B.Sc. Biotechnology		2018-2019
		SEMESTER - IV
18BTU403	GENOMICS AND PROTEOMICS	4H - 4C

Instruction Hours / week: L:4 T: 0 P: 0

#### Marks: Internal: 40 External: 60 Total: 100 End Semester Exam: 3 Hours

**Course Objectives:** This course is designed to import advance developments in the field of genome sequencing, genome data base, genome mapping and experimental proteomics to the students of the subjects.

**Course Outcomes:** The student will able to apply their knowledge to analyze genomic and proteomic data in large and as well will have an understanding on the application of various experimental techniques that are being currently used in the field of subject.

## UNIT-I

Introduction to Genomics, Gene and Pseudogenes, Gene structure, DNA sequencing methods – manual and automated: Maxam and Gilbert and Sangers method. Pyrosequencing, Genome Sequencing: Shotgun and Hierarchical (clone contig) methods, Computer tools for sequencing projects: Genome sequence assembly software.

## UNIT-II

Managing and Distributing Genome Data: Web based servers and software for genome analysis: ENSEMBL, VISTA, UCSC Genome Browser, NCBI genome. Selected Model Organisms' Genomes and Databases.

### UNIT-III

Genomic mapping: Genetic markers – VNTR, mini and micro satellites, STS, SNPs, ESTs. Types of genome maps, Mapping techniques – Physical and genetic mapping, Map resources, Practical uses genome maps.

## UNIT-IV

Introduction to protein structure, Chemical properties of proteins. Physical interactions that determine the property of proteins. Short-range interactions, electrostatic forces, van der Waal interactions, hydrogen bonds, Hydrophobic interactions. Determination of sizes -Sedimentation analysis, gel filteration, Native PAGE, SDS-PAGE. Determination of covalent structures – Edman degradation.

## UNIT-V

Introduction to Proteomics, Analysis of proteomes. 2D-PAGE. Sample preparation, solubilization, reduction, resolution. Reproducibility of 2D-PAGE. Mass spectrometry based methods for protein identification. De novo sequencing using mass spectrometric data.

### References

- 1. Devarajan Thangadurai, Jeyabalan Sangeetha, (2015). Genomics and Proteomics: Principles, Technologies, and Applications. CRC Press, Tylor & Francis Group
- 2. Lesk A.M., (2014). Introduction to Bioinformatics, (4th ed.). Oxford University Press, UK.
- 3. Bhat S., (2008). Genomics, Bioscience Publishing, New Delhi,
- 4. Timothy P., (2007). Proteomics, SPRINGER.
- 5. Glick, B.R., & Patten C. L. (2017). *Molecular Biotechnology- Principles and Applications of recombinant DNA*. (5th ed.). Washington: ASM Press.
- 6. Primrose, S.B., & Twyman, R.M. (2013). Principles of Gene Manipulation and Genomics (7th ed.). Wiley-Blackwell.
- 7. Tamarin, R. H. (2017). Principles of Genetics (7th ed.). McGraw Hill Education.
- 8. Glick, B.R., & Patten C. L. (2017). *Molecular Biotechnology- Principles and Applications of Recombinant DNA*. (5th ed.). Washington: ASM Press.



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# LECTURE PLAN DEPARTMENT OF BIOTECHNOLOGY

STAFF NAME SUBJECT NAME SUBJECT CODE SEMESTER CLASS : T. SIVARAMAN : GENOMICS AND PROTEOMICS : 18BTU403 : IV : II B.Sc., (BT)

S. No.	Lecture Duration Period	Topics to be Covered	Support Material/Page No.
		UNIT - I	
1	1	Introduction – Genes and pseudogenes, Gene Structure	T1: 74 - 91
2	1	Gene structures	T1: 74 - 91
3	1	Genome sequencing Maxam and Gilbert Method	T1: 91 - 98
4	1	Chain termination methods	R1: 1 - 36
5	1	Automated DNA Sequencing and Shotgun sequencing	R1: 1 - 36
6	1	Pyrosequencing	R1: 1 - 36
7	1	Hierarchical (clone contig) methods and computational tools	T1: 74 - 123
8	1	Genome sequence assembly software	T1: 74 - 123
	Total No. of I	Hours Planned For Unit I = 08	
		UNIT - II	
1	1	Web based servers and software for genome analysis	T3: 286 - 308
2	1	ENSEMBL	W1
3	1	VISTA	W2
4	1	UCSC Genome Browser	W3
5	1	NCBI genome	W4

