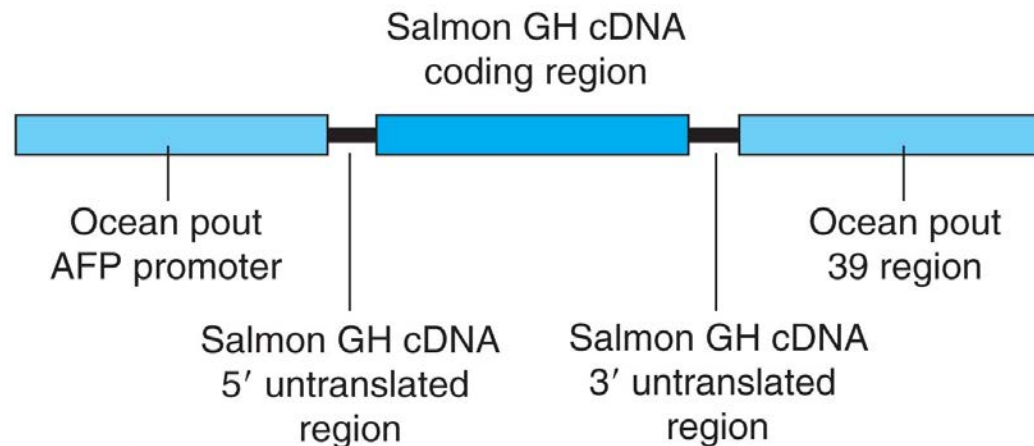
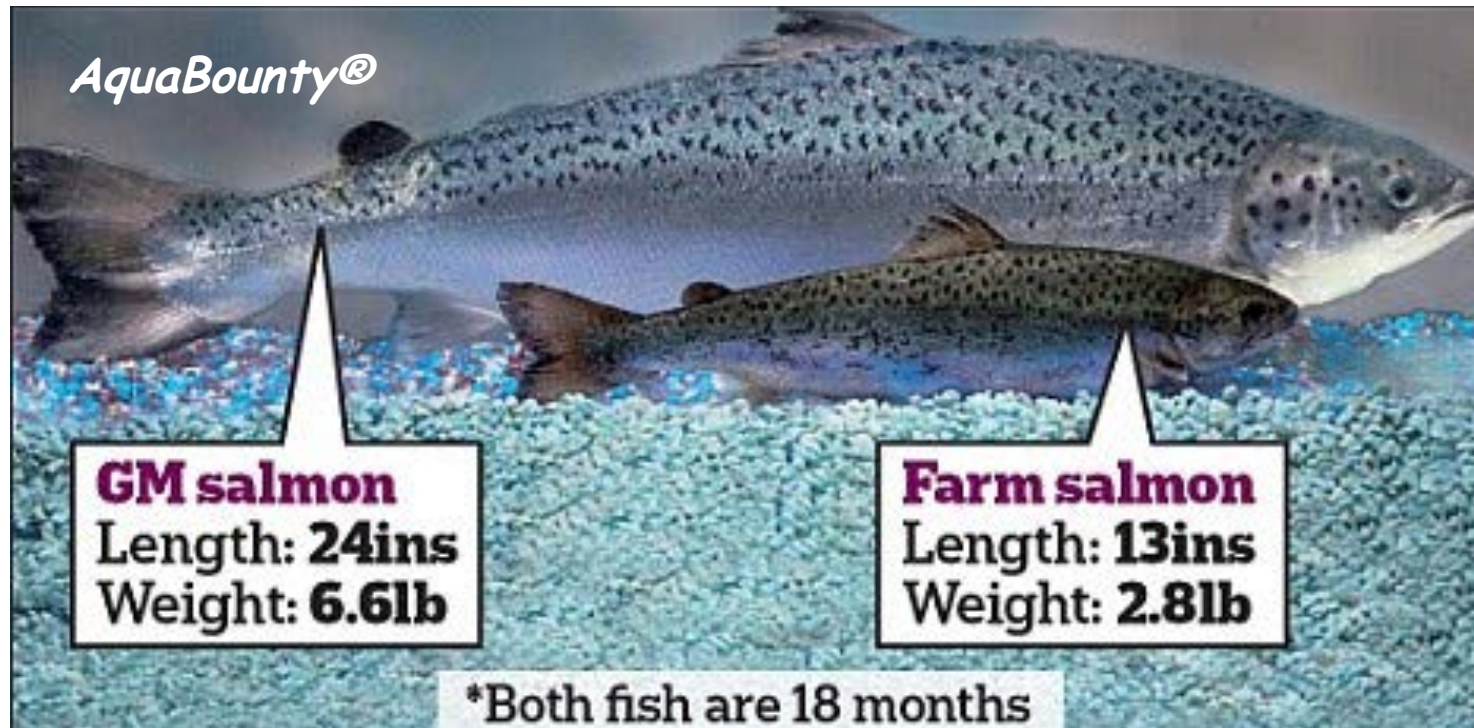
## Other Transgenic Animals Have Been Generated

TABLE 2.1 State of the art of transgenic technology for selected organisms.

