

of a Bacteria





Cloning: Ethical Issues and Future Consequences



Plants of Tomorrow

HC70A, SAS70A, & PLSS059 Winter 2019 Genetic Engineering in Medicine, Agriculture, and Law

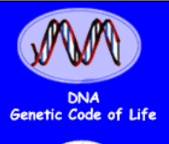
Professors Bob Goldberg, John Harada, & Channapatna Prakash

Lecture 5
The Nuts & Bolts of Genetic Engineering:
From Mutations to Pedigrees to Drug
The Factor XIII Story

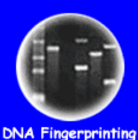














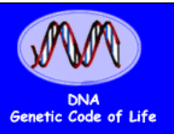
Cloning: Ethical Issues and Future Consequences

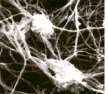


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#### **THEMES**

- 1. What is Hemophilia and How is it Inherited?
- 2. How Can a Disease Gene Be Found When It is Not Known Where the Gene is Expressed?
- 3. What Vectors Can Be Used For Cloning DNA?
- 4. What is the Advantage of Using a Virus Vector For Constructing Genome Libraries?
- 5. How To Make a Library of the Human Genome?
- 6. How Find a Gene With Only a Knowledge of the Protein Sequence?
- 7. How Use DNA Testing to Detect Factor VIII Disease Alleles?
- 8. How Isolate a Factor VIII cDNA Clone?
- 9. Genomic vs. cDNA Libraries
- 10. How Produce Factor VIII Protein For Use as a Drug?





## How Do We Treat a Genetic Disease? From Gene To Drug



Due to Mutations in a Different Class of Blood Proteins

## The Molecular Genetics of Hemophilia

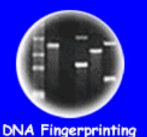
(Potentially Lethal Disease)

Hemophiliacs bleed because a defective gene deprives them of a key blood-clotting protein. The protein has now been made artificially by isolating the normal gene and then inserting it into cultured cells

by Richard M. Lawn and Gordon A. Vehar



of a Bacteria





Cloning: Ethical Issues and Future Consequences



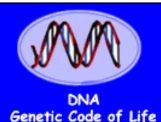
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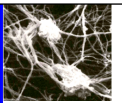
#### A Case Study of Cloning Genes and mRNAs

Reference: Scientific American, March 1, 1986





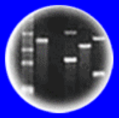








Entire Genetic Code of a Bacteria



**DNA Fingerprinting** 



Cloning: Ethical Issues and Future Consequences



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Founded in 1976 By Robert Swanson and Herb Boyer First IPO in 1980 for \$88/share Purchased by Hoffmann-La Roche in 2009 for \$47B





## Hemophilia Has Been Known As An Inherited Disease For >2500 Years!



#### First Reference to Hemophilia is in the Old Testament

#### Genesis 17:10-14

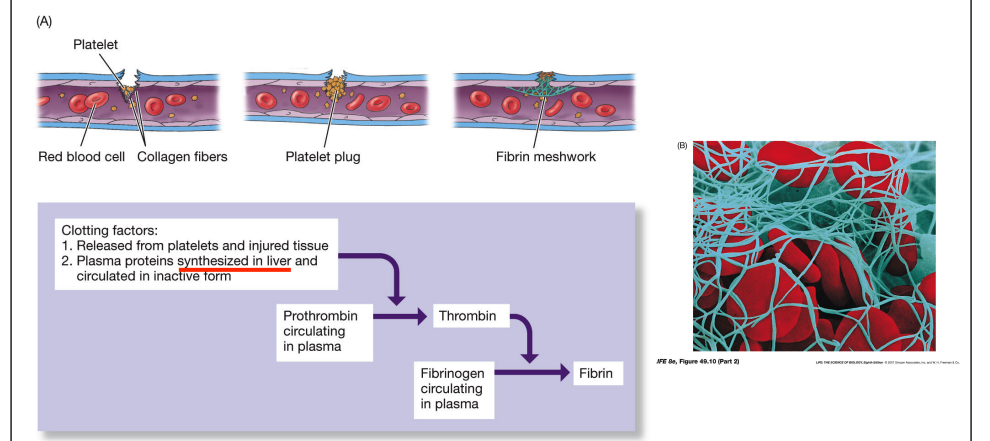
'This is My covenant that you shall keep between Me and you and your descendants after you: every male among you shall be circumcised. You shall circumcise the flesh of the foreskin.......At the age of eight days every male among you shall be circumcised throughout your generations............an uncircumcised male...that soul shall be cut off from its people, he has invalidated My covenant.'

The Talmud also makes reference to families in whom children have died as a result of circumcision (Babylonian Talmud, Chapter Yevamoth p64b) [6]. Should a mother lose two children or should two sisters lose a child each after circumcision, subsequent children of the woman, the two sisters or of any other sisters of the same family should not be circumcised until they are older, or possibly not at all. This is thought to be the earliest reference to haemophilia; it was recognized in the Talmud that this condition was transmitted by the mother.



Abraham was circumcised at 93 and gave birth to Isaac at 99. His wife - Sarah - was 90!

#### A Cascade Of Events After Wounding Leads to A Fibrin Clot

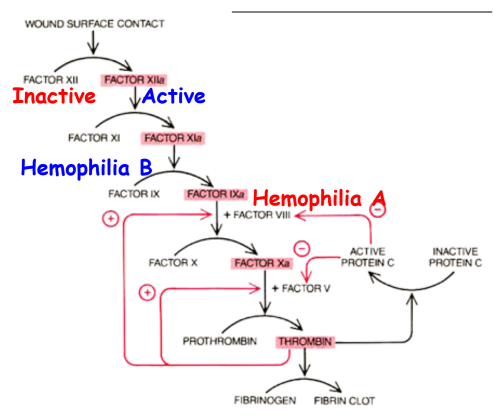


LIFE 8e, Figure 49.10 (Part 1)

LIFE: THE SCIENCE OF BIOLOGY, Eighth Edition @ 2007 Sinauer Associates, Inc. and W. H. Freeman & Co.

Clotting Factors Such As Factor VIII Play A Critical Role in This Process

#### Protein Factors in Blood Lead To Clotting



CLOTTING CASCADE begins when cell damage at a wound somehow activates the enzyme factor XII; it ends with the conversion of fibrinogen into fibrin by thrombin. At each step an inactive protein is converted into a protease, or protein-cutting enzyme (color), which activates the next protein. Some steps require cofactors such as factors VIII and V. The cascade includes positive- and negative-feedback loops (colored arrows). Thrombin activates factors VIII and V; it also deactivates them (by activating protein C), which helps to halt clotting. Some 85 percent of hemophiliaes lack factor VIII. The rest lack factor IX.

Eight
Proteins/Genes
Required:

- 1. Factor VII
- 2. Factor XI
- 3. Factor IX
- 4. Factor VIII
- 5. Factor X
- 6. Protein C
- 7. Prothrombin
- 8. Fibrinogen

What Happens If Any of These Proteins, or Genes, are Mutated?



No Blood Clot!

