The Gene Gateway Workbook

A collection of activities derived from the tutorials at Gene Gateway, a guide to online data sources for learning about genetic disorders, genes, and proteins.



Using hereditary hemochromatosis as a model, access a variety of Web sites and databases to

- Learn about a genetic disorder and its associated gene.
- Identify mutations that cause the disorder.
- Find the gene on a chromosome map.
- Examine the gene's sequence and structure.
- Access the amino acid sequence of a gene's protein product.
- Explore the 3-D structure of the gene's protein product.

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Introduction

The Gene Gateway Workbook is a collection of activities with screenshots and step-by-step instructions designed to introduce new users to genetic-disorder and bioinformatics resources freely available on the Web. It should take about 3 hours to complete all five activities.

The workbook activities were derived from more detailed guides and tutorials available at the Gene Gateway Web site (<u>http://genomics.energy.gov/genegateway/</u>). The Gene Gateway Web site was created as a resource for learning more about the genes, traits, and disorders listed on the Human Genome Landmarks (HGL) poster, but it can be used to investigate any gene or genetic disorder of interest.

Many guides to genome Web resources are designed for bioscience researchers and are too technical for nonexperts. This workbook and other Gene Gateway resources target a more general audience: teachers, high school and college students, patients with disorders and their families, and anyone else who wants to learn more about how life works at a molecular level.

This workbook shows you how to get started using bioinformatics resources that often intimidate and overwhelm new users. It also demonstrates how information from one resource, such as annotated protein sequence data from Swiss-Prot, can be used to reinforce and clarify information available from another resource, such as three-dimensional (3-D) structures from Protein Data Bank (PDB). Gene Gateway provides users with a systematic approach to using multiple bioinformatics databases to gain a better understanding of how genes and proteins can contribute to the development of a particular genetic condition.

Using the genetic disorder hereditary hemochromatosis as a model, this workbook shows you how to access:

- Online Mendelian Inheritance in Man (OMIM) and GeneReviews to learn about a genetic disorder, its associated gene or genes, and common disease-causing mutations
- NCBI Map Viewer to find a gene locus on a chromosome map
- Entrez Gene and GenBank to examine the sequence and structure of a gene
- Swiss-Prot to find the annotated amino acid sequence of a gene's protein product
- Protein Data Bank and Protein Workshop to view and modify the 3-D structure of the gene's protein product

Skills gained by working through the activities in this workbook can be applied to learning about other genetic disorders, genes, and proteins.

This workbook and other genome science resources are available from the Web site for the genome programs of the Office of Biological and Environmental Research, U.S. Department of Energy Office of Science (<u>http://genomics.energy.gov/</u>).

Why use hereditary hemochromatosis as a model?

- Hereditary hemochromatosis, a disorder in which too much iron accumulates in certain tissues and organs, is caused by changes in the DNA sequence of a single gene, so the genetic basis of this condition is easier to understand than more complex disorders caused by alterations in multiple genes.
- The gene and its protein product are relatively well studied. Three-dimensional structures of the protein product are available in PDB, the international repository for macromolecular structure data.
- Hereditary hemochromatosis is the most common autosomal recessive disorder affecting individuals of Northern European descent (about 1 in 200 Caucasians develop hereditary hemochromatosis).
- Effective methods for treatment are available with early diagnosis.

Some basic concepts to understand before starting

- Genes are the basic physical and functional units of heredity. Each gene is located on a
 particular region of a chromosome and has a specific ordered sequence of nucleotides (the
 building blocks of DNA).
- Central dogma of molecular biology: DNA \rightarrow RNA \rightarrow Protein
 - Genetic information is stored in DNA.
 - Segments of DNA that encode proteins or other functional products are called genes.
 - Gene sequences are transcribed into messenger RNA intermediates (mRNA).
 - mRNA intermediates are translated into proteins that perform most life functions.
- Eukaryotic genes have introns and exons. Exons contain nucleotides that are translated into amino acids of proteins. Exons are separated from each other by intervening segments of DNA called introns. Introns do not code for protein, and they are removed when eukaryotic mRNA is processed. Exons make up segments of mRNA that are spliced back together after the introns are removed; the intron-free mRNA is used as a template to make proteins.
- Special cellular components (ribosomes) use the triplet genetic code to translate the nucleotides of a mRNA sequence into the amino acid sequence of a protein. A Table of Standard Genetic Code is provided in the back of this workbook.
- There are 20 different amino acids. Proteins are created by linking amino acids together in a linear fashion to form polypeptide chains. See the Table of Standard Genetic Code in the back of this workbook for single-letter and three-letter abbreviations for the 20 different amino acids.
- Protein polypeptide chains fold into 3-D structures that can associate with other protein structures to perform specific functions.

Activity 1 Online Resources: OMIM and GeneTests

- Learn about the genetic disorder and its associated gene.
- Identify mutations that cause the disorder.

Online Mendelian Inheritance in Man (OMIM)

OMIM is a large, searchable, up-to-date database of human genes, genetic traits, and disorders created and edited by researchers at Johns Hopkins University. The OMIM database is accessible through the National Center for Biotechnology Information (NCBI) suite of online resources. Each record in OMIM summarizes research defining what is currently known about a particular gene, trait, or disorder.

To access OMIM, let's go to the NCBI Web site (<u>http://www.ncbi.nlm.nih.gov/</u>), and then click on **OMIM** above the search box at the top.

NCBI HomePage - Mi	crosoft Internet Explorer							
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Address 🕘 http://www.ncb	i.nlm.nih.gov/	💌 🔁 Go						
Since National Center for Biotechnology Information National Library of Medicine National Library of Medicine National Institutes of Health Health PubMed All Databases BLAST								
Search All Database	es for Go							
SITE MAP Alphabetical List	What does NCBI do? Established in 1988 as a national resource	Hot Spots Assembly Archive						
Resource Guide About NCBI	for molecular biology information, NCBI creates public databases, conducts research in computational biology, develops software tools for analyzing genome data, and disseminates biomedical information – all for	Clusters of orthologous groups						
An introduction to NCBI GenBank	the better understanding of molecular processes affecting human health and disease. <u>More about NCBI</u>	Coffee Break, Genes & Disease, NCBI Handbook						
Sequence submission	Genome Reference Consortium	Electronic PCR						
support and software	The <u>Genome Reference Consortium</u> (GRC) has been formed to continue the improvement of the human and mouse	Entrez Home						
Literature databases PubMed, OMIM, Books, and PubMed	genome reference assemblies. The goal of the GRC is to fix the small number of loci that may be misrepresented in the reference assembly, fill the remaining gaps, and to produce alternate representations of	 Gene expression omnibus (GEO) 						
Central	complex loci.	Human genome resources						
Molecular databases	PubMed Central	▶ Influenza Virus Resource						
sequences, structures,	<u>PubMed Central</u> is an archive of biomedical and life sciences journals.	► Map Viewer						
æ		🔮 Internet						

A screenshot of the OMIM home page is shown below.



URL for OMIM home page: http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db=OMIM

Although the easiest way to search OMIM is to simply type a disorder name in the search box at the top, another option for searching OMIM is to use search field qualifiers. By adding search field qualifiers in square brackets to each search term and combining terms using Boolean operators (OR, AND, or NOT), you can execute a much more specific search in a single step.

This activity demonstrates only how to use a couple of OMIM's field qualifiers. More information about field qualifiers and other advanced search options is available from OMIM Help (<u>http://www.ncbi.nlm.nih.gov/Omim/omimhelp.html</u>). In addition to OMIM, field qualifiers can be used to search other NCBI information systems such as PubMed (a resource for accessing bibliographic citations from biomedical literature) and nucleotide and protein sequence databases.

Most genes, disorders and traits listed on the Human Genome Landmarks (HGL) poster were taken from the title fields of OMIM records. The field qualifier for the title field is [TI] or [TITL]. Since we selected our disorder from the HGL poster, we also know that hemochromatosis is found on chromosome 6. The field qualifier for specifying a particular chromosome is [CH] or [CHR].

1. To use a field qualifier in your search, simply add the qualifier to the end of your search term. For example, to search for hemochromatosis on chromosome 6 enter **hemochromatosis[TI] AND 6[CHR]** as shown in the search box below. Be sure to capitalize any Boolean operator (AND, OR, and NOT) you use in your search statements. Click **Go** to submit your search.

All Databases	PubMed	Nucleotide	Protein	Genome	Structure	PMC	Taxonomy
Search OMIM		💽 for h	emochromat	osis[TI] AND (6[CHR]		Go Clear
							

NOTE: Limiting a search to a particular chromosome may not work for disorders caused by alterations in multiple genes, such as breast cancer or diabetes. These disorders are linked to genes on several different chromosomes; therefore, limiting your search to just one chromosome may not yield the best results.

2. The search should return one result. Clicking on the MIM number ± 235200 opens the full OMIM record for hemochromatosis shown below.

DMIM - HEMOCHROMA	ATOSIS; HFE - Microsoft Internet Explorer	
File Edit View Favorites	Tools Help	27
Address a http://www.ncbu	.nlm.nih.gov/entrez/dispomim.cgPld=235200 👻	🔁 Go
S NCBI	Deline Mendelian Inheritance in Man	2
MIM +235200 Description Clinical Features Other Features Inheritance Mapping Heterogeneity Molecular Genetic	All Databases PubMed Nucleotide Protein Genome Structure PMC OMIM Search OMIM v for Go Clear Limits Preview/Index History Clipboard Details Display Detailed v Show 20 v Send to v	
Genotype/Phenot Correlations Diagnosis	+235200 GeneTests, Link HEMOCHROMATOSIS; HFE	s
Population Geneti Pathogenesis	Alternative titles; symbols	
Gene Structure Nomenclature	HLAH HEMOCHROMATOSIS, HEREDITARY; HH HFE GENE, INCLUDED; HFE, INCLUDED	
Animal Model History	Gene map locus <u>20p12, 6p21.3</u>	
Allelic Variants View List	TEXT	
References	DESCRIPTION	
Creation Date	The clinical features of hemochromatosis include cirrhosis of the liver, diabetes, hypermelanotic pigmentation of the skin, and heart failure. Primary hepatocellular carcinoma (HCC; <u>114550</u>),	~
ð	🌒 Internet	

- 3. Let's examine some of the features of this record:
 - Each record includes a blue navigation menu on the left with quick links to different sections within the record.
 - Each OMIM record is assigned a unique six-digit MIM number located at the top of each entry. For hereditary hemochromatosis, the MIM number is 235200. As a unique identifier, the MIM number can be used to search other databases for information about a particular disorder. Clicking on the MIM number link will open the record in a simpler, frame-free format more suitable for printing.
 - The plus sign (+) in front of the MIM number means that this entry refers to a phenotype associated with a gene of known sequence. In other records, a number sign (#) in front of the six-digit MIM number means that a phenotype may be associated with multiple loci. For additional information about MIM number symbols, see OMIM Frequently Asked Questions (http://www.ncbi.nlm.nih.gov/Omim/omimfaq.htmlmim_number_symbols).