Organism	Transfection	Viral vectors	Transposon	ES cells	Nuclear transfer
Mouse	4 <sup>a</sup>	2	1	4 <sup>a</sup>	2
Cow	3	1	0	0	2
Sheep	3	0	0	0	2
Goat	3	0	0	0	2
Pig	3	0	0	0	2
Rabbit	3	0	0	1	0
Chicken	1	2	1	0	0
Atlantic salmon	3	0	0	0	0
Channel catfish	2	0	0	0	0
<i>Tilapia</i>	3	0	0	0	0
<i>Zebrafish</i>	1	0	0	1	1
Crustaceans	1	1	0	0	0
Mollusks	1	1	0	0	0
<i>Drosophila</i>	2	2	2	2	0
Mosquito	1	0	2	0	0

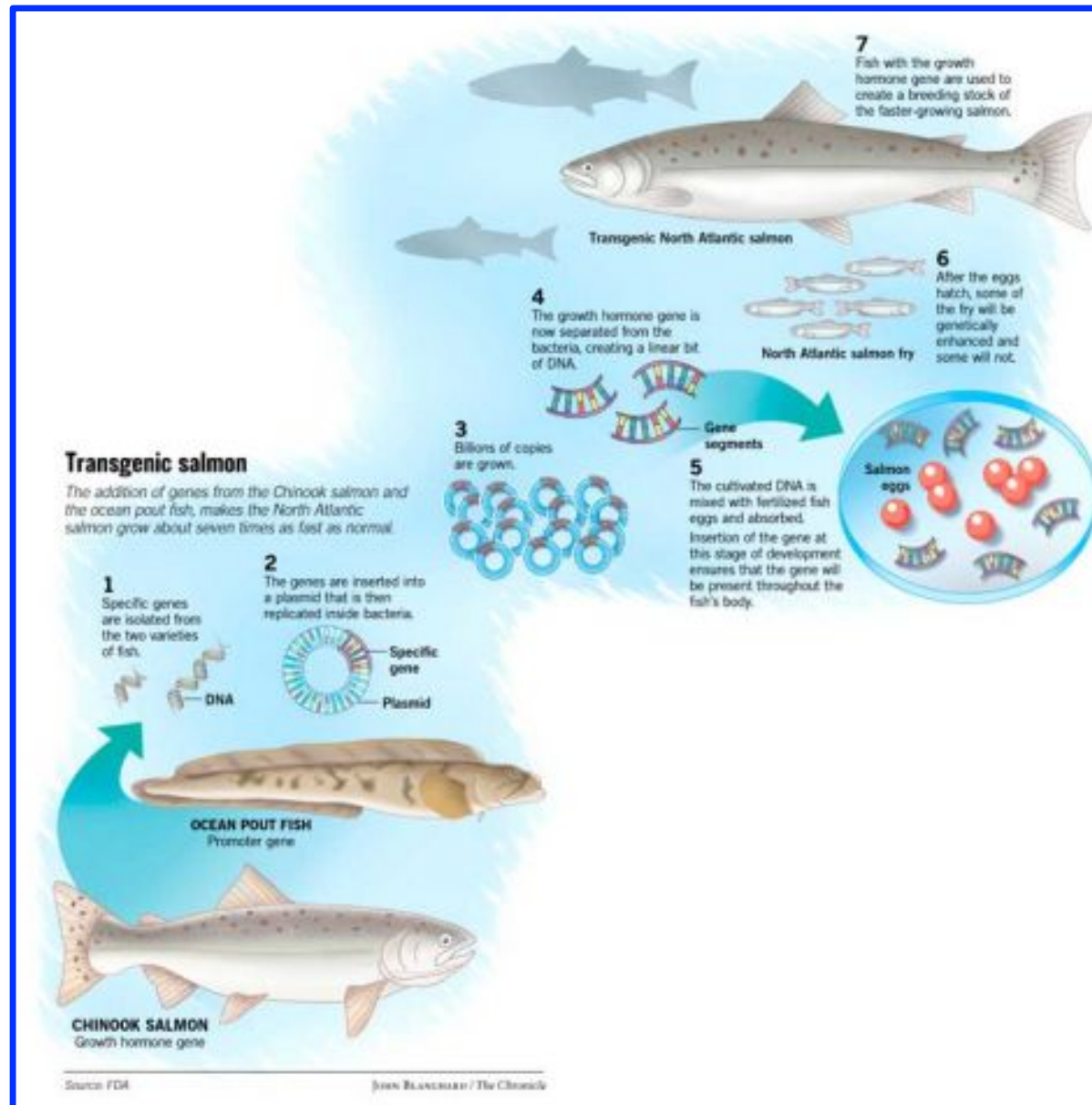
NOTE: 0: No significant progress.  
 1: Has been accomplished experimentally (proof of concept).  
 2: Routine experimental use.  
 3: Commercialization sought.  
 4: Widespread production.  
<sup>a</sup> For experimental uses.  
 See (Dove, 2000)