#### Hemophiliacs Have Mutations in Factor VIII, Factor IX, or Factor XI Genes

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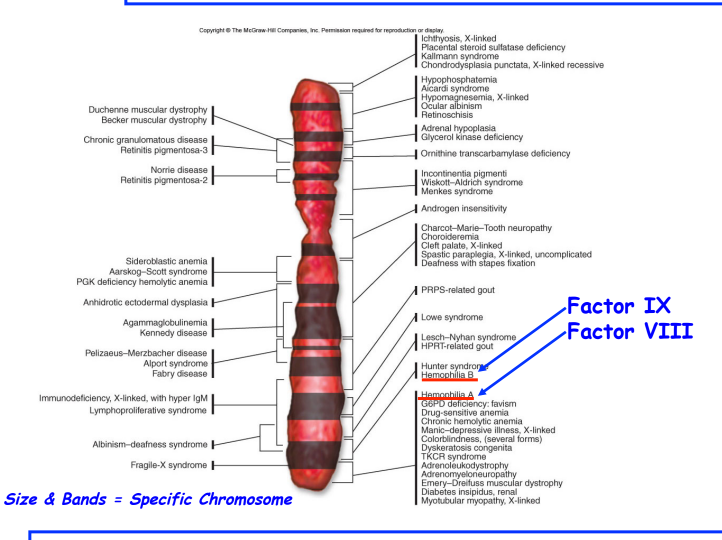
TABLE 13.2	Some Important	Genetic Disorders		
Disorder	Symptom	Defect	Dominant/ Recessive	Frequency Among Human Births
Cystic fibrosis	Mucus clogs lungs, liver, and pancreas	Failure of chloride ion transport mechanism	Recessive	1/2500 (Caucasians)
Sickle cell anemia	Blood circulation is poor	Abnormal hemoglobin molecules	Recessive	1/600 (African Americans)
Tay-Sachs disease	Central nervous system deteriorates in infancy	Defective enzyme (hexosaminidase A)	Recessive	1/3500 (Ashkenazi Jews)
Phenylketonuria	Brain fails to develop in infancy	Defective enzyme (phenylalanine hydroxylase)	Recessive	1/12,000
Hemophilia	Blood fails to clot	Defective blood-clotting factor VIII	X-linked recessive	1/10,000 (Caucasian males)
Huntington disease	Brain tissue gradually deteriorates in middle age	Production of an inhibitor of brain cell metabolism	Dominant	1/24,000
Muscular dystrophy (Duchenne)	Muscles waste away	Degradation of myelin coating of nerves stimulating muscles	X-linked recessive	1/3700 (males)
Hypercholesterolemia	Excessive cholesterol levels in blood lead to heart disease	Abnormal form of cholesterol cell surface receptor	Dominant	1/500

18,000 People in US Have Hemophilia & 400 Babies/Year Are Born With Disorder Prior to 1960s - Average Life Span Was 11 Years

Hemophilia A	Defective Factor VIII Gene	1/10,000 males	80%
Hemophilia B	Defective Factor IX Gene	1/30,000 males	20%
Hemophilia C	Defective Factor XI Gene	Autosomal	<1%

Both Factor VIII & IX Genes on X-Chromosome  $(9 \rightarrow 3)$  s)

## Factor VIII and Factor IX Genes are Closely Linked on the X Chromosome





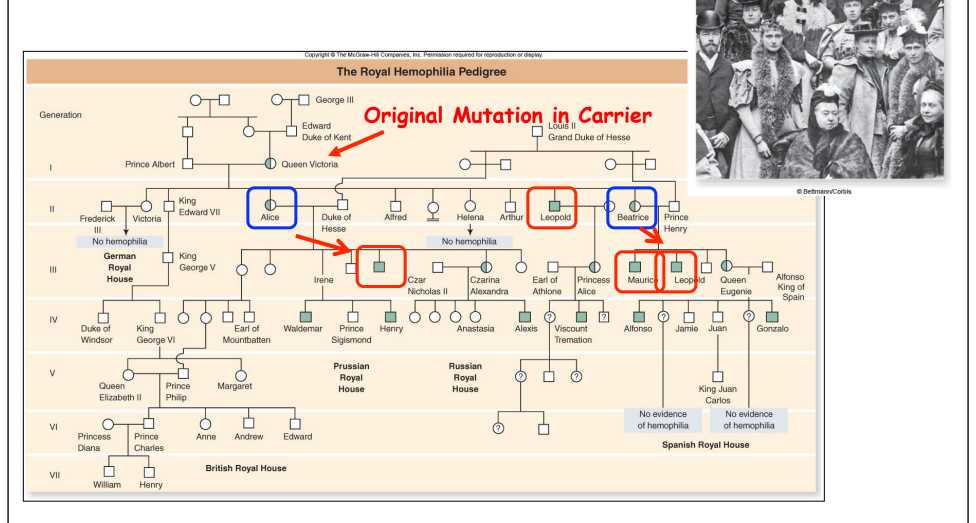
The X chromosome has ~1098 Genes and 150,000,000 bp (150 Mb). 168 Mendelian Diseases Explained by 113 X-Linked Genes



Pedigrees Can Be Used To Determine If a Trait is Dominant or Recessive

Each Type of Inheritance Predicts Specific Results in Each Generation

## Hemophilia A and B Genes Are Sex Linked & Recessive Traits When Mutated



- Note: 1. Males Obtain Detective Gene From Mothers
  - 2. 50% of Sons Of A Maternal Carrier Have The Defective Gene

#### Hemophilia A and B Sex-Linked Inheritance

#### Carrier Female

Healthy Male

Egg	X	X
Sperm		
X	XX	XX
	♀ <i>Carrier</i>	♀ Healthy
Y	XY	XY

#### Sex-Linked Inheritance

♀ Carriers → 1/2 Sons Afflicted + No Daughters! Only One X-Chromosome is in 3

## What Was Known About Factor VIII Before Gene Cloned?

- 1. Blood Protein (But Perhaps Synthesized Elsewhere!)
- 2. Could be purified in small amounts from >20 Liters of human blood + cow blood + pig blood
- 3. Short Stretch of <u>Protein</u> Sequenced = Known Protein Sequence!
- \*4. Hemophilia A could be treated by <u>blood transfusions</u> from normal individuals, ∴ clotting factor <u>in blood</u>
- 5. 1980s AIDS Epidemic Caused Many Hemophiliacs to Get HIV/AIDS (~50% of hemophiliacs got AIDS in 1985)
  - .. How to go From Protein to Gene

#### The Problem

## For Factor VIII- Not Known Where Gene Was Expressed: Must Use Genome Library

Analysis of presence or absence of

Wild-type

person Hemophiliac

blood-clotting factors

VIII

Early 1980's

(b) Vessel damage

Inactive XII - Active XII

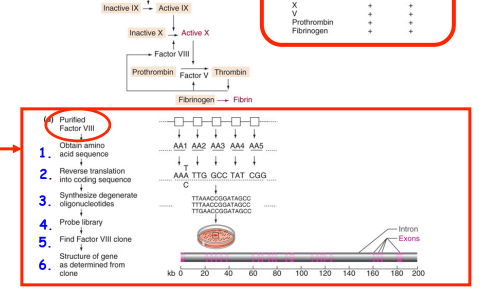
Inactive XI - Active XI

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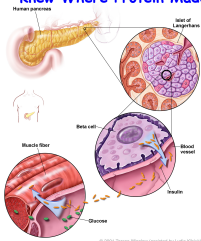
Key Concept

How Clone A Gene When You Don't Know Where it is Expressed!

Key:
Protein
Sequence
Known



Different Than Insulin Knew Where Protein Made!



mRNA -> Drug

How Find Gene & cDNA?

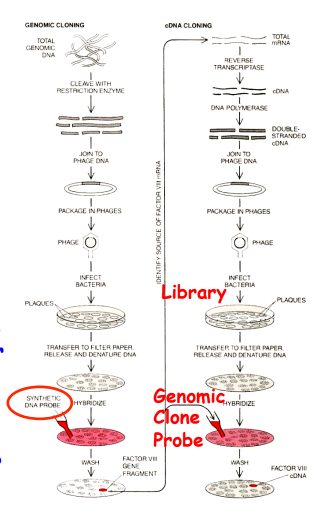
Protein → Gene → mRNA → Drug!