Below the MIM number, you will find the disorder name and the official gene symbol. The
official gene symbol, which is HFE for hemochromatosis, serves as a unique identifier for a
gene. To be "official," a gene symbol must have been approved by the HUGO Gene
Nomenclature Committee (<u>http://www.genenames.org/</u>). The gene symbol is especially
useful when searching other databases (such as sequence, genome-mapping, and
structure databases) for gene-specific information.



NOTE: For single-gene disorders like hemochromatosis, the official gene symbol usually will be included in the record title. For complex disorders like breast cancer, official symbols for associated genes will be described in the first paragraph of text.

- The gene map locus describes where a gene can be found on a chromosome. For the gene locus **6p21.3**, 6 is the chromosome number, p indicates the short arm of the chromosome, and 21.3 is a number assigned to a particular region of the chromosome. The gene map locus links to OMIM's Gene Map, a table of genes organized by cytogenetic location.
- The amount of text within an OMIM record varies according to what is known about a particular gene, disorder, or trait. Since hemochromatosis is well studied, a lot of information is known about this disorder and its gene. Some different types of information that may be included in an OMIM record are disorder description, inheritance, genotype and phenotype correlations, diagnosis, population genetics, gene structure, gene function, and animal models.
- Selecting the **Gene Structure** link (in the blue navigation column on left) provides information about the size and number of exons in the gene.
- Although not a part of every OMIM record, another useful section is Allelic Variants (see link in the blue navigation column on left). This section typically describes some of the most notable gene mutations associated with the development of disorders. Select the View List link under Allelic Variants to see a listing of important mutations identified for the HFE gene. At the top of the list of allelic variants is the most common mutation known to cause hereditary hemochromatosis. The standard notation for this allelic variant is CYS282TYR. This means that a mutation occurs in the DNA sequence that changes the amino acid at position 282 of the gene's protein product from cysteine to tyrosine.



4. Another way you can modify your OMIM search is to use **Limits**. Under the OMIM search box near the top of the page, click on the **Limits** tab (shown below).



5. The Limits page provides a variety of options that you can use to narrow your search. For example, instead of using the search field qualifier [CHR] to narrow your search to genes on chromosome 6, you could select the chromosome from the Limits page. You also can search by MIM number or limit your search terms to the title or other field of an OMIM record.

6. Let's use options on the Limits page to determine how many genes in the human genome have been described in OMIM. Put a check beside the **MIM Number Prefix** options for **gene with known sequence** and **gene with known sequence and phenotype** as shown in the screenshot below. Then click the **Go** button beside the search box at the top of the page.



7. You should retrieve over 12,000 search results. Of the estimated 20,000 to 25,000 genes in the human genome, about 12,000 genes have records in OMIM. You may want to test your new search skills by using OMIM to search for other genes or genetic conditions. In addition to OMIM, another good resource for learning about genetic disorders and associated genes is the GeneTests Web site, which is described in the next part of this activity.

GeneTests

The GeneTests Web site is a medical genetics information resource developed by researchers and healthcare professionals and funded by the National Institutes of Health. In addition to providing up-to-date, authoritative reports (GeneReviews) on genetic disorders, the site also includes educational materials (e.g., fact sheets on genetic testing and counseling, PowerPoint slides, and an illustrated glossary) and online directories of genetic laboratories and clinics.

This activity focuses on accessing and using genetic disorder information available from GeneReviews. All entries are written and reviewed by physicians, so the language is similar to that of medical text. While the amount and kind of content can vary greatly from record to record in OMIM, all reports in GeneReviews will provide similar kinds of information and share the same organizational structure.

Let's go to the GeneTests Web site (<u>http://www.genetests.org/</u>) to find a GeneReview for hereditary hemochromatosis.



1. Click on **Reviews** in the navigation bar at the top.

2. Once you get to the **Search by Disease** screen at *GeneReviews*, enter **hemochromatosis** into the search box.

3. Beside the search result "HFE-Associated Hereditary Hemochromatosis," select the *Reviews* link to access the hereditary hemochromatosis review shown below.

HFE-Associated Her	editary Hemochromato	sis GeneReviews NCBI B	ookshelf - Microsoft Inte	rnet Explorer			
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Kris V Ko	wdley, MD				<u>Summary</u>		
Professor of	Medicine, Gastro	penterology/Hepatol	ogy		<u>Diagnosis</u>		
Founder and University of	l Director, Iron O Washington	verload Clinic			Clinical Description		
Seattle, WA					Differential Diagnosis		
<u>kkowdley@u</u>	washington.edu	ĺ.			Management		
Jonathan	F Tait, MD, P	hD			Genetic Counseling		
Professor, D	epartment of Lab	poratory Medicine		Γ	Molecular Genetics		
University of	Washington Sch	ool of Medicine					
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http://www.ncbi.nlm.nih	.gov/entrez/utils/fref.fcgi?Prid	1=5504&uid=9999&db=books&url	=http://www.genetests.org/se	ervlet/access?id=8		Internet	>
Tup.//www.ncbi.nim.nin	.gov/enu/ez/udis/frei.tcgi?Prid	1=2204&uiu=3333&uD=D00KS&UN	=nup.//www.genetests.org/se	n viet/access?id=8		Internet	

4. Access the **Molecular Genetics** section for a brief overview of this disorder's molecular basis. This section provides the official symbol for the gene associated with this disorder, the gene's chromosomal locus, name of the gene's protein product, links to records for this gene in other databases, descriptions of mutations known to cause the disorder, and summaries of the protein's normal function and structure. Other sections in this report describe disease characteristics, diagnosis and testing, treatments, and genetic counseling issues. Use the information in GeneReviews and OMIM to answer the Questions for Activity 1 on the Hereditary Hemochromatosis Worksheet included in the back of this workbook.

Activity 2 Online Resource: NCBI Map Viewer

- Find the hereditary hemochromatosis gene on a chromosome map.

NCBI Map Viewer is a Web-based tool for viewing and searching an organism's complete genome. Users also can view maps of individual chromosomes and zoom in to specific regions within chromosomes to explore the genome at the sequence level.

Map Viewer provides access to several different types of maps for different organisms. Many of these maps are meaningful only to scientific researchers. A discussion of all the different types of maps and genomic data is beyond the scope of this activity, which will focus only on how to locate a specific gene locus on a chromosome map.

From the NCBI home page (<u>http://www.ncbi.nlm.nih.gov/</u>), select **Map Viewer** from the alphabetized list of "Hot Spots" on the right. A screenshot of the **NCBI Map Viewer** home page is shown below.

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SNCBI Home GenBar	k BLAST					^
Map Viewer Home	and the first sector of the sector of	1972 3107 197 197 197	Tel et anno 1991 *		Help	
The Map View	wer provides a wide va	riety of genome mapping and sequen	cing data. More		¥ A	
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	Mammals				(13)	
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for:	Scientific n	ame Common name	Build	Tools		
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Tools Legend 🔻	Macaca mu	latta rhesus macaque	Build 1.1	9 B G		
Search or Browse the Genome	Pan troglody	tes chimpanzee	Build 2.1	9 B G		
B BLAST	Rodents				(2)	
G Genome Resources page	Scientific n	ame Common name	Build	Tools		
News 🔻	Mus muscu	lus laboratory mouse	Build 37.1 Build 36.1	9 B G 9 B		
Annotation update released for human Mar 24	Rattus norve	egicus rat	RGSC v3.4	9 B G		
genome build 36	► Monotreme	5			(1)	
An annotation update for the numan genome (NCB) Build 36.3) more	► Marsupials				(1)	
Show all	Other Mamr	nals			(6)	
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NODUL	Invertebrates				(9)	
NCBI Home NCBI Web Search	Protozoa	B			(17)	
NCBI Site map	► Plants 🔍				(45)	
Taxonomy	🔻 Fungi 🔍	B			(17)	
Entrez (Global Query)	Scientific nam	e Commo	on name Build	Tools	-	
Map Viewer FTP	Aspergillus clar	vatus	Build 1.1	9 B G		~
http://www.ncbi.nlm.nih.gov/Genbank/index.html				🌍 Intern	iet	.:

URL for NCBI Map Viewer: http://www.ncbi.nlm.nih.gov/mapview/

On the Map Viewer home page, in the list of **Primates**, click on the *Homo sapiens* (human) **Build 36.3** link to view the entire human genome. This will launch the *Homo sapiens* genome view shown in the following screenshot.

Entrez Genome view	ew - Microsoft Internet Explorer							
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Address 🕘 http://www.ncb	cbinim.nih.gov/projects/mapview/map_search.cgi?taxid=9606							
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Human Genome Resources NCBI Handbook	Human Genome Resources NCBI Handbook NCBI Handbook							
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Homo sapiens genome view: http://www.ncbi.nlm.nih.gov/mapview/map_search.cgi?taxid=9606

In Activity 1, we learned that the official symbol for the hereditary hemochromatosis gene is HFE, and its locus is 6p21.3. Let's find the HFE gene on chromosome 6.