# Genetic Engineering Fast Growing Salmon



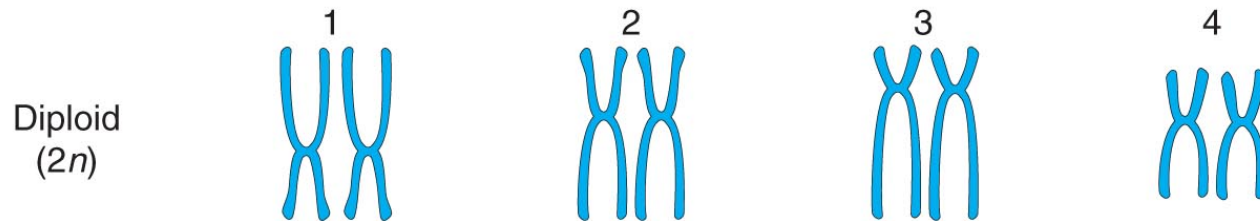


# Producing Genetically Engineered Salmon

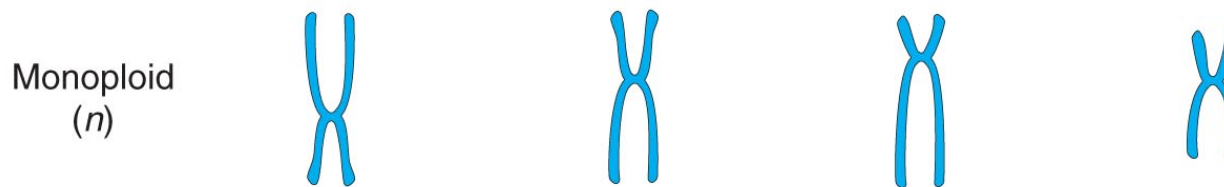


# Genetically Engineered Salmon Are Sterile

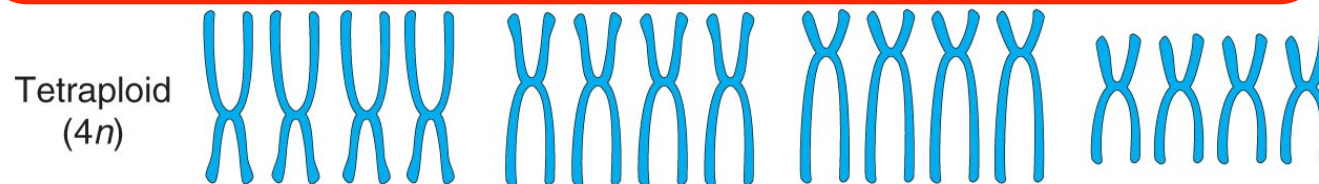
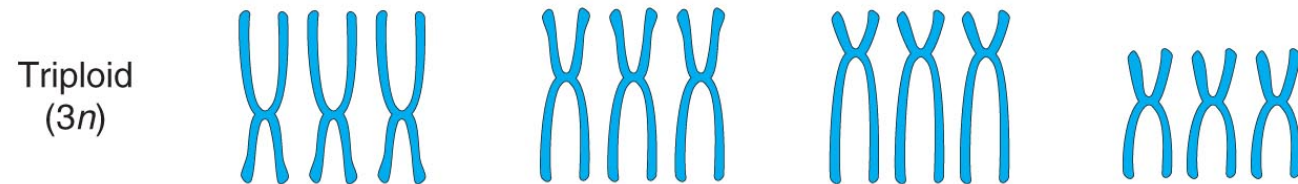
Normal chromosome complement in a somatic cell