Steps Required to Clone Factor VIII

Gene and cDNA

Gene

- 1. Make Genome Library Because Factor VIII Gene in Genome!
- 2.Purify Protein from Blood- that's where it works (wasn't known where made)
- 3. Reverse Translate using the genetic code a portion of the protein sequence
- 4. Synthesize a DNA probe complementary to Factor VIII gene corresponding to protein sequence
- 5. Screen Genome Library Entire Gene on The Clone?



cDNA

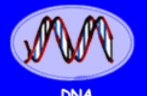
- 1.Use Gene probe to screen cDNA library for Factor VIII cDNA clone
- 2. How know what mRNA to use to make cDNA library?
- 3.Use gene probe to probe RNA blots containing mRNA from all major organs (liver, kidney, blood, etc.)
- 4. Find Factor VIII
  mRNA in livermale, liver- secrete
  into blood
  Why Need cDNA?
  Story continued

Want cDNA to Manufacture Factor VIII as a Drug to Treat Hemophilia A!

#### Step One

How to Construct a Human Genome Library to Find the Factor VIII Gene?

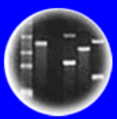
If It is Not Known Where Gene is Active Can "Look" to Genome Instead of mRNA to Find + Clone Gene!



Genetic Code of Life



Entire Genetic Code of a Bacteria



**DNA Fingerprinting** 



Cloning: Ethical Issues and Future Consequences



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#### Vectors Used in Genetic Engineering Have Similar Conceptual Properties But are Used in Different Situations

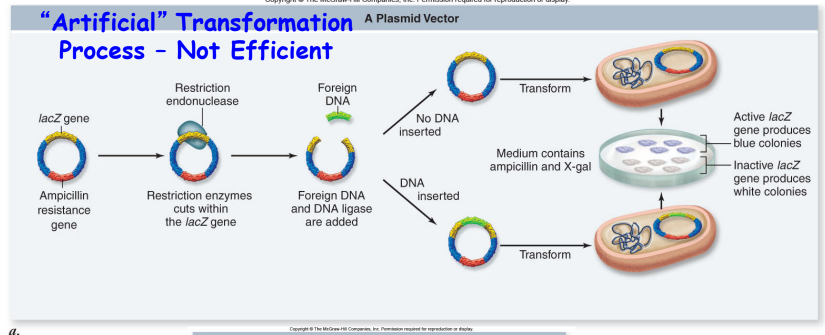
Vector Type	Maximum Insert Size (kb)	Applications	Limitations
Bacterial plasmid vectors (circular)	≃6-12	DNA cloning, protein expression, subcloning, direct sequencing of insert	Restricted insert size; limited expression of proteins; copy number problems; replication restricted to bacteria
	DNA	DNA	
Bacteriophage vectors (linear)	~25	cDNA, genomic and expression libraries	Packaging limits DNA insert size; host replication problem
Cosmid (circular)	~35	cDNA and genomic libraries, cloning large DNA fragments	Phage packaging restrictions not ideal for protein expres- sion; cannot be replicated in mammalian cells
Bacterial artificial chromosome (BAC, circular)	~300	Genomic libraries, cloning large DNA fragments	Replication restricted to bacteria; cannot be used for protein expression
Yeast artificial chromosome (YAC, circular)	200–2,000	Genomic libraries, cloning large DNA fragments	Must be grown in yeast; canr be used in bacteria
Ti vector (circular)	Varies depending on type of Ti vector used	Gene transfer in plants	Limited to use in plant cells only; number of restriction sites randomly distributed; large size of vector not easily manipulated

#### Properties of All Vectors

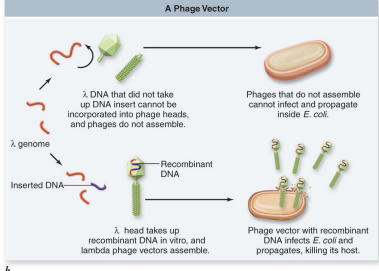
- 1. Replicate
- 2. Selectable
- 3. Can Be Used To Insert Foreign Genes/Restriction Sites
- 4. Easily Isolated + Transferred Back To Cells

## Plasmid vs. Bacteriophage Vectors for Cloning DNA Fragments

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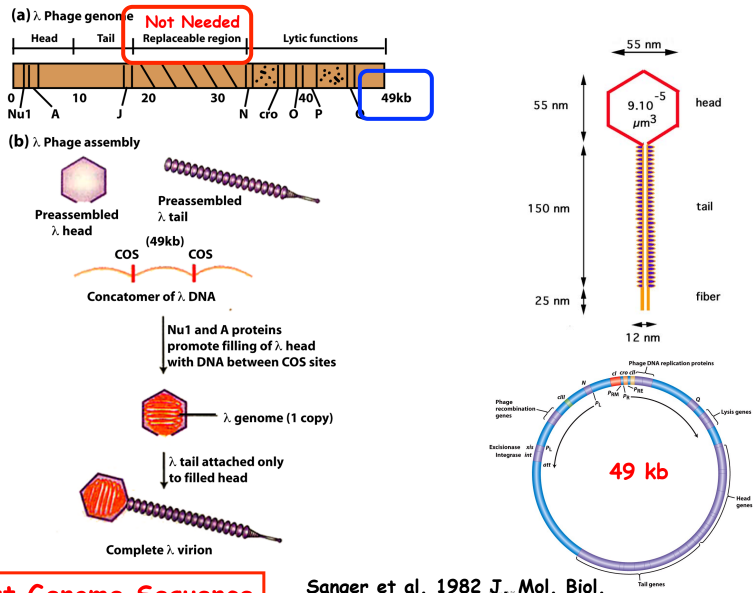


"Natural"
Infection Process



- Much More Efficient
- Can Use Less DNA
- Get Lots More Clones
- Need Lots of Clones For Large Genome

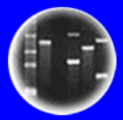
#### Structure of the $\lambda$ Phage and Its Genome



First Genome Sequence

Sanger et al. 1982 J. Mol. Biol. 162: 729-773.

# DNA Genetic Code of Life Entire Genetic Code of a Bacteria



**DNA Fingerprinting** 

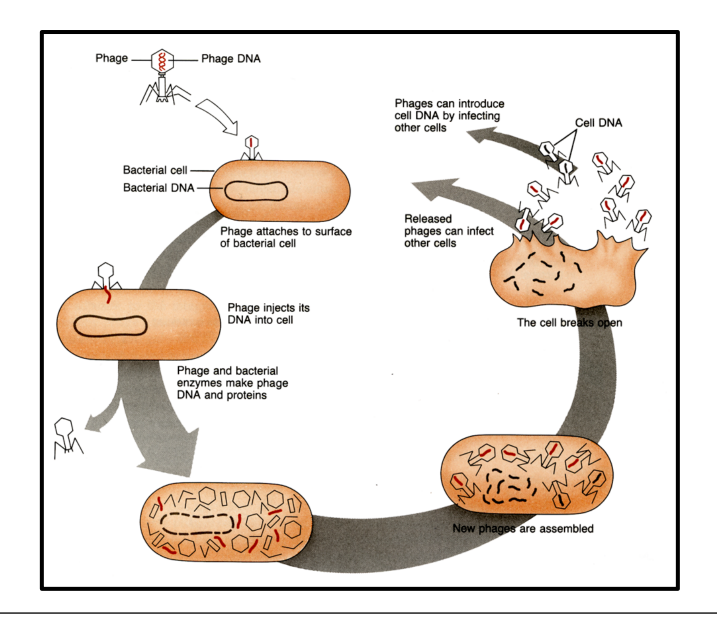


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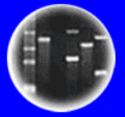
#### λ Phage Infects E.coli & Destroys (Lyses) Cells



## DNA Genetic Code of Life



Entire Genetic Code of a Bacteria



**DNA Fingerprinting** 

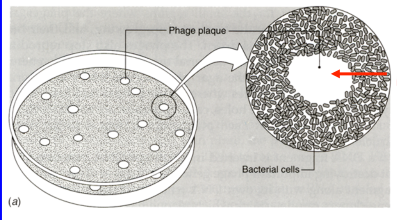


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#### Lysed Cells Can Be Seen as Clear Plaques on Agar Plates



Plaque

Plaque

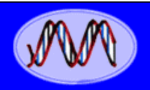
1. Each <u>Plaque</u> is a Virus Clone Representing One Viral Infection!