1. In the search box at the top of the page, enter **HFE[sym]** as shown below. The [sym] search field qualifier specifies your search so that only hits for a gene with the symbol "HFE" are generated for your query.

PubMed	Nucleotide	Protein	Genome	Gene	Structure	PopSet	Τε
Search for	HFE[sym]	on chrom	osome(s)		assembly All	•	Find

2. Red tick marks should be displayed on chromosome 6 in the genome view, indicating the approximate location of the HFE gene in the middle of the short arm of chromosome 6. The "44" below chromosome 6 (see screenshot below) indicates the number of hits for our query. About 44 different maps in Map Viewer include the gene symbol "HFE."

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3. In the genome view, click on the number $\underline{6}$ link below the chromosome. This will open a view of chromosome 6 that should look like the screenshot below. In the next step we will modify this view so we can see an ideogram showing the region of chromosome 6 where the HFE gene can be found.



4. Let's modify the display options by clicking on **Maps & Options**. This will open a window for customizing map options. Make the following adjustments. Before you click the **Apply** button, your options window should resemble the screenshot below.

- Remove all maps listed under **Maps Displayed (left to right)** except the **Gene** map. To remove a map, select it with your mouse and then click the **REMOVE** button.
- Under **Available Maps** select **Ideogram** (you will need to scroll through more than half of the available maps) and click the **ADD** button.
- The **Maps Displayed** list should look like the screen shot below. The **Gene** map should be designated as your master map. To make a map the master, select it with your mouse and then click the **Make Master/Move to Bottom** button. In the chromosome view, a master map is shown at the right edge of the display along with its details and descriptive text.
- Under **More Options** near the bottom of the window, change **Page Length** from 30 to 10. The Page Length option is highlighted in the screenshot below. This will display 10 labeled genes (rather than 30) in the master map.

http://www.ncbi.nlm.nih	gov - Map Viewer - Microsoft Intern	et Explorer		
Organism: Homo	o sapiens			<u>Help</u>
Chromosome: 6	Region Shown:	21200353.75 382	291906.25	
Available Maps:		Maps Dis	played (left to right):	
Org: human 🖌	Assembly: reference	~	Change Assembly	
Sequence Maps Ab initio		[] Ideogram [R] Gene	Move UP Move DOWN	
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More Options:				
■ Show Connection	ons Verbose Mode			
Compress Map: of	f 🔽 Auto Compress if >	350 px		
Thumbnail View:	 ● default (ideogram) ○ 1 	master		
OK Apply Close				
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How the Maps & Options window should look

• Click **Apply** at bottom and close screen.

About the maps

Ideogram – Shows the G-banding pattern of a chromosome at 850-band resolution.

Gene – Includes genes identified on segments of genomic sequence called contigs. A contig is a group of cloned (copied) pieces of DNA representing overlapping regions of a particular chromosome.

5. The new map of chromosome 6 should resemble the following screenshot. Notice that the red dots indicating the position of the HFE gene on the sequence maps appear to line up with the ideogram at the 6p22.2 chromosome band, not 6p21.3.

🙆 Map Viewer - Micros	soft Internet Explorer
File Edit View Favorites	s Tools Help
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PubMed	Entrez BLAST OMIM Taxonomy Structure
Search	Find Find in This View Advanced Search
Human genome	Homo sapiens (human) Build 36.3 (Current) BLAST The Human Genome
overview page	Chromosome: <u>1 2 3 4 5 [6] 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 X Y MT</u>
(Build 30.3) Human genome	Query: HFE[sym] [clear]
overview page	Master Map: Genes On Sequence Summary of Maps Maps & Options
	Region Displayed: 21M-38M bp Download/View Sequence/Evidence
Map Viewer Home	Ideogram
Map Viewer Help	HGNC svprdlevmmhmsts SNP best RefSeq 6p22.3-1
	⁶ ¢ ²²⁺² + <u>OMIMHGNC svprdlevmmhmsts SNP</u> best RefSeq 6p21.3
Human Maps	⁶ / ₆ ^{22.1} + <u>OMIMHGNC svprdlevmmhmsts SNP</u> best RefSeq 6p21.3
БТР	BTN3A1 + HGNC svprdl evmmhmsts SNP best RefSeq 6p22.1
Data As Table	$\underline{LOC340192}$ + \underline{sv} \underline{dlevmm} sts best RefSeq 6p22.1
View	^{6+21.32} + <u>OMIMHGNC svprdlevmmhmsts SNP</u> best RefSeq 6p21.3
Maps & Options	<u>OR5U1</u> + <u>HGNC svprdlevmmhm</u> <u>SNP</u> best RefSeq 6p22.1
Compress	^{6+21.31} ^{36#} <u>ATP6V1G2</u> * <u>OMIMHGNC svprdlevmmhmsts SNP</u> best RefSeq 6p21.3
Map 🗖	<u>IHPK3</u> • <u>OMIMHGNC svprdlevmmhmsts SNP</u> best RefSeq 6p21.31
Region Shown: 21M	PPARD + OMIMHGNC svprdlevmmhmsts SNP best RefSeq 6p21.2-1
38M Go	Summary of Maps:
	Map 1: Ideogram
out	Region Displayed: 6p22.3-6p21.2
2000 B in	Map 2: Genes On Sequence <u>Table View</u>
You are here:	Region Displayed: 21M-38M bp Download/View Sequence/Evidence
Ideogram	Total Genes On Chromosome: 1843 [16 not localized]
	Genes Labeled: 10 Total Genes in Region: 675
662112 III	
ê	🔮 Internet

Features of the Genes-seq map (the master map in the screenshot above):

- The portion of chromosome 6 displayed in Map Viewer is highlighted on the ideogram in the blue navigation column on the left. Rounding to the nearest million, the region displayed begins at about the 21 millionth nucleotide and ends at about the 38 millionth nucleotide of the DNA sequence of chromosome 6. The total DNA sequence for chromosome 6 is about 171 million base pairs long.
- Clicking on the Ideogram or Genes_seq maps (not the labels) will open a pop-up window with options for zooming in on the displayed maps. You can also zoom in and out using the zoom option in the blue navigation column.
- Map Viewer displays 10 labeled genes on the Genes_seq map. To see a more complete listing of genes in this region of the chromosome, select the Data As Table View link above Maps & Options in the blue navigation column on the left. The Data As Table View shows where genes start and stop in the chromosome's DNA sequence.

- The Genes_seq map provides links to gene-specific entries in other databases.
 - HFE Links to the HFE entry in NCBI's Entrez Gene database that brings together a variety of gene-specific information together in one interlinked system.
 - OMIM Links to the HFE entry in the Online Mendelian Inheritance in Man (OMIM) database covered in Activity 1.
 - HGNC Links to the gene symbol report maintained by the HUGO Gene Nomenclature Committee.
 - sv The Sequence Viewer link lets you drill down to the genome sequence level. This link takes you to a graphic showing the gene's position within the genomic sequence.
 - o pr Links to the reference sequence of the gene's protein product.
 - \circ dl Links to a page for downloading the sequence data for a particular chromosome region.
 - ev Links to Evidence Viewer, which provides biological evidence supporting a
 particular gene model showing exons and other features of a gene. It displays all
 RefSeq models, GenBank mRNAs, known or potential transcripts, and ESTs
 (expressed sequence tags) that align to the area of interest.
 - mm Links to Model Maker, which allows you to view the evidence used to build a gene model based on assembled genomic sequence. You can also create your own version of a model by selecting exons of interest.
 - hm Links to Homologene, a resource for comparing genes in homologous segments of DNA from different organisms.
 - sts Links to UniSTS, a comprehensive database that integrates genetic marker and mapping information. A sequence tagged site (STS) is a short (200 to 500 base pairs) DNA sequence that has a single occurrence in the human genome. Detectable by polymerase chain reaction (PCR), STSs are useful for localizing and orienting the mapping and sequence data reported from many different laboratories and serve as landmarks on the developing physical map of the human genome.
- 6. Let's zoom out to view the entire chromosome using the Maps & Options window.
 - Click on Maps & Options again to open the options window.
 - Delete the numbers defining the **Region Shown** at the top of the options window. This will modify the display so it shows the entire chromosome.
 - Under **More Options** near the bottom of the window, change **Page Length** from 10 to 20. The Page Length option is highlighted in the screenshot on the next page. This will display 20 labeled genes in the master map and should provide enough space on the screen to view the entire chromosome with readable labels for the chromosome bands.
 - Once the Maps & Options window resembles the screenshot on the following page, click the **Apply** button at the bottom and close the box.

http://www.ncbi.nlm.nih	n.gov - Map Viewer - Microsoft Intern	net Explorer						
Organism: Homo Chromosome: 6	o sapiens Region Shown:			<u>Help</u>				
Available Maps:		Maps Dis	played (left to right):					
Org: human 💌	Assembly: reference	~	Change Assembly					
Sequence Maps Ab initio Assembly Celera Genes Celera Transcripts	ADD>>	[] Ideogram [R] Gene	Move UP Move DOWN Make Master/Move to Bot	ttom				
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Thumbhan view.		master						
OK Apply Close								
🛃 Done			🏟 Internet	.:				

7. Your view of chromosome 6 should resemble the following screenshot. Scroll down to the bottom of the map to examine the **Summary of Maps** section and use this information and the map of chromosome 6 to answer questions for Activity 2 on the Hereditary Hemochromatosis Worksheet in the back of this workbook.