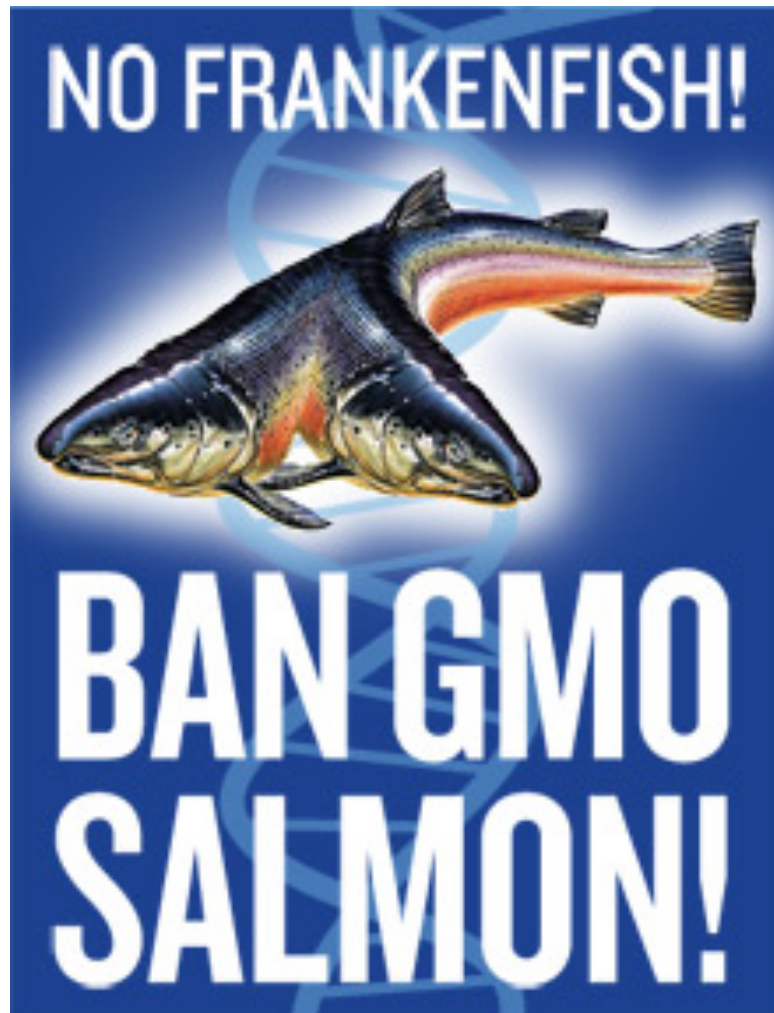
Haploid ( $n$ ) number of chromosomes in a gamete (egg or sperm)



Polyploid cells contain three or more sets of chromosomes



## Another GMO "Controversy"





**FOOD SCIENCE**

# Politics holds back animal engineers

*Funds and approvals lag for transgenic livestock in US.*

**OFF THE TABLE**

A brief history of some of the genetically engineered food animals submitted to the US Food and Drug Administration (FDA) for review. No such animal has yet been approved.

Animal	Purpose	Created	History
Salmon	Grows to market size faster than conventional salmon	1989 (Massachusetts)	1995 FDA receives application 2008 Fish farm moved to Panama 2010 Cleared by FDA scientific advisory panel
Pig	Produces more milk to nurse healthier young	1993 (Illinois)	1999 FDA receives application
Goat	Milk has human lysozymes to treat diarrhoeal disease	1999 (California)	2003 Funding denied by USDA 2008 FDA receives application 2011 Research moved to Brazil
Pig	Efficiently digests plant phosphorus, reducing pollution	1999 (Ontario, Canada)	2007 FDA receives application 2012 Pigs killed owing to lack of commercial interest
Cow, sheep, goat, pig	Increased muscle mass without reduced fertility	2010 (Texas)	2009 FDA receives application