2. Selectable <u>Marker</u> is Bacterial Cell Destruction & Plaque Formation

## Advantages of $\lambda$ Virus as a Vector for Cloning DNA

- 1. Long DNA Segments Can Be Cloned (~20kb) Need Fewer Clones For Whole Genome!
- 2. Can Clone DNA Segments In Viral Genome & Self-Assemble With Viral Proteins Into Virus In A Test Tube!
  - .. Make Recombinant Viruses In The Lab!
- 3. <u>Use "Natural" Infection</u> Process To Generate Large Number Of Clones For A Eukaryotic Genome Library.
  - Much Higher Efficiency For Getting Recombinant DNA

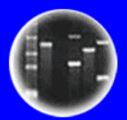
    →Bacterial Cells Compared With Dna Transformation.
  - .. Set More Clones Per Amount Of Recombinant DNA!



DNA Genetic Code of Life



Entire Genetic Code of a Bacteria



**DNA** Fingerprinting

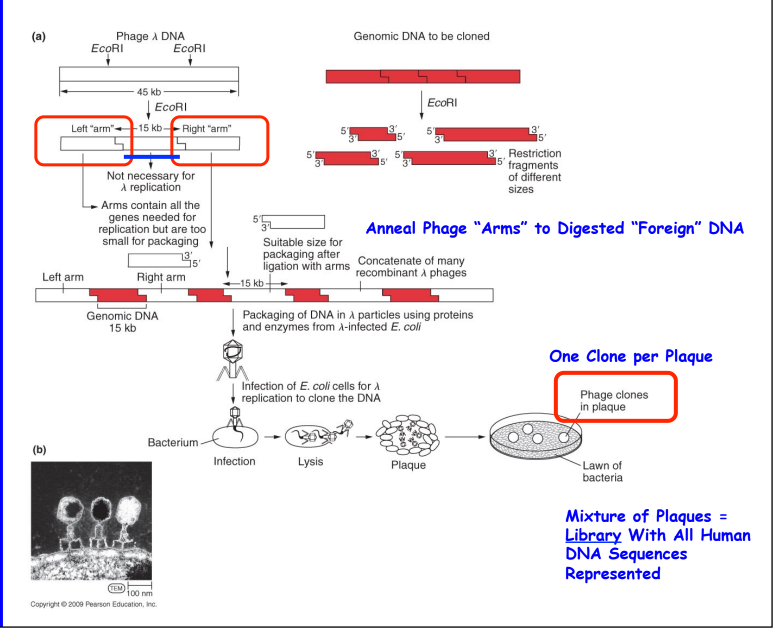


Cloning: Ethical Issues and Future Consequences

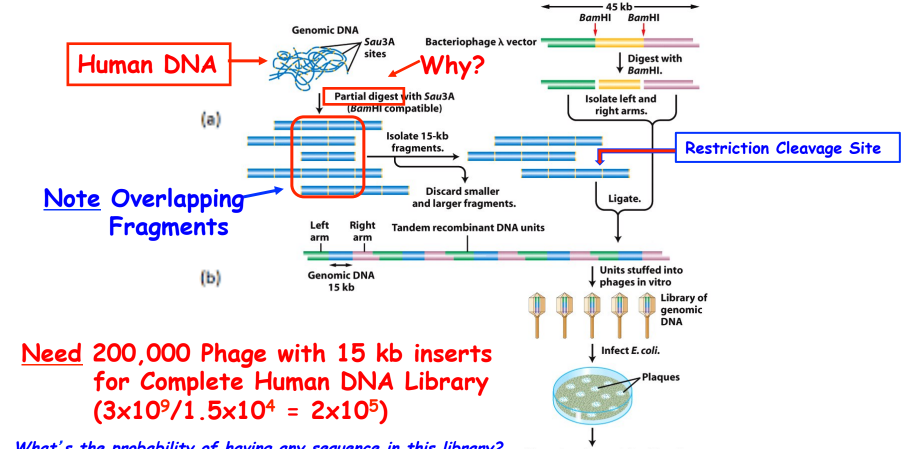


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## Using a Bacterial Virus To Clone the Human Genome



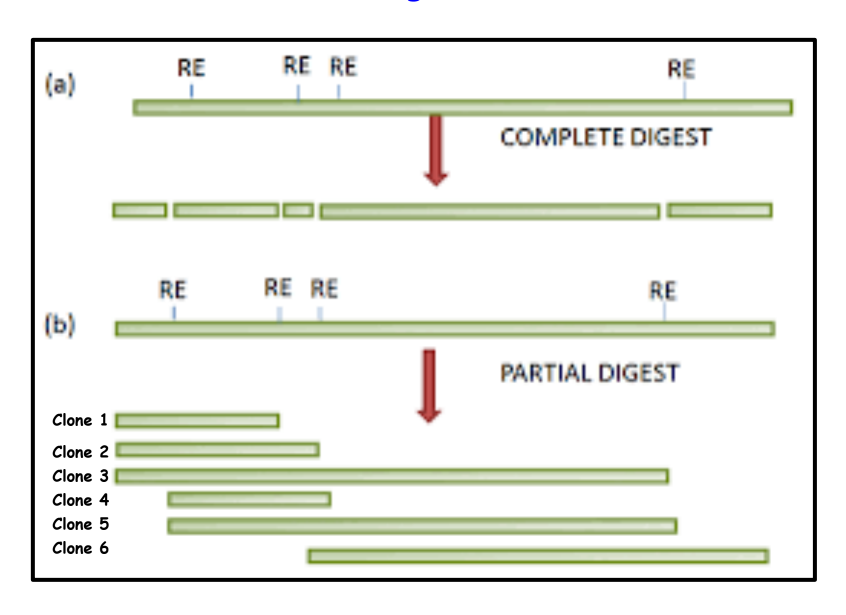
#### Cloning the Human Genome and Screening for the Factor VIII Gene



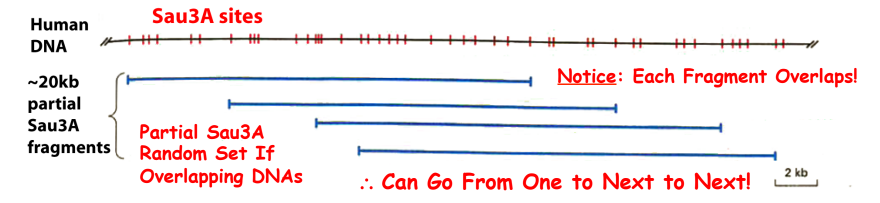
Introduction to Genetic Analysis, Ninth Edition © 2008 W. H. Freeman and Company

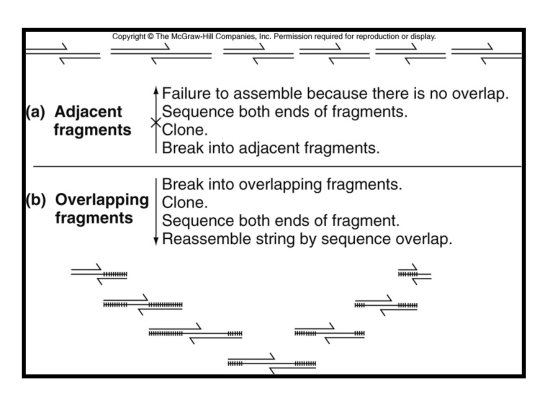
Why Partial Digestion? An Important Concept! What is Complete & Partial Digestion?