ap Viewer - Micros	oft Internet Explorer						
Edit View Favorites	Tools Help						
BubMad	.nlm.nlh.gov/projects/mapview/maps.cg	PTAXID=96068.CHR=68.BEG=8	NEND-SMAP	P0=ideogr%2Cgenes-r&VERBOSE=ON&COMPRESS=o	fl&WIDTH=3508/SIZE=20&OV	R=&QSTR=HFE%5Bsym%5	iD&QUERY=uid%2
earch	Endez		Fin	d Find in This View	Taxonomy	ranced Search	suucture
earen			1 114				6
man genome	Homo sapiens (hu	man) Build 36.	3 (Cu	<u>rrent)</u>	Ŀ	SLAST The Hu	man Genom
erview page	Chromosome: 12	345[6]789	10 11 1	12 13 14 15 16 17 18 19 20 21	22 X Y MT		
man genome	Query: HFE[sym]	[clear]					
erview page	Master Map: Gene	s On Sequence	е	Summ	ary of Maps		Maps & Options
and 35.1)	Region Displayed: 0-1	71M bp			Down	load/View Sequ	ence/Evidend
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p Viewer Help		PRPF4B	+ ON	MIMHGNC svprdlevmmhmsts	SNP best RefSeq	6p25.2	PRP4 pre-
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<u>60</u>	6-21 -	MRPL2	+	HGNCsvprdlevmmhmsts	SNP best RefSeq	6p21.3	mitochon
out	6422.01 6422.01	C6orf138	•	HGNC svprdl evmmhmsts	SNP protein	6p12.3	chromoso
200M	6+22+32 - 6+22-33 -	- ORC3L	• <u>ON</u>	MIMHGNC svprdl evmmhmsts	SNP best RefSeq	6q14.3-q16.1	origin reco
In horo:	6425.1 - 1599 6427.2 -	NCOA7	+ <u>ON</u>	MIMHGNC svprdl evmmhmsts	SNP best RefSeq	6q22.31-q22.3	2 nuclear re
eogram	6423-3 - 6424-1 - 6424-2 -	ECHDC1	•	HGNC svprdl evmnhmsts	SNP best RefSeq	6q22.33	enoyl Coe
	9121-0 - 6425-1 - 6425-2 -	OR2A4	•	HGNC svprdlevmmhm	SNP best RefSeq	6q23	olfactory
	6425.0 - 1649-	CCRL1P	+	HGNC svprdlevmm	SNP best RefSeq	6q24.1	chemokin
	6427 - 🕒 17 mil 🧌 🖾	FNDC1	+ <u>ON</u>	MIMHGNC svprdlevmmhmsts	SNP best RefSeq	6q25	fibronectin
6415 - 6415 -		SLC22A1	+ <u>ON</u>	MIMHGNC svprdlevmmhm	SNP best RefSeq	6q26	solute carr
422 -	Summary of Maner						
6420 -	Summary of Maps:						

Gene Gateway: A Web Companion to the Human Genome Landmarks Poster http://genomics.energy.gov/genegateway/

Activity 3 Online Resources: Entrez Gene and GenBank

- Examine gene sequence and structure.

This activity covers how to use NCBI's Entrez Gene to access the genomic DNA sequence of the hereditary hemochromatosis gene. We will examine some features of a record from NCBI's GenBank and learn about the structure (e.g., intron and exon composition, coding sequence) of a gene.

In sequence databases such as GenBank, genomic DNA sequences from eukaryotic organisms contain both exons and introns, while mRNA sequences are intron-free DNA sequences. All sequences in GenBank and similar repositories use the DNA bases adenine (A), cytosine (C), guanine (G), and thymine (T) to represent each nucleotide. Even mRNA sequence records use A, C, G, and T where T is used to replace each uracil (U) in the mRNA sequence.

Entrez Gene is a NCBI resource that serves as a single-query interface for accessing sequence and other biological information for specific genes from a variety of sequenced organisms.

To begin, let's go to the Entrez Gene home page.

Gene Home - Microso	oft Internet Explorer				
File Edit View Favorites	Tools Help	N			
Address a http://www.ncbl	nlm.nlh.gov/sites/entrez?db=gene	🛩 🛃 Go			
S NCBI	Entrez Gene	My NCBI F7 [Sign In] (Register)			
All Databases	PubMed Nucleotide Protein Genome	e Structure PMC Taxonomy Books OMIM			
Search Gene	💌 for	Go Clear			
	Limits Preview/Index History Clipboa	rd Details			
Entrez Gene Home	Entrez Gene is a searchable database of genes, from <u>RefSeq</u> genomes, and defined by sequence and/or located in the NCBI Map Viewer				
About FAQ Help Gene	News Limit by Chromosomal Region; Sort Options. News archives Sample Searches				
Handbook	Find genes by	Search text			
Downloads	free text	human muscular dystrophy			
(FTP)	partial name and multiple species	transporter[title] AND ("Drosophila melanogaster"[orgn] OR "Mus musculus"[orgn])			
Mailing Lists	chromosome and symbol	(II[chr] OR 2[chr]) AND adh*[sym]			
	associated sequence accession number	M11313[accn]			
Gene	gene name (symbol)	BRCA1[sym]			
Reisey	publication (PubMed ID)	11331580[PMID]			
Feedback	Gene Ontology (GO) terms or identifiers	"cell adhesion"[GO] 10030[GO]			
Help Desk Corrections	Genes with variants of clinical significance (under development)	gene snp clin[filter]			
ล		😵 Internet			

http://www.ncbi.nih.gov/entrez/query.fcgi?db=gene

1. In the search box at the top of the page, enter **HFE[sym] AND Human[orgn]**. Be sure to capitalize any Boolean operator (AND, OR, and NOT) you use in your search statements.

All Databa	ises PubMed	Nucleotide	Protein	Genome	Structure	PMC	Taxonomy
Search Ger	ie 🔺	for HFE[sym]	AND Human	[orgn]		G0 CI	ear

Search Tip: Adding [sym] to the end of your query term tells Entrez Gene that you are searching by gene symbol only. If you do not specify that you want to search the gene symbol field, the search will return multiple records that include the query term anywhere within its text. Adding [orgn] to a search term limits the search to genes from a specific organism. For more information on options for refining your search, see the Search Field Descriptions and Qualifiers section of Entrez Help: http://www.ncbi.nlm.nih.gov/entrez/query/static/help/Summary_Matrices.html

2. Submitting this search should retrieve a single result. The HFE record is shown below.

Entrez Gene: HFE hemochromatos	is [Homo sapiens] - Microsoft Internet Explorer	
le Edit View Favorites Tools Help		729
idress an http://www.ncbunim.nih.gov/ste	s/entrez	2
S NCBI	Entrez Gene	My NCBI E
All Databases	PubMed Nucleotide Protein Genome Structure PMC Taxon	omy Books OMIM
Search Gene	for HFE[sym] AND Human[orgn] Go Clear Save Search	
Limits Preview/In	dex History Clipboard Details	
Display Full Report	 Show 20 Sort by Relevance Send to 	
All: 1 Current On	ly: 1 Genes Genomes: 1 SNP GeneView: 1 😿	
1: HFE hemochrom	atosis [Homo sapiens]	f Entrez Gene Home
GeneID: 3077	updated 13-Jun-200	8 Table Of
Summary		Contents
Official Symbol		Summary
Official Symbol	provided by <u>HGNC</u>	Genomic regions,
Official Full Name	hemochromatosis	Genomic context
Drimony course	provided by HGNC	Bibliography
Primary source	HGNC:4886	HIV-1 protein
See related	Ensembl:ENSG00000010704; HPRD:01993; MIM:235200	Interactions
Gene type		General gene
RefSeq status	REVIEWED	information
Organism	Homo sapiens	General protein
Lineage	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini; Catarrhini; Hominidae; Homo	Reference Sequences Related Sequences
Also known as	HH; HFE1; HLA-H; MGC103790; dJ221C16.10.1	Additional Links
Summary	The protein encoded by this gene is a membrane protein that is similar to MHC class I-type proteins and associates with beta2-microglobulin (beta2M). It is thought that this protein functions to regulate iron absorption by	▼ Links Order cDNA cione
		internet

3. In the **Summary** section you can find information about the function of the gene's protein product. The HFE protein is thought to have a role in regulating iron transport into cells, and defects in the HFE gene can cause the iron absorption disorder hereditary hemochromatosis. Use information provided in the **Summary** section to answer Question 1 for Activity 3 in the Hereditary Hemochromatosis Worksheet in the back of this workbook.

4. Below the summary section is the **Genomic regions, transcripts and products** section. A graphic model has been created for each transcript where a thin line represents an intron that gets spliced out, and the thicker red and blue blocks represent exons. Here we see that the HFE gene has more than one mRNA transcript. For example, an exon included in one transcript might be left out in another transcript. The **Genomic context** section shows where the HFE gene is located within a portion of the chromosome 6 DNA sequence.



5. Select the **Related Sequences** link in the Table of Contents on the right side of the screen to access sequence information for the HFE gene.