## FDA expected to approve Genetically Modified Salmon

AQUAADVANTAGE SALMON | JANUARY 3, 2013 | BY: MARK WACHTLER |

## How To Assess Risk?

**TABLE 2-1 Systematic Risk Assessment and Management**

Step	Key questions
Hazard identification	What event posing harmful consequences could occur?
Risk analysis	<p>How likely is the hazard?</p> <p>What would be the harms from realization of the hazard, and how severe are they, taking into account social values?</p> <p>What is the risk assessment as shown on a matrix of risk (likelihood of harm) plotted against severity of harm; see Figure 2-1, above)? Each cell of the matrix should be accompanied by a qualitative assessment of the response and a quantification of assurance needed to reduce harm if the cell's conditions were to occur.</p> <p>How well established is the knowledge used to identify the hazard, estimate its risk, and predict harms?</p>
Risk reduction planning and implementation	What can be done (including bioconfinement and other confinement) to reduce risk, either by reducing the likelihood or mitigating the potential harms? Are there steps that can be taken to prepare for remediation?
Risk tracking (monitoring)	<p>How effective are the implemented measures for risk reduction?</p> <p>Are they as good as, better than, or worse than planned?</p> <p>What follow-up, corrective action, or intervention will be pursued if findings are unacceptable?</p> <p>Did the intervention adequately resolve the concern?</p>
Remedial action	What remedial action should be taken?
Transparency and public participation	How transparent should the entire process be? How much and what type of participation should there be in the steps above (and in risk characterization) by the public at large, by experts, and by interested and affected parties?

# What Are The Issues?

**MYTH 1:** *Transgenic salmon grow much larger than other salmon - so much so that they could gain a mating advantage or outcompete native salmon for food or space.*

**FACT:** Transgenic salmon grow **faster** than other salmon but they do not grow any larger by the time they reach maturity.

**FACT:** Male salmon do not gain a mating advantage because of size. In fact, "precocious parr," only 6 inches in length, father about one-fifth of each new generation before they go to sea. Studies of escaped farmed salmon, which are almost always larger than wild fish, have found them to mate successfully only **3 percent** as often as native salmon.

**FACT:** Farmed salmon are trained to eat fish feed -- small, dry pellets that look exactly like the "dog chow" we feed our family pets. If they escape, they look for something similar. Most don't find it. More than 85 percent of the farm escapees caught off British Columbia and Alaska had **no food** in their bellies. In a 1999 study, the Washington State Department of Ecology found farm escapees to be **eating tree bark** in local rivers, because it apparently looked like fish feed. Transgenic salmon may forage even more poorly because they lack the critical swimming speed to pursue prey, deplete their energy reserves more quickly and expose themselves to predators more often in the search for food.

**MYTH 2:** *If transgenic salmon do breed successfully with native fish, their novel gene will escape into the wild gene pool and destroy native salmon populations. Researchers at Purdue University found that only 60 transgenic salmon could drive a wild population to extinction.*

**FACT:** The U.S. Food and Drug Administration will not approve the use of transgenic salmon unless they can be demonstrated to be **sterile**. Aqua Bounty Farms has stipulated that it will market **only sterile, all female transgenic salmon**. There can be no gene flow to wild salmon because sterile fish cannot reproduce.

**And will be grown in cages far from natural salmon habitats!**

**FACT:** Muir and Howard, the Purdue scientists who proposed the "Trojan Gene Hypothesis," did not study transgenic salmon. They designed a mathematical model based on the behavior of Japanese medaka, a small, freshwater fish that matures in 56 days and breeds daily until it dies. Salmon take three, five and even ten years to mature and most breed only once in their lifetimes. Sterile salmon do not breed at all.

**MYTH 3:** *Sterilization is not 100 percent effective so we can't be sure that transgenic salmon will really be sterile.*

**FACT:** Triploidy produces complete, 100 percent sterilization in female salmon because it prevents the development of the ovaries needed to produce eggs. The only uncertainties about the technique have been raised in the context of male salmon, grass carp and oysters. There is no scientific debate over the complete sterility of triploid female salmon.

**FACT:** Scientists can test for triploidy by scanning blood or embryonic fluids in a flow cytometer. The sterility of every batch of transgenic salmon eggs can be verified before they ever leave the hatchery.

**MYTH 4:** *Transgenic salmon are voracious predators that will consume all the available food in an ecosystem and will prey on native juveniles.*

**FACT:** Transgenic salmon actually consume **less food** per pound of weight gained because they process their food 10 to 30 percent more efficiently.

**FACT:** Transgenic salmon may be highly prone to starvation in the natural environment as they learn to identify and hunt for wild food. They maintain a higher metabolic level for a longer period of time in food deprivation studies, and deplete their energy reserves more quickly than do standard salmon.