## Partial Digestion Permits "Walking" From One DNA Region to the Next



### Constructing a Human Genome Library by Partial Digestion Creates a Set of Overlapping DNA Fragments/ Clones





#### Step Two

## How Find the Factor VIII Gene in a Human Genome Library?

#### The Genetic Code

Second Letter

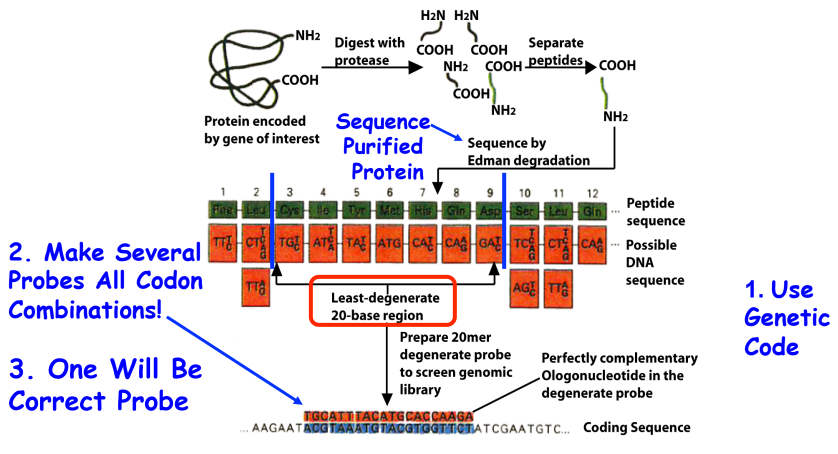
		u	J		3		A	9	}		_
	5	UUU UUC UUA UUG	Phe Leu	UCU UCC UCA UCG	Ser	UAU UAC UAA UAG	Tyr Stop Stop	UGU UGC UGA UGG	Cys Stop Trp	UCAG	
1st	С	CUU CUC CUA CUG	Leu	CCU CCC CCA CCG	Pro	CAU CAC CAA CAG	His Gln	CGU CGC CGA CGG	Arg	UCAG	3rd
letter	A	AUU AUC AUA AUG	lle Start Met	ACU ACC ACA ACG	Thr	AAU AAC AAA AAG	Asn Lys	AGU AGC AGA AGG	Ser Arg	UCAG	letter
	G	GUU GUC GUA GUG	Val	GCU GCC GCA GCG	Ala	GAU GAC GAA GAG	Asp Glu	GGU GGC GGA GGG	Gly	U C A G	

#### **Properties**

- Universal
- Three Nucleotides
  - Punctuation
  - Degenerate

#### Factor VIII Protein → Gene

Using the Factor VIII Protein Sequence and Genetic Code as a Guide to <u>Synthesize</u> a Factor VIII Probe



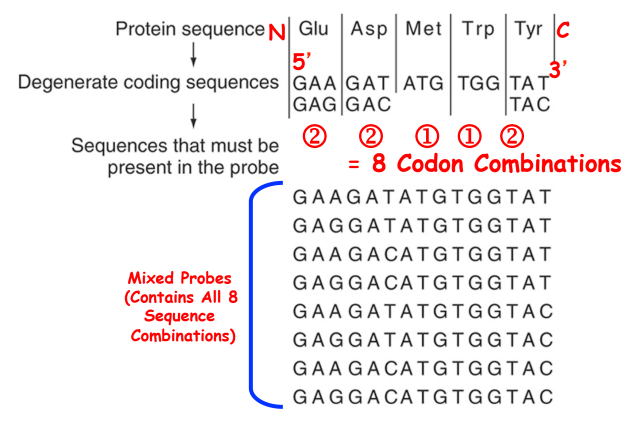
How many Combinations of **Synthetic** Probes?

2x3x2x1x2x2x2=96

## Using the Genetic Code to go From Protein Sequence to Gene Sequence

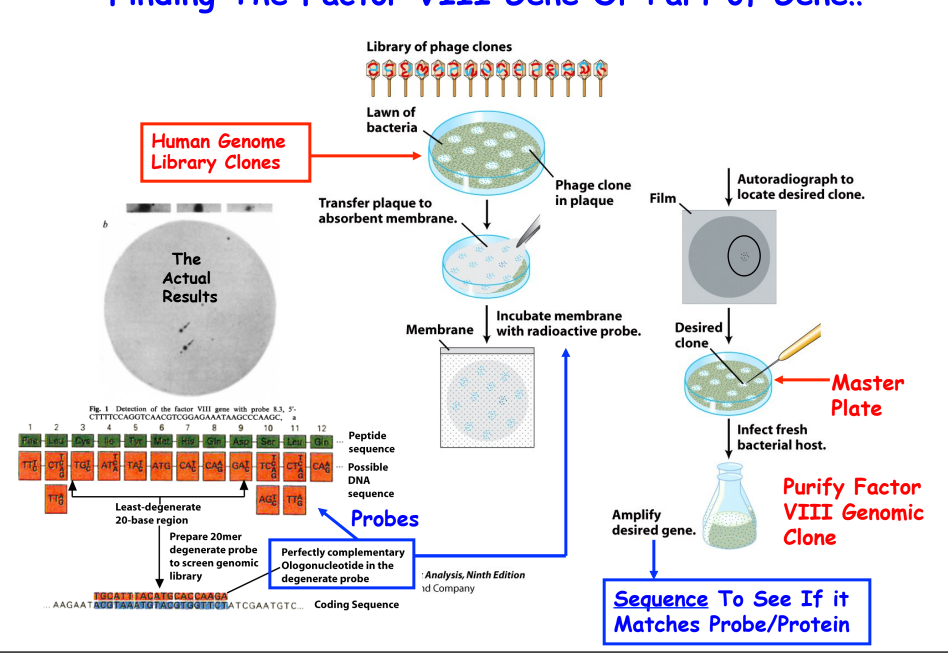
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(b) Synthesizing DNA probes based on reverse translation

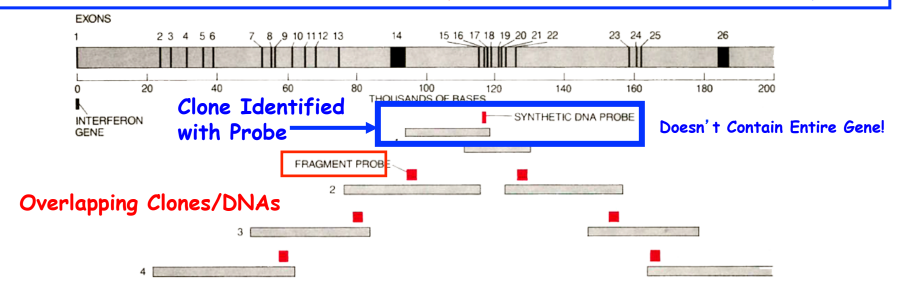


- 1. Need Amino Acid Sequence of Part of the Protein
- 2. Need DNA Sequences Representing all Codon Combinations
- 3. <u>Synthesize</u> DNA Sequence Probes!