Entrez Gene: HFE hemochromato:	sis [Homo sapiens] - Microsoft Internet Explorer	
File Edit View Favorites Tools Help		
odres ei nttp://www.ncbunm.nn.gov/ste	Secure Preseq	V [2] G
S NCBI	Entrez Gene	My NCBI El [Sign.In] (Register)
All Databases	PubMed Nucleotide Protein Genome Structure PMC Taxonomy	Books OMIM
Search Gene	IOP HPEIsymi AND Humanlorghi	
Limits Preview/In	dex History Clipboard Details	
Display Full Report	▼ Show 20 ▼ Sort by Relevance ▼ Send to ▼	
All: 1 Current On	ly: 1 Genes Genomes: 1 SNP GeneView: 1 😠	
1: HEE hemochrom	atosis [Homo sapiens]	Catrez Gran Home
GeneID: 3077	updated 13-Jun-2008	Table Of
Summary	*	2 Contents
Official Symbol	HFE provided by HGNC	Summary Genomic regions,
Official Full Name	hemochromatosis provided by HGNC	Genomic context
Primary source	HGNC:4886	HIV-1 protein
See related	Ensembl:ENSG00000010704; HPRD:01993; MIM:235200	interactions
Gene type	protein coding	Interactions
RefSeq status	REVIEWED	information
Organism	Homo sapiens	General protein
Lineage	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini; Catarrhini; Hominidae; Homo	information Reference Sequences Related Sequences
Also known as	HH; HFE1; HLA-H; MGC103790; dJ221C16.10.1	Additional Links
Summary	The protein encoded by this gene is a membrane protein that is similar to MHC class 1-type proteins and associates with beta2-microglobulin (beta2M).	Links
http://www.ncbi.nim.nih.gov/portal/query	Jcg?db=gene	💣 Internet

🗿 Entrez Gene: HFE hemochro	omatosis [Homo sap	iens] - Microsoft Internet	Explorer		
File Edit View Favorites Tools	Help				**
Address 💩 http://www.ncbi.nlm.nih.	gov/sites/entrez#relseq				🖌 🄁 Go
Related Sequer	nces			(2)	~
Nucleotid	e	Protein			
Genomic	AF184234.1	AAF01222.1			
Genomic	AF204869.1	None			
Genomic	AF331065.1	AAK16502.1			
Genomic	AF525359.1	AAM82608.1			
Genomic	AF525499.1	AAM91950.1			
Genomic	CH471087.1	EAW55516.1			
		EAW55517.1			
		EAW55518.1			
		EAW55519.1			
		EAW55520.1			
		EAW55521.1			
		EAW55522.1			
		EAW55523.1			
		EAW55524.1			
		EAW55525.1			
		EAW55526.1			
		EAW55527.1			
Genomic	CS187189.1	CAJ42862.1			
Genomic	<u>U80914.1</u>	AAD00449.1			
Genomic	<u>U91328.1</u>	AAB82083.1			
Genomic	<u>Y09801.1</u>	CAA70934.1			=
Genomic	<u>Z92910.1</u>	CAB07442.1			_
mRNA	AF079407.1	AAC62646.1			
mRNA	AF079408.1	AAC62647.1			
mRNA	AF079409.1	AAC62648.1			
mRNA	<u>AF109385.1</u>	AAD52104.1			~
A Done		0.811		Internet	.t

Related Sequences section of HFE record in Entrez Gene.

6. To find genomic sequence (including both introns and exons) for HFE, in the **Related Sequences** section, select the genomic sequence record <u>Z92910.1</u>. A screenshot of this GenBank record is shown on the following page.

How did you know which genomic sequence to select?

The problem with archival sequence databases like NCBI's GenBank is that they usually have multiple sequence records for the same gene. You may need to open each record individually and browse through definition, sequence annotation, and comments to determine how much of the gene's nucleotide sequence is contained within each record.

For example, the <u>U91328.1</u> record contains the sequence of a genomic segment that not only includes the HFE gene sequence but also sequences for other genes. <u>Y09801.1</u> contains only sequence information for the HFE promoter and the HFE gene's first exon. The genomic nucleotide sequence records beginning with "AF" contain only partial coding sequence (CDS) for the HFE gene. Of the genomic records listed, <u>Z92910.1</u> has the most complete sequence information for the HFE gene.

GenBank Record <u>Z92910.1</u> - The genomic sequence of the human HFE gene.

NCBI Sequence View	ewer v2.0 - Microsoft Internet Explorer	
File Edit View Favorite	tes Tools Help	27
Address a http://www.nd	icbl.nim.nih.gov/entrez/viewer.fcgi?db=Nucleotide&idopt=GenBank&val=1890179	💌 🔂 Go
S NCBI	Nucleotide	My NCBI
PubMed	Nucleotide Protein Genome Structure PMC Taxonomy	OMIM Books
Search Nucleotic	ide 🕐 for 🛛 🖓 Go Clear	
	Limits Preview/Index History Clipboard Details	
Display GenBan	nk 💌 Show 5 💌 Send to 💌 Hide: 🗖 sequence 🗖 all but gene, CDS and m	RNA features
Range: from	begin to end Creverse complemented strand Fe	eatures: + Refresh
□1: Z92910.	Reports Homo sapiens HFE [gi:1890179]	Links
-	2	a describiose
Features	Sequence	
LOCUS DEFINITION ACCESSION VERSION KEYWORDS SOURCE ORGANISM	292910 12146 bp DNA linear PRI 14-NOV-2006 Homo sapiens HFE gene. 292910 292910.1 GI:1890179 haemochromatosis; HFE gene. Homo sapiens (human) <u>Homo sapiens</u> Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Primates; Haplorrhini; Catarrhini; Hominidee; Homo.	
REFERENCE AUTHORS TITLE	<pre>1 (bases 1 to 858) Albig,W., Drabent,B., Burmester,N., Bode,C. and Doenecke,D. The haemochromatosis candidate gene HFE (HLA-H) of man and mouse is located in syntemic regions within the histone gene cluster I call Biochem 69 (2) 117-126 (1999)</pre>	
PUBMED REFERENCE AUTHORS	9548560 2 (bases 1 to 12146) Albig,W.	×
Done		🔮 Internet 🛒

7. Scroll down the sequence record to the **Features** section (shown below). The different features characterized for this gene are explained on the following page.

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FEATURES	Location/Outlines/Lifiers	
SOURCE	1, 12146	
Source	/organism="Nomo caniane"	
	/ mol time-"removie DNA"	
	/db yref="texon:9606"	
	/cbromsoma="6"	
	/man="6n"	
	/map 0p	
	/clone libe TCRE VIC-library"	
dene	1028 10637	
dene.	/dene="HFF"	
exon	10281324	
3223232	/gene="WFF"	
	/number=1	
CDS	join(1249, 1324, 4652, 4915, 5125, 5400, 6494, 6769,	
000	6928 7041,7995 8035)	
	/dene="HFF"	
	/function="iron metabolism"	
	/note="haemochromatosis candidate gene"	
	/codon start=1	
	/protein id="CAB07442.1"	
	/db xref="GI:1890180"	
	/db_xref="GDB:119309"	
	/db_xref="GOA:030201"	
	/db_xref="HGNC:4886"	
	/db xref="InterPro:IPR001039"	
	/db xref="InterPro:IPR003006"	
	/db_xref="InterPro: IPR003597"	
	/db xref="InterPro: IPR007110"	
	/db xref="InterPro: IPR013783"	
	/db_xref="PDB:1A6Z"	
	/db_xref="PDB:1C42"	
	/db xref="PDB: 1DE4"	
	/db xref="UniProtKB/Swiss-Prot:030201"	
	/translation="MGPRARPALLLLMLLOTAVLOGRLLRSHSLHYLFMGASEODLGL	
	SLFEALGYVDDOLFVFYDHESRRVEPRTPWVSSRISSOMWLOLSOSLKGWDHMFTVDF	
c		-

Gene Gateway: A Web Companion to the Human Genome Landmarks Poster http://genomics.energy.gov/genegateway/

Some features of the sequence in GenBank Record Z92910.1 include

source - The source feature must be included in each sequence record. The source provides the entire sequence length and the scientific name of the source organism. Other types of information in this feature may include chromosome number, map location, and clone or strain identification.

<u>gene</u> - Gives nucleotide numbers where the gene stops and starts. This link opens a new sequence record that shows only the gene sequence.

exon - Gives nucleotide numbers where each exon begins and ends. You will see several of these entries as you scroll down. Each exon is a sequence segment that codes for a portion of processed (intronfree) mRNA. The name of the gene to which the exon belongs and the exon number are provided. An "exon" link opens a new sequence record that shows only the exon sequence.

<u>CDS</u> - The coding sequence (CDS) consists of nucleotides that actually code for amino acids of the protein product. This feature includes the coding sequence's amino acid translation and may also contain gene name, gene product function, a link to protein sequence record, and cross-references to other database entries. A "CDS" link opens a new sequence record that shows only the coding sequence.

intron - Gives nucleotide numbers where each intron begins and ends. An intron is a segment of noncoding sequence that is transcribed but removed from the transcript by splicing together the exons (coding portions) on either side of it. **An "intron" link opens a new sequence record that shows only the intron sequence.**

What's the difference between exons and coding sequence?

Exons often are described as short segments of protein coding sequence. This is a bit of an oversimplification. Exons are segments of sequence spliced together after introns have been removed from pre-mRNA. Exons carry the coding sequence of a gene, but some exons may contain no coding sequence. Portions of exons or even entire exons may contain sequence that is not translated into amino acids. These are the untranslated regions (UTR) of mRNA. UTRs are found upstream and downstream of the proteincoding sequence. See diagram below.



8. Examine the reference section, features section, and sequence at the bottom of this record, and then answer questions 2–4 of the Questions for Activity 3 in the Hereditary Hemochromatosis Worksheet in the back of this workbook.

<u>Activity 4</u> Online Resource: Swiss-Prot

- Access the amino acid sequence of a gene's protein product.

This activity covers how to use the Swiss-Prot protein sequence database to learn about the amino acid sequence and other features of the hereditary hemochromatosis protein.

The protein sequence database Swiss-Prot was developed by groups at the Swiss Institute of Bioinformatics (SIB) and the European Bioinformatics Institute (EBI). Swiss-Prot is noted for its detailed annotation (descriptions of protein function and labeling of domains and other key features within proteins) of protein sequence data. TrEMBL is a computer-annotated database companion to Swiss-Prot that holds sequence data until it can be manually annotated, reviewed, and added to Swiss-Prot.