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6	1	Selected Model Organisms' Genomes and Databases	T2: 67 - 116
7	1	FlyBase	T2: 67 - 116
8	1	OMIM	T2: 67 - 116
	Total No. of H	Iours Planned For Unit II = 08	
		UNIT – III	
1	1	VNTR, mini and micro satellites	R1: 113 - 129 T3: 286 - 308
2	1	STS, SNP, ESTs.	R1: 113 - 129
3	1	Genome Mapping	R1: 98 - 99
4	1	Physical mapping	R1: 209 - 216
5	1	Genetic mapping	R1:209 - 216
6	1	Map resources	T3: 41 - 188
7	1	Practical uses - genome maps	R2: 143 - 148
8	1	Genome mapping and genetic markers overview	
	Total No. of H	ours Planned For Unit III = 08	
		UNIT - IV	
1	1	Introduction to protein structure	T4: 219
2	1	Chemical properties of proteins	T4: 220 - 230
3	1	Physical interactions and properties	T2: 67 - 116
4	1	Van der Waal interactions, hydrogen bonds, Hydrophobic interactions, electrostatic forces	T4: 258 - 263
5	1	Edman degradation	T4: 264 - 271
6	1	Sedimentation analysis, Gel filtration	W5
7	1	Native PAGE, SDS-PAGE	W6
8	1	Determination of covalent structures	T4: 231 - 257
	Total No. of H	Iours Planned For Unit V = 08	
	1	UNIT - V	
1	1	Introduction to Proteomics	W7

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3	1	Reproducibility of 2D-PAGE	T5: 1 - 34
4	1	EI-MS	T5: 35 - 74
5	1	ESI-MS	T5: 35 - 74
6	1	MALDI-MS	T5: 35 - 74
7	1	De novo sequencing using mass spectrometric data	T5: 35 - 74
8	1	ESE QP discussion	
	Total No. of	Hours Planned For Unit V = 08	
	Total No. of Planned Hours = 40		

## TEXT BOOKS

T1: Genomics and Cloning (East-West Press, 2004) – Kumar, HD.

T2: Introduction to Bioinformatics (Oxford University Press, Second edition, 2007) – Arthur M Lesk.

T3: Bioinformatics concepts, Skills and Applications (CBS Publishers & Distributors, Second edition, 2007) – Rastogi, SC., Namita, M and Parag, R.

T4: Biochemistry (John Wiley & Sons, 2011) – Donald Voet and Judith Voet.

T5: Proteomics (Kluwer Academic Publishers, 2002) – Timothy, P.

### **REFERENCE BOOKS**

R1: Genomics (Bioscience Publishers, 2008) – Bhatt, S.

R2: Principles of Genome analysis (Blackwell publishing, 2003), Primrose, SB & Twyman, RM.

## **WEBSITES**

W1: https://www.ensembl.org/index.html

- W2: http://genome.lbl.gov/vista/index.html
- W3: https://genome.ucsc.edu/
- W4: https://www.ncbi.nlm.nih.gov/
- W5: https://en.wikipedia.org/wiki/Size-exclusion\_chromatography
- W6: https://en.wikipedia.org/wiki/Polyacrylamide\_gel\_electrophoresis
- W7: https://en.wikipedia.org/wiki/Proteomics

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#### UNIT - I

#### **SYLLABUS**

**Introduction to Genomics:** Gene and Pseudogenes, Gene structure, DNA sequencing methods – manual and automated: Maxam and Gilbert and Sanger's method. Pyrosequencing, Genome Sequencing: Shotgun and Hierarchical (clone contig) methods, Computer tools for sequencing projects: Genome sequence assembly software.