#### Finding The Factor VIII Gene Or Part of Gene!!



## The Result-The Factor VIII Gene is Huge- 186,000 bp- The Probe Identified a Clone Containing Only One Part of Gene !!! Why?



How Find Clones with Rest of Gene?

Key Question!

Remember - the Library Contains Overlapping DNA Clones ∴ Can

Use One Part of First Clone to Re-Screen Library &

"Walk" to Other Gene Regions - Using Restriction Maps &

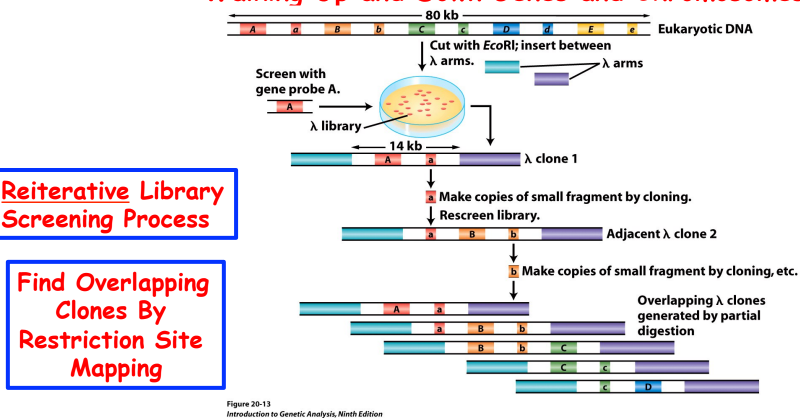
Sequencing (Compare With Protein Sequence) as Guides!

Sequence -----> GenBank

Step Three

#### Finding the Entire Factor VIII Gene? Walking & Sequencing

#### Walking Up and Down Genes and Chromosomes



Find Overlapping Clones By Restriction Site Mapping

Screening Process

Basis of Genome Projects & Whole Genome Sequencing

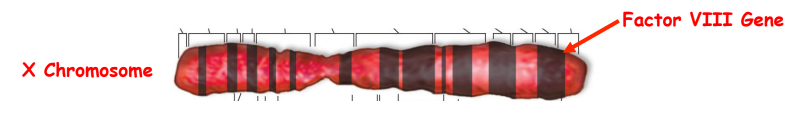
Key Concepts

How know Find Complete Factor VIII Gene?

Compare Protein & DNA Sequences

#### The Factor VIII Gene Was Found To Be Very Large

- 186,000 Nucleotides in Length (Won't Fit in One Phage Clone)
- 25 Introns
- 9,000 Nucleotide Coding Sequence (cDNA)
- · 2,351 Amino Acids in Protein



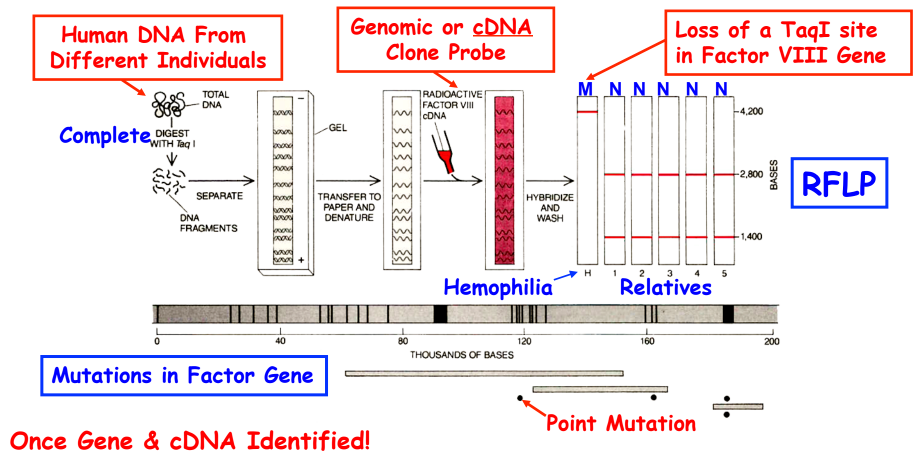
#### Factor VIII Mutations Occur Throughout the Gene

[*Haemophilia* 11, 481-491 (2005)]

VIII:C (%)	Family history	Consanguinity*	Inversion	Codon†	Mutation	Amino acid change	Exon	Conservation:
	Sporadic	NC	Normal	51	TTT → TCT§	Phe → Ser	2	FFFF, identical
.20	Sporadic	NC	Normal	80	$GTT \rightarrow GAT$	Val → Asp	3	VVVV, identical
	Sporadic	NC	Normal	102	GGT → GTT§	Gly → Val	3	GGGG, identical
!	Sporadic	NC	Normal	104	TCC → CCC§	Ser → Pro	3	SSSS, identical
	Sporadic	NC	Normal	143	GAG → AAG§	Glu → Lys	4	EEEE, identical
	Sporadic	NC	Normal	233	delCA§	Thr → fs (TGA-264)	6	
70	Inherited	NC	Normal	321	$GAA \rightarrow AAA$	Glu → Lys	8	EEEE, identical
)	Sporadic	NC	Normal	372	$CGC \rightarrow CAC$	Arg → His	8	RRRR, identical
	Inherited	NC	Normal	527	$CGG \rightarrow TGG$	Arg → Trp	11	RRRR, identical
L	Sporadic	NC	Normal	528	TGC → TAC§	Cys → Tyr	11	CCCC, identical
L	Inherited	NC	Normal	592	CAA → TAA	Gln → Stop	12	QQQQ, identical
l	Inherited	NC	Normal	864	delGACA	Gly → fs [TAA-867]	14	
					insCAATTAAATGAGAA§			
l	Sporadic	NC	Normal	948	insA§	Lys $\rightarrow$ fs (TGA-984)	14	
	Sporadic	NC	Intron 1	1107	AGG → TGG§	Arg → Trp	14	RGKK, dissimilar
	Sporadic	NC	Normal	1107	AGG → TGG§	Arg → Trp	14	RGKK, dissimilar
	Inherited	NC	Normal	1191-1194	delA	Ile → fs (TAG-1198)	14	
.40	Sporadic	NC	Normal	1191-1194	insA	Ile → fs (TAA-1220)	14	
	Sporadic	C	Normal	1227	delC§	Leu → fs (TGA-1231)	14	
.10	Sporadic	NC	Normal	1241	$GAC \rightarrow GAG$	Asp → Glu	14	DGGE, similar
	Sporadic	NC	Normal	1392	1392dcl1418§	Pro → fs (TAG-1446)	14	
	Incrited	C	Normal	1392	1392del1418§	Pro → fs (TAG-1446)	14	
	Sporadic	NC	Normal	1441	insA\$		14	
	Incrited	C	Normal	1441	insAS			
	Inherited	NC	Normal	1.502	CAG → TAG§	Gln → Stop	14	QREQ, dissimilar
	Inherited	NC	Normal	1504	delGT§	Val → fs (TGA-1517)	14	
	Sporadic	NC	Normal	1535	$TGG \rightarrow TGA$	Trp → Stop	14	WLWM, dissimilar
hibitor 96 BU								
	Sporadic	NC	Normal	1571	TAT → TAAS	Tyr → Stop	14	Y-YY, dissimilar
	Sporadic	NC	Normal	1.581	AAA → TAAS	Lys → Stop	14	KEKK, dissimilar
1,20	Sporadic	NC	Normal	1696	$CGA \rightarrow GGA$	Arg → Gly	14	RRRR, identical
.80	Sporadic	NC	Normal	1729	delAS	Gln → fs (TAA-1752)	1.5	
	Inherited	NC	Normal	1751	GAA → AAA§	Glu → Lys	15	EEEE, identical
	Sporadic	NC	Normal	1775	TTC → TCC§	Phe → Pro	16	FFFF, identical
	Sporadic	NC	Normal	1835	TGG → TGAS	Trp → Stop	16	WWWW, identical
.60	Sporadic	C	Normal	1882	ATC → ATAS	lle → Ile	17	IIII, identical
	Inherited	C	Normal	1966	CGA → CAA	Arg → Glu	18	RRRR, identical
	Sporadic	NC	Normal	1966	CGA → TGA	Arg → Stop	18	RRRR, identical