Let's start by going to the Swiss-Prot home page.

http://us.expasy.org/sprot/

ExPASy - UniProt Knowledgebase:	Swiss-Prot and TrEMBL - Microsoft Internet Explorer	
File Edit View Favorites Tools Help		4
Address Dhttp://us.expasy.org/sprot/		🛩 🛃 Go
	Search Swiss-Prot/TrEMBL r for	Go Clear
swissprot	Swiss-Prot Protein knowledgebase TrEMBL Computer-annotated supplement to S	Valk
The UniProt Knowled • UniProtKB/Swis (such as the des variants, etc.), a References / Lin • UniProtKB/TrEl nucleotide seque	dgebase consists of: ss-Prot; a curated protein sequence database cription of the function of a protein, its domair minimal level of redundancy and high level of king to Swiss-Prot / User manual / Recent cha MBL; a computer-annotated supplement of Sv ence entries not yet integrated in Swiss-Prot.	e which strives to provide a high level of annotation is structure, post-translational modifications, integration with other databases [More details / anges / Disclaimer]. viss-Prot that contains all the translations of EMBL
These databases are	e developed by the Swiss-Prot groups at SIB a	and at EBI.
UniProt Knowledge UniProtKB/Swiss-F entries (More statis UniProtKB/TrEMBL entries (More statis	ebase Release 13.5 consists of: Prot Release 55.5 of 10-Jun-2008: 389046 stics) - Release 38.5 of 10-Jun-2008: 5906286 stics)	> Swiss-Prot headlines Over 100 cross-references in UniProtKB/Swiss- Prot (Read more)
	Access to the UniProt Kno	wledgebase
 SRS - Access to System Full text search i Advanced search 	o UniProtKB/Swiss-Prot, UniProtKB/TrEMBL at in the UniProt Knowledgebase h in the UniProt Knowledgebase by descriptio	nd other databases using the Sequence Retrieval
a		🌍 Internet

1. Scroll down to **Access to UniProt Knowledgebase** section and select <u>Advanced search in</u> <u>the UniProt Knowledgebase</u>. A screenshot of the advanced search page is shown on the next page.

Vew Favorites Tools Help				
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http://us.expasy.org/sprot/sprot-search.html				
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	Search Swiss-Prot/TrEMBI	for	Go Clear	
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t" (this is not possible in Si	RS). Example queries:			
To retrieve all AP1 comple	ex proteins from mouse (AP1	S1, AP1G1, etc. but n	ot MIAP1, IQGAP1,),	specify Gene
Name: ap1*, Organism: N	lus, and deselect "Append a	nd prefix * to query ter	ms".	
To retrieve the three hum	an beta-adrenergic receptor	proteins in UniProtKB/	Swiss-Prot, but not the b	eta-adrenergic
receptor kinases specify	Description: beta&adrenergi	c&recentor/kinase_On	anism. Homo saniens	and select "Append
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URL for Swiss-Prot/TrEMBL Advanced Search: http://us.expasy.org/sprot/sprot-search.html

2. Scroll down to the search boxes. Remove the check in the box next to **UniProtKB/TrEMBL**. We want only sequences from Swiss-Prot. In the **Gene name** search box enter **HFE**. In the **Organism** box enter **human**. To make sure that only one record for the gene with the exact symbol "HFE" is retrieved, deselect **Append and prefix * to query terms**. The advanced search page should resemble the screenshot above. Submit your query.

3. You should retrieve one result. Select the AC number <u>Q30201</u> for the HFE_HUMAN entry to open the record for the HFE protein.

	sGNc=AND&GN=HFE&OC=human&view=ful8z	sum=100		
📥 ExPASy Home page	Site Map	Search ExPASy	Contact us	Swiss-Prot
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Swiss-Prot record for the human HFE protein.

niProtKB/Swiss-Prot entry Q30201 [HFE_HUMAN] Her	editary hemochromatosis protein - Microsoft Internet Explorer				
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lote: most headings are clickable, even	i f they don't appear as links. They link to the user manual or other docu	iments.			
ntry name	HFE HUMAN				
Primary accession number	Q30201				
secondary accession numbers	075929 075930 075931 Q17RT0 Q96KU5 Q96KU7 Q96K Q9HC70 Q9HC83	<u8 q9hc64="" q9hc68<="" td=""></u8>			
ntegrated into Swiss-Prot on	November 1, 1997				
equence was last modified on	November 1, 1997 (Sequence version 1)				
nnotations were last modified on	June 10, 2008 (Entry version 98)				
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ame and origin of the protein					
Protein name	Hereditary hemochromatosis protein [Precursor]				
Protein name Synonym	Hereditary hemochromatosis protein [Precursor] HLA-H				
Protein name Synonym Gene name	Hereditary hemochromatosis protein [Precursor] HLA-H Name: HFE				
vontein name Synonym Gene name	Hereditary hemochromatosis protein [Precursor] HLA-H Name: HFE Synonyms: HLAH				
Protein name Synonym Sene name	Hereditary hemochromatosis protein [Precursor] HLA-H Name: HFE Synonyms: HLAH Homo sapiens (Human) [TaxID: 9606]				

4. Look at the **Protein Name** field. Notice that this protein is designated as a precursor protein. This means that part of the protein chain needs to be cut off by a proteolytic enzyme to form the "mature" functional protein.

5. Using navigation links at the top of the record, go to the **Features** section. The **Features** section of the HFE protein record is shown below.

UniProtKB/S	wiss-Pro	t entry	y Q30201	[HFE_HUMAN] Hereditary hemochromatosis protein - Microsoft Internet Explorer		٦
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Features						Г
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SIGNAL	1	22	22			
CHAIN	23	348	326	Hereditary hemochromatosis protein.	PR0_0000018892	
TOPO_DOM	23	306	284	Extracellular (Potential).		
TRANSMEN	307	330	24	Potential.		
TOPO_DOM	331	348	18	Cytoplasmic (Potential).		
DOMAIN	207	298	92	Ig-like C1-type.		
REGION	23	114	92	Alpha-1.		
REGION	115	205	91	Alpha-2.		
REGION	206	297	92	Alpha-3.		
REGION	298	306	9	Connecting peptide.		
CARBOHYD	110	110		N-linked (GlcNAc) (Potential).		
CARBOHYD	130	130		N-linked (GlcNAc) (Potential).		
CARBOHYD	234	234		N-linked (GlcNAc) (Potential).		
DISULFID	124	187				
DISULFID	225	282				
VAR_SEQ	2.6	114		RSHSLHYLFMGASEQDLGLSLFEALGYVDDQLFVFYDHES	VSP_003218	
				RRVEPRTPWVSSRISSQMWLQLSQSLKGWDHMFTVDFWTI MENHNHSKE -> Q (in isoform 2 and isoform 4).		
VAR_SEQ	26	49		RSHSLHYLFMGASEQDLGLSLFEA -> P (in isoform 5).	VSP_003219	
VAR_SEQ	27	206		Missing (in isoform 6).	VSP_003220	
					Internet	Ì

6. Select the <u>Feature aligner</u> link. This will open a new screen with a list of selected features within the HFE protein. See the screenshot below.

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				GTLV	KQFHDAKEFE	PRDVLPNGD	, IIGG@IIFY	FFGELQKIIC	QVENFGLDQF	LIVIWEPSP:	,
	TRANSMEM	307-330	24	(Potential)							
				IGVISGIAVF	VVILFIGILF	IILR					
	TOPO_DOM	<u>331-348</u>	18	Cytoplasmic	(Potential)						
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7. Notice that the protein chain includes only amino acids 23–348. The first 22 amino acids are not associated with any domains (functional units within a protein). This portion of protein sequence is cleaved from the larger precursor sequence to make the mature, functional HFE protein.

8. Swiss-Prot records are known for their detailed sequence annotation. Notice how each domain is broken down into segments of corresponding amino acids within the protein chain. Select the 23-348 position link to access a new page showing this portion within the entire protein sequence (see screenshot on the next page).

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121	ILG	CEMQI	EDN	STEG	YWKY	GY DO	GQDH.	LEFCI	? DT.	LDWR	AAEP	RAWI	PTKLI	SWE	RHKIR	ARQNR	180	
181	AYLI	ERDCI	PAQ	LQQL.	LELGI	RG V.	LDQQ	VPPL	7 KV	I.HHA.	TSSV	TTL	RCRAI	JNY	YPQNI	TMKWL	240	
241	KDK	2PMD/	AKE .	FEPK	DVLPI	IG D	GTYQ	GWITI	AV	PPGE	EQRY	TCQ	/EHP(GLD	QPLIV	IWEPS	300	
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16	Lou	Dho	Glu	Ala	Lou	Cly	Tur	Val	Aen	Aen	Cln	Leu	Dho	Val	Dhe	40		
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01	τY.	usb	1113	Gru	Det	Arg	Arg	val	Gru	FIO	Arg	1111	210	TTD	var	15		

9. The selected section of the protein sequence is highlighted in red. Another nice feature is the representation of protein sequence using both one-letter and three-letter amino acid abbreviations.

10. Select the <u>Q30201</u> link at the top of the page to return to the main Swiss-Prot record for the HFE protein.

11. Return to the Features section of the record. Scroll down to the part that describes the amino acid position of the protein's secondary structures (e.g., STRAND, TURN, HELIX). You can use this information to figure out which segments of protein sequence form beta-strands, alpha helices, or the turns between these units of secondary structure.

12. In addition to detailed protein sequence annotation available from the Features section, other useful sections are **Comments** and **Cross-references**. The Comments section will provide brief descriptions of protein function, tissues in which the protein is expressed, and associated disease phenotypes. The Cross-references section links to related records found in many different bioinformatics resources. If a protein has structural information deposited in the Protein Data Bank, it will be noted in the Cross-references section.

13. The sequence and feature information presented in this record will help you gain a better understanding of the protein structure examined in Activity 5. Continue with Activity 5 before answering the questions for activities 4 and 5 in the worksheet in the back of this workbook.

Gene Gateway: A Web Companion to the Human Genome Landmarks Poster http://genomics.energy.gov/genegateway/

<u>Activity 5</u> Online Resources: Protein Data Bank and Protein Workshop

- Explore the sequence and structure of the gene's protein product.

This activity demonstrates how to find and view a protein structure using tools and resources available from the Protein Data Bank (PDB). PDB is an international archive of 3-D structural information for biological macromolecules. PDB's structure records provide access to several interactive molecular graphics program. This activity uses Protein Workshop, a tool for viewing and generating high-quality images of molecular structures available from PDB.

Before You Begin

Many features of the PDB Web site require newer Web browsers with JavaScript and cookies enabled, and pop-ups should not be blocked. Internet Explorer 6 was used to create this activity. For more information on system requirements see PDB Frequently Asked Questions (<u>http://www.rcsb.org/pdb/static.do?p=home/faq.html</u>).