#### Genome

Genomics is an interdisciplinary field of science within the field of molecular biology. A genome is a complete set of DNA within a single cell of an organism, and as such, focuses on the structure, function, evolution, and mapping of genomes. Genomics aims at the collective characterization and quantification of genes, which direct the production of proteins with the assistance of enzymes and messenger molecules. Genomics also involves the sequencing and analysis of genomes. Advances in genomics have triggered a revolution in discovery-based research to understand even the most currently complex biological systems such as the brain. In contrast to genetics, which refers to the study of individual genes and their roles in inheritance, genomics uses high throughput DNA sequencing and bioinformatics to assemble, and analyze the function and structure of entire genomes. The field also includes studies of intragenomic (within the genome) phenomena such as heterosis (hybrid vigour), epistasis (effect of one gene on another), pleiotropy (one gene affecting more than one trait) and other interactions between loci and alleles within the genome. Advances in genomics have triggered a revolution in systems biology which facilitates the understanding of complex biological systems such as the brain. From the Greek FEN gen, "gene" (gamma, epsilon, nu, epsilon) meaning "become, create, creation, birth", and subsequent variants: genealogy, genesis, genetics, genic, genomere, genotype, genus etc. While the word genome (from the German Genom, attributed to Hans Winkler) was in use in English as early as 1926, <sup>1</sup>the term *genomics* was coined by Tom Roderick, a geneticist at the Jackson Laboratory (Bar Harbor, Maine), over beer at a meeting held in Maryland on the mapping of the human genome in 1986.

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## **Genes**

A gene is the basic physical and functional unit of heredity.

Genes, which are made up of DNA, act as instructions to make molecules called proteins. In humans, genes vary in size from a few hundred DNA bases to more than 2 million bases. The Human Genome Project has estimated that humans have between 20,000 and 25,000 genes.

Every person has two copies of each gene, one inherited from each parent.

Most genes are the same in all people, but a small number of genes (less than 1 percent of the total) are slightly different between people.

Alleles are forms of the same gene with small differences in their sequence of DNA bases.

These small differences contribute to each person's unique physical features.

Genes are a section of DNA that are in charge of different functions like making proteins. Long strands of DNA with lots of genes make up chromosomes. DNA molecules are found in chromosomes. Chromosomes are located inside of the nucleus of cells.

Each chromosome is one long single molecule of DNA. This DNA contains important genetic information.

Chromosomes have a unique structure, which helps to keep the DNA tightly wrapped around the proteins called histones. If the DNA molecules were not bound by the histones, they would be too long to fit inside of the cell.

Genes vary in complexity. In humans, they range in size from a few hundred DNA bases to more than 2 million bases.

Different living things have different shapes and numbers of chromosomes. Humans have 23 pairs of chromosomes, or a total of 46. A donkey has 31 pairs of chromosomes, a hedgehog has 44, and a fruit fly has just 4.

DNA contains the biological instructions that make each species unique.

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DNA is passed from adult organisms to their offspring during reproduction. The building blocks of DNA are called nucleotides. Nucleotides have three parts: A phosphate group, a sugar group and one of four types of nitrogen bases.

A gene consists of a long combination of four different nucleotide bases, or chemicals. There are many possible combinations.

The four nucleotides are:

- 1. A (adenine)
- 2. C (cytosine)
- 3. G (guanine)
- 4. T (thymine)

Different combinations of the letters ACGT give people different characteristics. For example, a person with the combination ATCGTT may have blue eyes, while somebody with the combination ATCGCT may have brown eyes.

Genes decide almost everything about a living being. One or more genes can affect a specific trait. Genes may interact with an individual's environment too and change what the gene makes.

Genes affect hundreds of internal and external factors, such as whether a person will get a particular color of eyes or what diseases they may develop.

Some diseases, such as sickle-cell anemia and Huntington's disease, are inherited, and these are also affected by genes.

#### Genes consist of three types of nucleotide sequence:

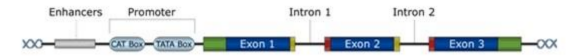
coding regions, called exons, which specify a sequence of amino acids non-coding regions, called introns, which do not specify amino acids

regulatory sequences, which play a role in determining when and where the protein is made (and how much is made)

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A human being has 20,000 to 25,000 genes located on 46 chromosomes (23 pairs). These genes are known, collectively, as the human genome.