**FACT:** Any food competition would occur in the marine environment because sterile transgenic salmon cannot produce the juveniles that occupy freshwater habitat. In the marine life stages, transgenic salmon would compete with older native salmon of about the same size. Because food availability is not limiting in the marine environment, transgenic salmon would gain no advantage from their higher feeding motivation.

**FACT:** Sterile female salmon do not engage in spawning behaviors and almost never return to freshwater habitat after they begin to feed at sea. Native juveniles are confined to freshwater habitat. Any predation risk would, therefore, be lower than now occurs in conventional salmon aquaculture. There is no evidence of predation by current farm escapees on native juveniles.

**MYTH 5:** *Transgenic salmon produce antifreeze proteins and excessive amounts of growth hormone.*

**FACT:** Transgenic salmon produce no antifreeze proteins. Only the molecular "switch" from the antifreeze gene is used.

**FACT:** Transgenic salmon produce the same amount and kind of circulating growth hormone as wild-type salmon, but they produce it through the entire year.



# GM salmon: FDA's assessment of environmental risks



DEPARTMENT OF HEALTH AND  
HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2011-N-0899]

Draft Environmental Assessment and  
Preliminary Finding of No Significant  
Impact Concerning a Genetically  
Engineered Atlantic Salmon;  
Availability

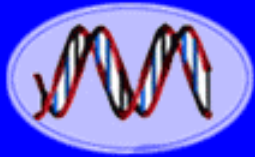
AGENCY: Food and Drug Administration,  
HHS.

ACTION: Notice.

SUMMARY: The Food and Drug  
Administration (FDA, the Agency) is  
announcing the availability for public  
comment of the Agency's draft  
environmental assessment (EA) of the  
proposed conditions of use specified in  
materials submitted by AquaBounty  
Technologies, Inc., in support of a new  
animal drug application (NADA)  
concerning a genetically engineered  
(GE) Atlantic salmon. Also available for  
comment is the Agency's preliminary  
finding of no significant impact (FONSI)  
for those specific conditions of use.

Both documents -- an **environmental assessment** and preliminary "**finding of no significant impact**," known by the policy wonks as a FONSI -- will be published Dec. 26 in the Federal Register and be available for public comment for 60 days. 12/26/12

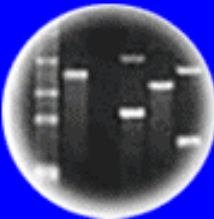
The assessment focused on the environmental questions. Food safety came earlier. Back in 2010, **the FDA concluded** that the salmon "is as safe as food from conventional salmon, and there is a reasonable certainty of no harm from consumption." For example, the flesh of the fish contain no more growth hormone than regular Atlantic salmon, the FDA said -- a concern of opponents to the fish because of the manner in which they were genetically engineered.



DNA  
Genetic Code of Life



Entire Genetic Code  
of a Bacteria



DNA Fingerprinting



Cloning: Ethical Issues  
and Future Consequences

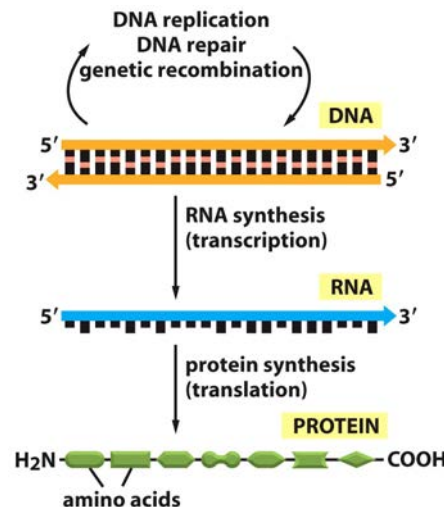


Plants of Tomorrow

# How Genes Work & What Are Genes In Context of...



## Thinking About The Consequences of GMOs



Need Science-  
Based Questions &  
Science-Based  
Solutions-NOT  
OPINIONS!

1. What is a Gene?
2. What is the Anatomy of a gene?
3. How Does the Gene Replicate?
4. How Does the Gene Direct Synthesis of a Protein?
5. Does the Gene Work Independently of other Genes?
6. What is the Sequence & Structure of the Protein?
7. How does it work in cell?
8. Does the Protein Structure imply any Potential "Harm"?
9. Does the Gene Change the Organism? Fitness?
10. Is the Environment Harmed?
11. Does the "Benefit" Outweigh the Cost?





*Professor Frank Furedi, University  
Of Kent, England*