Need To Screen Across the Gene for Markers -- Family Specific

#### Factor VIII Gene Probes/ Sequence Can Be Used to Characterize Mutant Genes & Do DNA Testing for Carriers

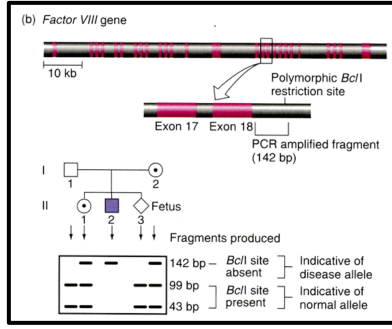


Use DNA Gel Blots (or PCR) & Factor VIII
Probes to Investigate Presence of
Mutant Alleles in Families (carriers)

Mutations Arise Independently in Families

## Using PCR and RFLPs (Markers) to Screen For the Hemophilia A Disease Allele/Gene

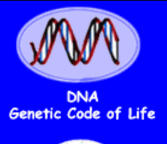
- 1. Use PCR To Amplify A Specific Factor VIII Gene Region
- 2. Use Restriction Enzyme
  (Bcl I) To Distinguish
  Between Normal Allele
  (1 Site) & Disease
  Allele (No Site)
  - = Normal Allele
  - = Disease Allele



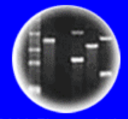
The 21<sup>st</sup>
Century
Approach!

- 1. Sequence the Entire Gene & Find Mutation
  - 2. Then
    Synthesize
    Primers to
    Test Family
    Members
    Using PCR

Only Can Do This With a Knowledge of DNA Sequence of Wild-type (Normal) and Disease Genes (Can Vary Family to Family)







**DNA Fingerprinting** 



Cloning: Ethical Issues and Future Consequences



Plants of Tomorrow

#### Step Four

How Find Factor VIII mRNA to Generate a cDNA for Protein Production in Host Cells?

Recall: Eukaryotic Genes Provide
Obstacles for Efficient Protein
Production in Genetically
Engineered Cells!

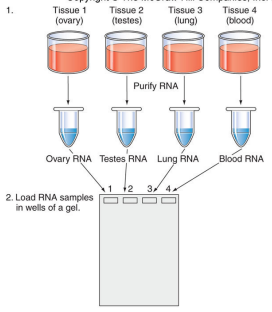
Introns! Switches!

#### Making the Drug

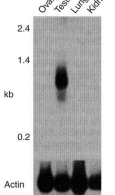
#### Need cDNA Not Gene

#### Factor VIII Gene Can Be Used to Find Out Where It is Active Using RNA Blots

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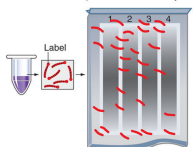


4. Wash away unhybridized probe. Make autoradiograph.



RNA Blot Is Like a DNA Blot Except That RNA is on Gel & Blotted

Separate RNA samples by gel electrophoresis.
 Blot onto filter. Expose filter to labeled hybridization probe.

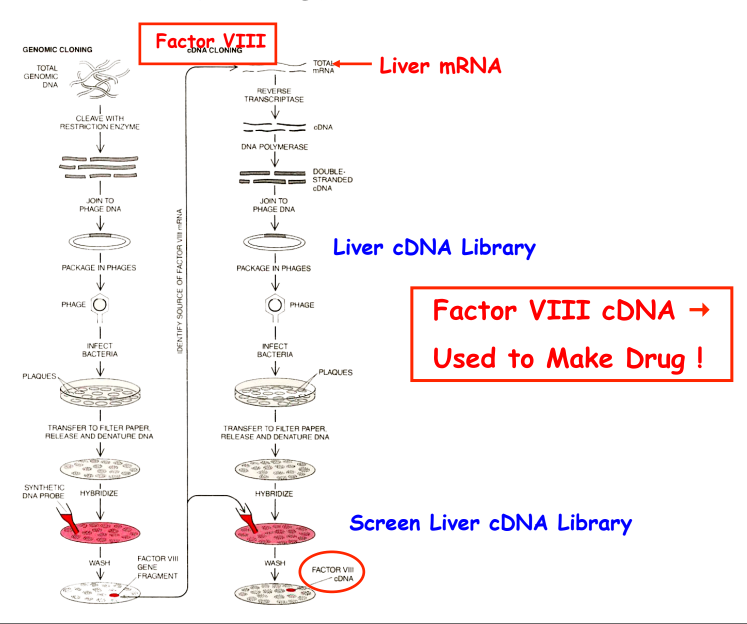


Factor VIII
Gene Is Highly
Active in Liver!

Could Also Use PCR (RT-PCR)

(4): Reprinted with permission from Nature 1990 Jul 19; 346(6281):216-7, Sinclair et al. © 1990 Macmillian Magazines Limited

## Using Factor VIII Gene Probe to Identify Factor VIII cDNA clone

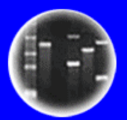


## DNA

#### DNA Genetic Code of Life



Entire Genetic Code of a Bacteria



**DNA** Fingerprinting

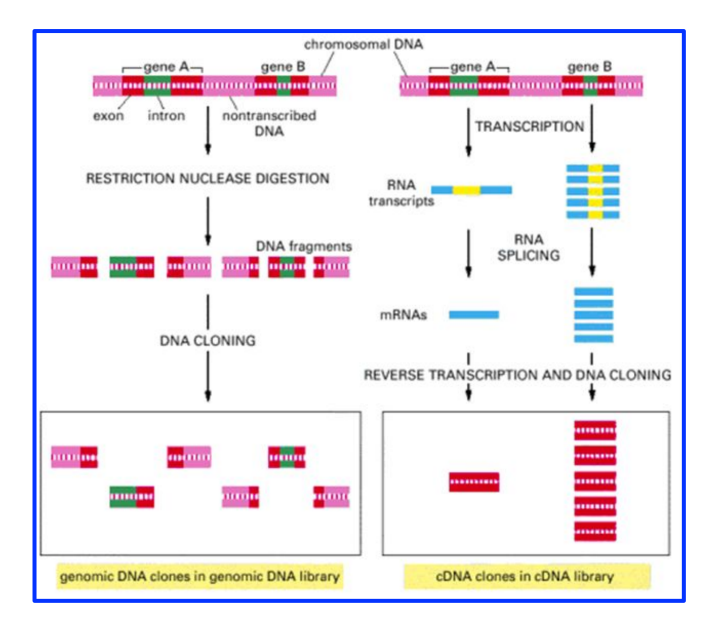


Cloning: Ethical Issues and Future Consequences

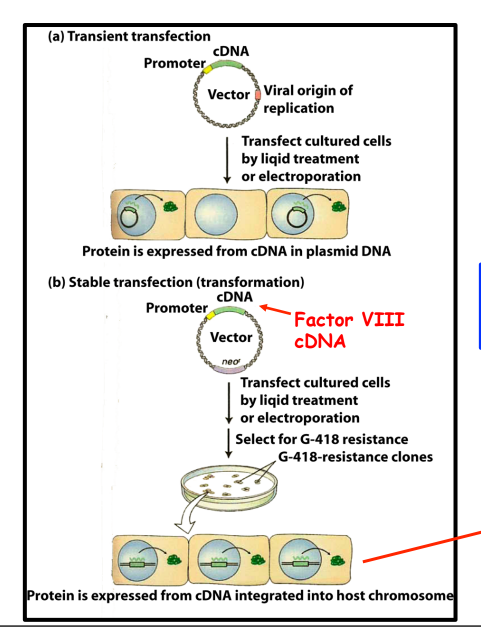


Plants of Tomorrow

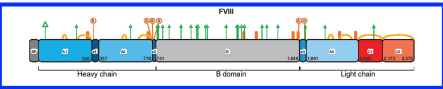
#### Genomic Libraries vs. cDNA Libraries A Review