The structural components of a gene

# The Human Genome Project (HGP)

The Human Genome Project (HGP) is a major scientific research project. It is the largest single research activity ever carried out in modern science.

It aims to determine the sequence of the chemical pairs that make up human DNA and to identify and map the 20,000 to 25,000 or so genes that make up the human genome.

The HGP has opened the door to a wide range of genetic tests.

The project was started in 1990 by a group of international researchers, the United States' National Institutes of Health (NIH) and the Department of Energy.

The goal was to sequence 3 billion letters, or base pairs, in the human genome, that make up the complete set of DNA in the human body.

By doing this, the scientists hoped to provide researchers with powerful tools, not only to understand the genetic factors in human disease, but also to open the door for new strategies for diagnosis, treatment, and prevention.

The HGP was completed in 2003, and all the data generated is available for free access on the internet. Apart from humans, the HGP also looked at other organisms and animals, such as the fruit fly and E. coli.

Over three billion nucleotide combinations, or combinations of ACGT, have been found in the human genome, or the collection of genetic features that can make up the human body.

Mapping the human genome brings scientists closer to developing effective treatments for hundreds of diseases.

The project has fueled the discovery of more than 1,800 disease genes. This has made it easier

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for researchers to find a gene that is suspected of causing an inherited disease in a matter of days. Before this research was carried out, it could have taken years to find the gene.

### **Pseudogenes**

Pseudogenes are segments of DNA that are related to real genes. Pseudogenes have lost at least some functionality, relative to the complete gene, in cellular gene expression or proteincoding ability. Pseudogenes often result from the accumulation of multiple mutations within a gene whose product is not required for the survival of the organism, but can also be caused by genomic copy number variation (CNV) where segments of 1+ kb are duplicated or deleted. Although not *fully* functional, pseudogenes may be functional, similar to other kinds of noncoding DNA, which can perform regulatory functions. The "pseudo" in "pseudogene" implies a variation in sequence relative to the parent coding gene, but does not necessarily indicate pseudo-function. Despite being non-coding, many pseudogenes have important roles in normal physiology and abnormal pathology. Although some pseudogenes do not have introns or promoters (such pseudogenes are copied from messenger RNA and incorporated into the chromosome, and are called "processed pseudogenes"), others have some gene-like features such as promoters, CpG islands, and splice sites. They are different from normal genes due to either a lack of protein-coding ability resulting from a variety of disabling mutations (e.g. premature stop codons or frameshifts), a lack of transcription, or their inability to encode RNA (such as with ribosomal RNA pseudogenes). The term "pseudogene" was coined in 1977 by Jacq et al.Because pseudogenes were initially thought of as the last stop for genomic material that could be removed from the genome, they were often labeled as junk DNA. Nonetheless, pseudogenes contain biological and evolutionary histories within their sequences. This is due to a pseudogene's shared ancestry with a functional gene: in the same way that Darwin thought of two species as possibly having a shared common ancestry followed by millions of years of evolutionary divergence, a pseudogene and its associated functional gene also share a common ancestor and have diverged as separate genetic entities over millions of years.

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Pseudogenes are usually characterized by a combination of homology to a known gene and loss of some functionality. That is, although every pseudogene has a DNA sequence that is similar to some functional gene, they are usually unable to produce functional final protein products. Pseudogenes are sometimes difficult to identify and characterize in genomes, because the two requirements of homology and loss of functionality are usually implied through sequence alignments rather than biologically proven.

Homology is implied by sequence identity between the DNA sequences of the pseudogene and parent gene. After aligning the two sequences, the percentage of identical base pairs is computed. A high sequence identity means that it is highly likely that these two sequences diverged from a common ancestral sequence (are homologous), and highly unlikely that these two sequences have evolved independently (see Convergent evolution).

Non-functionality can manifest itself in many ways. Normally, a gene must go through several steps to a fully functional protein: Transcription, pre-mRNA processing, translation, and protein folding are all required parts of this process. If any of these steps fails, then the sequence may be considered nonfunctional. In high-throughput pseudogene identification, the most commonly identified disablements are premature stop codons and frameshifts, which almost universally prevent the translation of a functional protein product.