Some Protein Structure Basics

- Proteins are created by linking amino acids in a linear fashion to form polypeptide chains. The amino acid sequence of a polypeptide chain is the **primary structure** of a protein. See the Table of Standard Genetic Code in the back of this workbook for single-letter and three-letter abbreviations for the 20 different amino acids.
- Amino acids have different chemical properties. For example, some amino acid residues are strictly hydrophobic ("water fearing") and must be protected from aqueous environments, while other amino acids are hydrophilic ("water loving"). The substitution of just one amino acid for another with very different chemical properties can have serious consequences for a protein's structure and function.
- The folding of regions within the polypeptide chain into alpha helices and beta sheets is a protein's **secondary structure**.
- The packing of the entire polypeptide chain into a three-dimensional globular unit is a protein's **tertiary structure**.
- If a protein molecule is a complex of more than one polypeptide chain, then the complete structure of this molecule is called a protein's **quaternary structure**.
- A domain is a discrete portion of a protein with its own function and specific threedimensional structure. The combination of domains in a single protein determines its overall function.
- Different parts of a polypeptide chain can be linked by disulfide bridges that form between two cysteine residues. Disulfide bridges (or disulfide bonds) stabilize a protein's three-dimensional structure. The loss of a disulfide bridge would be detrimental to a protein's overall structure.

Finding a Structure Record in PDB

To begin, we need to access the Protein Data Bank (http://www.rcsb.org/pdb/).



Note: If you are new to PDB, be sure to check out **General Education** in the light blue column on the left of the screen. Under Educational Resources you can find

- General educational resources introducing molecular structure basics
- Molecule of the Month (a collection of vignettes, each featuring a different molecular structure and its importance to human welfare)
- Education Corner (learn how different educators are using PDB in the classroom)
- PDB newsletters
- Tutorials and other resources.

1. Beside the search box at the top of the PDB home page, select Advanced Search.

2. On the Advanced Search page, from the drop box **Choose a Query Type** select **Swiss-Prot ID(s)**. In Activity 4 we accessed the human hemochromatosis protein record Q30201 in Swiss-Prot. Enter **Q30201** in the search box. The advanced search page should look like the screenshot below. Select the **Evaluate Subquery** button to submit your search.

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Address 🕘 http://www.rcsb.org/pdb/search/advSearch.do		🛩 🛃 Go
PROTEIN DATA BANK	A MEMBER OF THE THE PDB An Information Portal to Biological Macromolecular Structures As of Tuesday Jun 24, 2008 Sthere are 51491 Structures @ PDB Statistics @	^
CONTACT US HELP PRINT PAGE	PDB ID or keyword Author Site Search Author	
Home Search	Are you missing data updates? The PDB archive has moved to ftp://ftp.wwpdb.org. For more information click here.	
Search Database	Match all 🔽 of the following conditions: Advanced Search Tutorial (Requires Flash)	
Sequence Ligands Models	Choose a Query Type:	
Unreleased Entries Structural Genomics Tarnete How to Search	Clear All Evaluate Query	
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3. The search should return two structures. Click on the search result to open a summary of the structure's PDB record.

Swige Bret ID(a)	0
Swiss-Prot ID(s) Swiss-Prot IDs Q30201 2 Structures Evaluate Subquery	-

4. A brief summary of each search result is displayed. The PDB ID for the HFE structure we want to open is 1A6Z. Click on **1A6Z** or the title **HFE (HUMAN) HEMOCHROMATOSIS PROTEIN** (highlighted in the screenshot below) to open the complete PDB record.



5. The complete record is shown on the following page. Note the **Molecular Description** near the bottom of the screenshot. This structure is a complex of four polypeptide chains: A, B, C, and D. A and C are identical HFE polypeptide chains, and B and D are identical chains of another protein called beta-2-microglobulin.

6. Note the primary citation in the 1A6Z record. The best way to learn about structure details is to access the article listed as the primary citation. Although the full text for some articles may be freely available online, many articles are accessible only by subscription. Some university research libraries may provide public access to their journal collections. The article for this structure has been accessed to reveal the following details:

- Only the soluble portion of the HFE polypeptide chain is included in the 1A6Z structure. The transmembrane domain is missing, so the HFE protein in this structure has only 275 of the 348 amino acids in the complete HFE protein sequence.
- The first 22 amino acids of the HFE polypeptide sequence have been excluded because they are not part of the mature, functional protein. Therefore, the first amino acid in this structure is really the 23rd, and cysteine 260 is the cysteine residue involved in the CYS282TYR mutation that we learned about in Activity 1.
- Each HFE polypeptide chain is complexed with another polypeptide chain called beta-2 microglobulin.
- The 1A6Z structure consists of two HFE-beta-2 microglobulin complexes.

7. Select the **Sequence Details** tab (highlighted in screenshot below) to examine the sequence and secondary structure details for this structure.

RCSB PDB : Structure Explorer - Mic	rosoft Internet Explorer					
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Structure Analysis Help	Primary Citation	Lebron, J.A., Ben J.N., Bjorkman, P its interaction with tran [Abstract]	mett, M.J., Vaughn, P.J. (1998) Crystal stru nsferrin receptor. Cell :	D.E., Chirino, A.J., S clure of the hemochroma 93: 111-123	Snow, P.M., Mintier, G.A., Feder, osis protein HFE and characterization of	6 MC
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of this structure click on the Sequence Details tab above the summary page.	Experimental Method	Type X-RAY DIFF	RACTION Data N/A			KING Jmol
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8. The Sequence Details for record 1A6Z are shown on the following page. HFE sequence information is presented first. Each letter in the protein sequence represents a different amino acid. C stands for cysteine. See the Table of Standard Genetic Code in the back of this workbook to determine which amino acid is represented by each letter.

9. Secondary structure details are mapped onto sequence details. Different graphical symbols are used to represent extended beta strands, helixes, and turns. Cysteines that form disulfide bonds are highlighted in yellow and connected by green dotted lines.

HFE Sequence Details in PDB Structure 1A6Z

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	FASTA Sequence	Description	HEE	1, III III III III III III III III III I
	Download Original Files	Identical chains	C	
	Display Files	[show all chains] Chain Type	nolvpentide(L)	
	Display Molecule	UniProt reference	Q30201	
	Structural Reports	Longth	[show this sequence below]	
	External Links	Length	275 residues	
	Structure Analysis	[hide] [reference]	d1a6za1 Hemochromatosis protein Hie, alpha-3 domain. 94 residues 🗠	
	Help	DSSP secondary structure [hide] [reference]	25% helical (8 helices; 70 residues) 39% beta sheet (20 strands; 109 residues)	
		More annotations		11
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	click here	SCOP	d1a6za2	
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		PDB RLLRSHSLHYLFN	4GASEQDLGLSLFEALGYVDDQLFVFYDHESRRVEPRTPWVSSRISSQ	
		PDB 4 10	20 30 40 50 60	
		5540		
		PDB MWLQLSQSLKGWE	2HMFTVDFWTIMENHNHSKESHTLQVILGCEMQEDNSTEGYWKYGYDG 80 90 100 110 120	
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		DSSP	→-///////////////	
		PDB QDHLEFCPDTLDW	VRAAEPRAWPTKLEWERHKIRARQNRAYLERDCPAQLQQLLELGRGVL	
		PDB 121 130	140 150 160 170 180	
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		PDB DQQVPPLVKVTHF PDB 181 190	200 Z10 Z20 Z30 Z40	
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10. By showing the UniProt reference sequence, (see screenshot below) we can see how the sequence of the PDB structure lines up with the Swiss-Prot protein sequence we just examined in Activity 4 (UniProt is another name for Swiss-Prot). Find cysteine 282 in the UniProt sequence. Cysteine 282 is the amino acid that is replaced by tyrosine in the CYS282TYR mutation. You will see that cysteine 282 is at position 260 in the PDB structure. Cysteine 260 forms a disulfide bond with cysteine 203. Disulfide bonds are critical to forming the proper structural arrangement needed to make a functional protein; therefore, the loss of cysteine 260 would be detrimental to protein structure. **Answer the first two questions for Activities 4 and 5 in the worksheet in the back of this workbook.**

it View Favorites Tools Help			
Display Files Display Molecule Structural Reports	Identical chains [show all chains] Chain Type UniProt reference	C polypeptide(L) Q30201 [show PDB sequence below]	
External Links Structure Analysis Help	Length SCOP domain assignment [hide] [reference] DSSP secondary structure [hide] [reference] More annotations	275 residues d1a8za1 Hemochromatosis protein Hfe, alpha-3 domain: 94 residues d1a8za2 Hemochromatosis protein Hfe, alpha-1 and alpha-2 domains: 178 residues 25% helical (8 helices; 70 residues) 39% beta sheet (20 strands; 109 residues)	a 1
Quick Tips : •• X Having trouble with the web site? Try the tutorial: click here	Select	dla5za2	
	DSSP UmProt PDB 61 70 SCOP DSSP UmProt PDB 443 150 1 UmProt PDB 121 130	20 110 120 130 140 VDFWTIMENHNHSKESHTLQVILGČEMQEDNSTEGYWKYGYDG 100 120 80 90 100 120 0 0 100 110 100 120 0 100 120 0 100 120 0 170 180 190 200 PRAMPTKLEWERHKIRARQNRAYLERDČPAQLQQLLELGROVI 140 150 160 170 180	
	SCOP DSSP UmiProt PDB 181 190 SCOP dla	dla6zal 	
	UMProt 263 270 2 UMProt TYQGWITLAVPGEEQR PDB 241 250	80 290 297 YTCOVELDOPLIVIW 260 270 275	

Viewing the Structure

11. Select the **Structure Summary** tab near the top of the Sequence Details page to return to the record summary. At the summary page select **MBT Protein Workshop** from display options in the **Images and Visualization** box (see screenshot below). If you are prompted to download a file, select "Open" to download the file.



12. A Protein Workshop window containing structure 1A6Z should open. You may want to maximize the window so that it fills your computer screen. If you have trouble opening this application, go to the Protein Workshop Help file available from PDB http://www.pdb.org/robohelp f/index.html#viewers/proteinworkshop.htm.