## Engineer Factor VIII cDNA to Produce Protein in Host Cell & Synthesize Factor VIII in Mammalian Cells

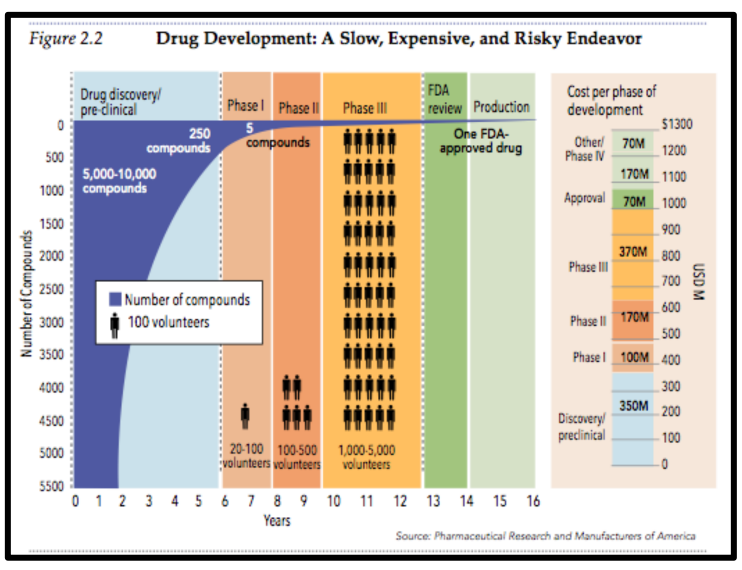


#### Why Mammalian Cells?



Purify Factor VIII Protein!

#### Need FDA Approval Before Recombinant DNA Drug Can Be Marketed and Used to Treat Patients



A Long and Expensive Process!

#### Recombinant Factor VIII



#### Bayer Biological Products EU



About Us

About Haemophilia For Kids

Research & Development

Press Releases

Recombinant Factor VIII

More Resources Haemophilia Centres in Europe

#### Related Links

Haemo-QoL Project Hemophilia Research Awards

#### Recombinant factor VIII

Recombinant factor VIII (rFVIII) is the antihaemophilic factor A, obtained using recombinant DNA technology. With this technology, pure protein is synthesized in the laboratory instead of being extracted from blood. In the following pages, it will be explained in detail how the knowledge and analysis of DNA, using the new instruments of molecular genetics, have represented both the beginning and follow-up stages in the development of recombinant FVIII.



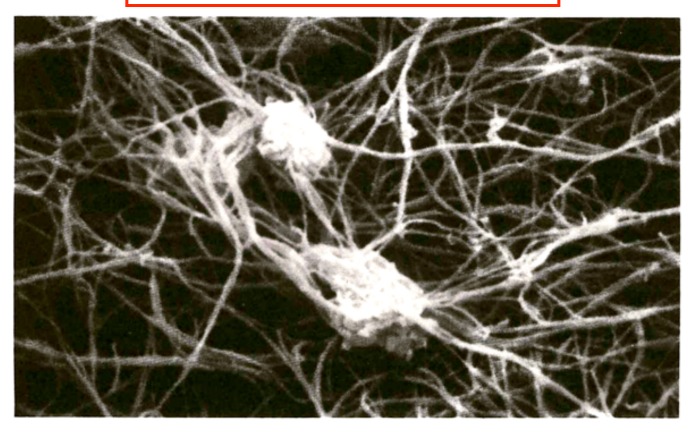
Prophylactic **Treatment** Costs \$300,000/ Yearl Most Hemophiliacs Use "On Demand" or As Needed

Factor VIII Gene Cloned In 1983

Factor VIII (Recombinant) Approved As Drug In 1993! Ten Years From Gene -> Drug! (Off Patent In 2011)

#### Using Factor VIII to Treat Hemophilia

#### Formation of a Blood Clot



FIBRIN STRANDS stabilize a blood clot at the site of a wound by trapping the platelets that form the bulk of the clot. The electron micrograph, which was made by Jon C. Lewis of Wake Forest University, shows a clot formed in a suspension of platelets and fibrin.

A clot in the bloodstream is the result of a complex cascade of enzymatic reactions culminating in the conversion of fibrinogen, a soluble protein, into insoluble fibrin strands. In hemophiliacs a crucial protein in the blood-clotting cascade is either missing or defective.

A Triumph of Genetic Engineering

#### The Future: Gene Therapy - A Permanent "Cure"

December 10, 2011

#### Treatment for Blood Disease Is Gene Therapy Landmark

By NICHOLAS WADE



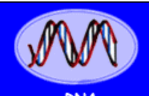
Gene Therapy Shows Promise for Treating Hemophilia

By ALICE PARK Monday, December 12, 2011

## The First Ever In-Human Gene Editing Will Try and Combat

Hemophilia

Factor IX - Hemoglobin B FDA-Approved Clinical Trial



Genetic Code of Life



Entire Genetic Code of a Bacteria



**DNA** Fingerprinting



Cloning: Ethical Issues and Future Consequences



Plants of Tomorrow

#### The Factor VIII Story -- A Summary

- 1. Purify Small Amounts of Factor VIII
- 2. Obtain Partial or Complete Amino Acid Sequence
- 3. Use the Genetic Code to Synthesize Degenerate DNA Probes
- 4. Isolate Factor VIII DNA Clones Complementary to Probe in Genome Library
- 5. Determine if Factor VIII Clones Contain the Complete Gene By Sequencing and Comparing With Protein Sequence
- 6. If Not, "Walk" to Obtain Overlapping DNA Clones That Collectively Contain the Factor VIII Gene
- 7. Sequence Clones To Determine Where the Factor VIII Gene Starts and Stops
- 8. Use Factor VIII Genome Probe to Find Out What Body Organ/Tissue Expresses the Factor VIII Gene
- 9. Make a cDNA Library From the Target Organ/Tissue and Isolate a Factor VIII cDNA Clone
- 10. Sequence the Factor VIII cDNA Clone and Compare With Factor VIII Gene Sequence to Map its Anatomy (I.e., introns, exons, swtiches) and Ensure That it Contains the Complete Protein Coding Sequence
- 11. Use Factor VIII cDNA and/or Genome Fragments as a Probe to Find RFLP Markers For Disease Alleles -- Or Sequence Disease Alleles to Find Relevant RFLP Markers By Comparison With Wild-Type Sequence
- 12. Insert Factor VIII cDNA Into an Expression Vector and Synthesize Factor VIII Protein in Host Cells (e.g., Mammalian Cells)