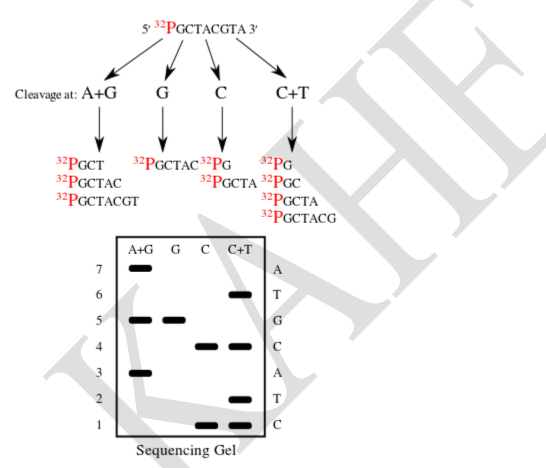
Pseudogenes for RNA genes are usually more difficult to discover as they do not need to be translated and thus do not have "reading frames". Pseudogenes can complicate molecular genetic studies. For example, amplification of a gene by PCR may simultaneously amplify a pseudogene that shares similar sequences. This is known as PCR bias or amplification bias. Similarly, pseudogenes are sometimes annotated as genes in genome sequences. Processed pseudogenes often pose a problem for gene prediction programs, often being misidentified as real genes or exons. It has been proposed that identification of processed pseudogenes can help improve the accuracy of gene prediction methods. Recently 140 human pseudogenes have been shown to be translated. However, the function, if any, of the protein products is unknown.

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## Maxam-Gilbert sequencing

Maxam–Gilbert sequencing is a method of DNA sequencing developed by Allan Maxam and Walter Gilbert in 1976–1977. This method is based on nucleobase-specific partial chemical modification of DNA and subsequent cleavage of the DNA backbone at sites adjacent to the modified nucleotides.



An example Maxam–Gilbert sequencing reaction. Cleaving the same tagged segment of DNA at different points yields tagged fragments of different sizes. The fragments may then be separated by gel electrophoresis. Maxam–Gilbert sequencing was the first widely adopted method for DNA sequencing, and, along with the Sanger dideoxy method, represents the first

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generation of DNA sequencing methods. Maxam–Gilbert sequencing is no longer in widespread use, having been supplanted by next-generation sequencing methods.

Although Maxam and Gilbert published their chemical sequencing method two years after Frederick Sanger and Alan Coulson published their work on plus-minus sequencing, Maxam-Gilbert sequencing rapidly became more popular, since purified DNA could be used directly, while the initial Sanger method required that each read start be cloned for production of single-stranded DNA. However, with the improvement of the chain-termination method (see below), Maxam-Gilbert sequencing has fallen out of favour due to its technical complexity prohibiting its use in standard molecular biology kits, extensive use of hazardous chemicals, and difficulties with scale-up. Maxam-Gilbert sequencing requires radioactive labeling at one 5' end of the DNA fragment to be sequenced (typically by a kinase reaction using gamma-<sup>32</sup>P ATP) and purification of the DNA. Chemical treatment generates breaks at a small proportion of one or two of the four nucleotide bases in each of four reactions (G, A+G, C, C+T). For example, the purines (A+G) are depurinated using formic acid, the guanines (and to some extent the adenines) are methylated by dimethyl sulfate, and the pyrimidines (C+T) are hydrolysed using hydrazine. The addition of salt (sodium chloride) to the hydrazine reaction inhibits the reaction of thymine for the C-only reaction. The modified DNAs may then be cleaved by hot piperidine; (CH<sub>2</sub>)<sub>5</sub>NH at the position of the modified base. The concentration of the modifying chemicals is controlled to introduce on average one modification per DNA molecule. Thus a series of labeled fragments is generated, from the radiolabeled end to the first "cut" site in each molecule. The fragments in the four reactions are electrophoresed side by side in denaturing acrylamide gels for size separation. To visualize the fragments, the gel is exposed to X-ray film for autoradiography, yielding a series of dark bands each showing the location of identical radiolabeled DNA molecules. From presence and absence of certain fragments the sequence may be inferred.