Gene Gateway: A Web Companion to the Human Genome Landmarks Poster http://genomics.energy.gov/genegateway/

13. Some basics for PC users interacting with the structure:

- Click and drag left mouse button to rotate the structure.
- Press Shift + click and drag left mouse button to zoom in and out.
- Click and drag right mouse button to move the structure.

14. At the top of the control panel, you should see four tabs: Tools, Shortcuts, Options, and Help and Credits. If you need to reset the structure to its original configuration at any time during this activity, select the **Options** tab and click **Reset**.



15. Let's explore options in the **Tools** control panel. Using **Tools** involves a four-step process: 1) select your tool; 2) choose what you want the tool to affect (**Atoms and Bonds** selected by default); 3) change the tool's options; and 4) select structure portion you want to modify by clicking in the structure tree at the bottom of the control panel or by clicking on the structure.

16. Chains A, B, C, and D should be displayed. Earlier in the activity we learned that A and C are identical HFE chains and chains B and D are beta-2-microglobulin. Let's use the color and visibility tools to modify the display so that only HFE chain A is visible.

17. First let's color Chain A blue so that we can distinguish it from other chains. The **Colors** tool should be selected. In step 2, choose to modify **Ribbons**. In step 3, click in the **Active Color** box to pick a dark shade of blue from the color palette. Click **OK** to close the **Color** window that pops up. Then select Chain A from the structure tree at the bottom of the control panel (see screenshot to right). Use your mouse to zoom and adjust the position of your structure (see step 13). Your structure should look something like the image below.



Tools Shortcuts Op	tions Help and Credits						
1) Select your tool.							
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2) Choose what you Atoms and Bond	2) Choose what you want the tool to affect. Atoms and Bonds Ribbons						
-3) Change the tool's Active Color: Sample	options, if necessary.						
4) Choose items from	n the tree or 3d viewer.						
4) Choose items from the tree or 3d viewer.							

18. Select the **Visibility** tool. Make sure tool options are set to change visibility of the structure's **Ribbons**. Then select Chain B from the structure tree at the bottom of the control panel (see screenshot to right). Repeat for chains C and D.

19. Select the **Shortcuts** tab. Under **Recolor the backbone by**, select **Conformation type** and click the **Enact** button to color the protein's secondary structure (e.g., helixes are green, beta strands are purple). Chain A should look something like the structure below. Note that another shortcut can be used to change the display area's background.



20. Return to the Tools tab. Let's recolor cysteine 260 and cysteine 203, two residues that form a disulfide bond connecting two different portions of the HFE polypeptide chain. To change the color of the cysteine residues, select the **Colors** tool, choose Ribbons, and pick red from the active color palette. In the tree, expand Chain A and scroll until you can select Cys 260 (selecting the plus sign in front of a chain in the tree will drop a list of all amino acid residues in the chain). See panel to the right. You may need to rotate your structure to locate the red cysteine 260. Repeat for cysteine 203 using dark blue or another color besides red. Rotate the structure to examine the positions of these residues within the chain. The structure should resemble the image on the following page (another graphics package was used to add readable labels to the cysteine residues). Although disulfide bonds are not displayed in this structure, you can see that a bond between cysteines 203 and 260 would keep two different strands parallel to one another within the protein.

Tools Shortcuts Opti	ions Help and Credits						
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H 260	CYS
H 🖌 261	GLN
🗉 🐱 262	VAL
🗉 🐱 263	GLU
E 10 264	HIS
■ 1/2 265	PRO
🖽 😾 26E	GLY
E 267	LEU
🗉 📈 268	ASP
■ 269	GLN 💷
🔳 🐱 270	IPRO V

21. Select the **Options** tab at the top of the control panel. Your structure can be saved as a graphic file using the **Save Image** option. If you click the **Advanced Image Editor** button, a **PDB Image Workbench** window will open. Using the menu options at the top of this window, you can edit the structure and add labels, text, arrows, and other features to your structure and save it as a graphic file (e.g., PNG, TIFF, or JPEG files)



Protein Structure and Hereditary Hemochromatosis Development

By examining the HFE protein's sequence and structure, we discover that the cysteine lost in the CYS282TYR mutation has an important role in establishing the correct threedimensional HFE structure. In this mutation, a cysteine residue is replaced by another amino acid, tyrosine, and the disulfide bond between two cysteines in the polypeptide chain is lost. This is detrimental to the protein's structure. As a result, the HFE protein can no longer perform its normal function of regulating iron uptake, and cells become overloaded with iron. This buildup of iron in cells, if untreated, can lead to organ damage and other complications.

	Т	С	Α	G
т	TTT Phe (F)	TCT Ser (S)	TAT Tyr (Y)	TGT Cys (C)
	TTC Phe (F)	TCC Ser (S)	TAC	TGC
	TTA Leu (L)	TCA Ser (S)	TAA STOP	TGA STOP
	TTG Leu (L)	TCG Ser (S)	TAG STOP	TGG Trp (W)
С	CTT Leu (L)	CCT Pro (P)	CAT His (H)	CGT Arg (R)
	CTC Leu (L)	CCC Pro (P)	CAC His (H)	CGC Arg (R)
	CTA Leu (L)	CCA Pro (P)	CAA GIn (Q)	CGA Arg (R)
	CTG Leu (L)	CCG Pro (P)	CAG GIn (Q)	CGG Arg (R)
Α	ATT IIe (I)	ACT Thr (T)	AAT Asn (N)	AGT Ser (S)
	ATC IIe (I)	ACC Thr (T)	AAC Asn (N)	AGC Ser (S)
	ATA IIe (I)	ACA Thr (T)	AAA Lys (K)	AGA Arg (R)
	ATG Met (M) START	ACG Thr (T)	AAG Lys (K)	AGG Arg (R)
G	GTT Val (V)	GCT Ala (A)	GAT Asp (D)	GGT Gly (G)
	GTC Val (V)	GCC Ala (A)	GAC Asp (D)	GGC Gly (G)
	GTA Val (V)	GCA Ala (A)	GAA Glu (E)	GGA Gly (G)
	GTG Val (V)	GCG Ala (A)	GAG Glu (E)	GGG Gly (G)

Table of Standard Genetic Code for DNA Sequence

Key to the Table of Standard Genetic Code

Alanine	ALA	Α	Arginine	ARG	R				
Asparagine	ASN	Ν	Aspartic acid	ASP	D				
Cysteine	CYS	С	Glutamic acid	GLU	Е				
Glutamine	GLN	Q	Glycine	GLY	G				
Histidine	HIS	Н	Isoleucine	ILE	L				
Leucine	LEU	L	Lysine	LYS	Κ				
Methionine	MET	М	Phenylalanine	PHE	F				
Proline	PRO	Р	Serine	SER	S				
Threonine	THR	Т	Tryptophan	TRP	W				
Tyrosine	TYR	Υ	Valine	VAL	V				
STOP = Termination Signal - signifies the end of a polypeptide chain									

Hereditary Hemochromatosis Worksheet

This worksheet provides questions to be answered as you complete the activities in the Gene Gateway Workbook.

Questions for Activity 1

1) What are some symptoms of hereditary hemochromatosis? How is it treated?

2) What is the official gene symbol of the hereditary hemochromatosis gene?

3) Which allelic variant (genetic mutation) can cause hereditary hemochromatosis?

Questions for Activity 2

1) On the diagram to the right, mark the general region where the HFE gene can be found on chromosome 6.

2) About how many genes are on chromosome 6?

3) How long is the DNA sequence for chromosome 6?



Questions for Activity 3

1) Using the summary provided in Entrez Gene for HFE, briefly describe the function of the gene's protein product.

Use the GenBank sequence record Z92910.1 to answer questions 2–6.

2) In the Features section of record Z92910.1, select the <u>gene</u> link. How many base pairs (bp) are in the genomic sequence of the HFE gene?

3) Scroll through the Features section of the <u>gene</u> sequence in Z92910.1. How many exons have been identified in this sequence?

4) Return to the main record Z92910.1. Select the <u>CDS</u> link. How many base pairs are in the coding sequence?

Questions for Activities 4 and 5

1) Examine HFE's amino acid sequence from Swiss-Prot (shown below). Find cysteine 282, the amino acid that is replaced by tyrosine in the CYS282TYR mutation. Refer to the Table of Standard Genetic Code for help with the single-letter amino acid abbreviations.

10	20	30	40	50	60
MGPRARPALL	LLMLLQTAVL	QGRLLRSHSL	HYLFMGASEQ	DLGLSLFEAL	GYVDDQLFVF
70	80	90	100	110	120
YDHESRRVEP	RTPWVSSRIS	SQMWLQLSQS	LKGWDHMFTV	DFWTIMENHN	HSKESHTLQV
120	140	150	160	170	100
130	140	120	100	1/0	100
	STEGYWKYGY				RHKIRARONR
	DIEGIMICIGI		DIDDWIGHI		
190	200	210	220	230	240
		1	1	1	1
AYLERDCPAQ	LQQLLELGRĠ	VLDQQVPPLV	KVTHHVTSSV	TTLRCRALNY	YPQNITMKWL
250	260	270	280	290	300
KDKQPMDAKE	FEPKDVLPNG	DGTYQGWITL	AVPPGEEQRY	TCQVEHPGLD	QPLIVIWEPS
310	320	330	340		
FPGILPATGAT	SGIAVFVVIL	ғ төтрғ ттрк	KKQGSKGAMG	HIVLAERE	

2) Compare the amino acid sequence above with the HFE sequence details provided for PDB structure 1A6Z. In question 1, underline the portion of the amino acid sequence included in the PDB structure.

3) Why is the cysteine residue affected in the CYS282TYR mutation important?

Contact Information

This document was produced by the Genome Management Information System at Oak Ridge National Laboratory, Oak Ridge, Tennessee, July 2003. The content was last updated June 24, 2008.

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For more information

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