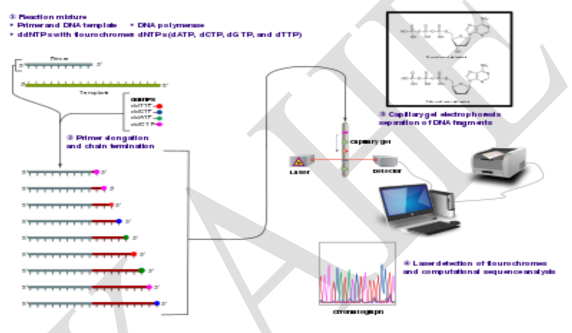
#### Sanger sequencing

Sanger sequencing is a method of DNA sequencing first commercialized by Applied Biosystems, based on the selective incorporation of chain-terminating dideoxynucleotides by

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DNA polymerase during in vitro DNA replication. Developed by Frederick Sanger and colleagues in 1977, it was the most widely used sequencing method for approximately 40 years. More recently, higher volume Sanger sequencing has been supplanted by "Next-Gen" sequencing methods, especially for large-scale, automated genome analyses. However, the Sanger method remains in wide use, for smaller-scale projects, validation of Next-Gen results and for obtaining especially long contiguous DNA sequence reads (> 500 nucleotides).

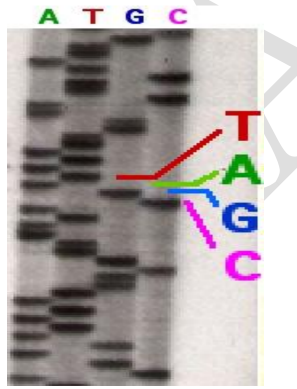


The classical chain-termination method requires a single-stranded DNA template, a DNA primer, a DNA polymerase, normal deoxynucleosidetriphosphates (dNTPs), and modified dideoxynucleotidetriphosphates (ddNTPs), the latter of which terminate DNA strand elongation. These chain-terminating nucleotides lack a 3'-OH group required for the formation of a phosphodiester bond between two nucleotides, causing DNA polymerase to cease extension of DNA when a modified ddNTP is incorporated. The ddNTPs may be radioactively or fluorescently labeled for detection in automated sequencing machines. The DNA sample is divided into four separate sequencing reactions, containing all four of the standard deoxynucleotides (dATP, dGTP, dCTP and dTTP) and the DNA polymerase. To each reaction is

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added only one of the four dideoxynucleotides (ddATP, ddGTP, ddCTP, or ddTTP), while the other added nucleotides are ordinary ones. The dideoxynucleotide is added to be approximately 100-fold lower in concentration than the corresponding deoxynucleotide (e.g. 0.005mM ddATP : 0.5mM dATP) allowing for enough fragments to be produced while still transcribing the complete sequence. Putting it in a more sensible order, four separate reactions are needed in this process to test all four ddNTPs. Following rounds of template DNA extension from the bound primer, the resulting DNA fragments are heat denatured and separated by size using gel electrophoresis. In the original publication of 1977,<sup>[2]</sup> the formation of base-paired loops of ssDNA was a cause of serious difficulty in resolving bands at some locations. This is frequently performed using a denaturing polyacrylamide-urea gel with each of the four reactions run in one of four individual lanes (lanes A, T, G, C). The DNA bands may then be visualized by autoradiography or UV light and the DNA sequence can be directly read off the X-ray film or gel image.



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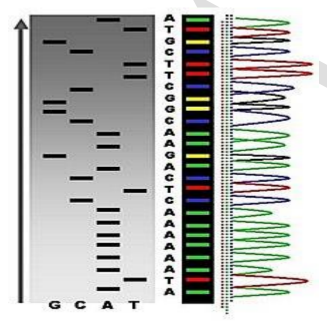
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In the image on the right, X-ray film was exposed to the gel, and the dark bands correspond to DNA fragments of different lengths. A dark band in a lane indicates a DNA fragment that is the result of chain termination after incorporation of a dideoxynucleotide (ddATP, ddGTP, ddCTP, or ddTTP). The relative positions of the different bands among the four lanes, from bottom to top, are then used to read the DNA sequence. DNA fragments are labelled with a radioactive or fluorescent tag on the primer (1), in the new DNA strand with a labeled dNTP, or with a labeled ddNTP.

Technical variations of chain-termination sequencing include tagging with nucleotides containing radioactive phosphorus for radiolabelling, or using a primer labeled at the 5' end with a fluorescent dye. Dye-primer sequencing facilitates reading in an optical system for faster and more economical analysis and automation. The later development by Leroy Hood and coworkers of fluorescently labeled ddNTPs and primers set the stage for automated, high-throughput DNA sequencing.



Sequence ladder by radioactive sequencing compared to fluorescent peaks

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Chain-termination methods have greatly simplified DNA sequencing. For example, chain-termination-based kits are commercially available that contain the reagents needed for sequencing, pre-aliquoted and ready to use. Limitations include non-specific binding of the primer to the DNA, affecting accurate read-out of the DNA sequence, and DNA secondary structures affecting the fidelity of the sequence.

*Dye-terminator sequencing* utilizes labeling of the chain terminator ddNTPs, which permits sequencing in a single reaction, rather than four reactions as in the labelled-primer method. In dye-terminator sequencing, each of the four dideoxynucleotide chain terminators is labelled with fluorescent dyes, each of which emits light at different wavelengths.

Owing to its greater expediency and speed, dye-terminator sequencing is now the mainstay in automated sequencing. Its limitations include dye effects due to differences in the incorporation of the dye-labeled chain terminators into the DNA fragment, resulting in unequal peak heights and shapes in the electronic DNA sequence trace chromatogram after capillary electrophoresis (see figure to the left). This problem has been addressed with the use of modified DNA polymerase enzyme systems and dyes that minimize incorporation variability, as well as methods for eliminating "dye blobs". The dye-terminator sequencing method, along with automated high-throughput DNA sequence analyzers, is now being used for the vast majority of sequencing projects.

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Automated DNA-sequencing instruments (DNA sequencers) can sequence up to 384 DNA samples in a single batch. Batch runs may occur up to 24 times a day. DNA sequencers separate strands by size (or length) using capillary electrophoresis, they detect and record dye fluorescence, and output data as fluorescent peak trace chromatograms. Sequencing reactions (thermocycling and labelling), cleanup and re-suspension of samples in a buffer solution are performed separately, before loading samples onto the sequencer. A number of commercial and

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non-commercial software packages can trim low-quality DNA traces automatically. These programs score the quality of each peak and remove low-quality base peaks (which are generally located at the ends of the sequence). The accuracy of such algorithms is inferior to visual examination by a human operator, but is adequate for automated processing of large sequence data sets. Common challenges of DNA sequencing with the Sanger method include poor quality in the first 15-40 bases of the sequence due to primer binding and deteriorating quality of sequencing traces after 700-900 bases. Base calling software such as Phred typically provides an estimate of quality to aid in trimming of low-quality regions of sequences.

In cases where DNA fragments are cloned before sequencing, the resulting sequence may contain parts of the cloning vector. In contrast, PCR-based cloning and next-generation sequencing technologies based on pyrosequencing often avoid using cloning vectors. Recently, one-step Sanger sequencing (combined amplification and sequencing) methods such as Ampliseq and SeqSharp have been developed that allow rapid sequencing of target genes without cloning or prior amplification. Current methods can directly sequence only relatively short (300-1000 nucleotides long) DNA fragments in a single reaction. The main obstacle to sequencing DNA fragments above this size limit is insufficient power of separation for resolving large DNA fragments that differ in length by only one nucleotide.

### Shotgun sequencing

Shotgun sequencing involves randomly breaking up DNA sequences into lots of small pieces and then reassembling the sequence by looking for regions of overlap.

Large, mammalian genomes are particularly difficult to clone, sequence and assemble because of their size and structural complexity. As a result clone-by-clone sequencing, although reliable and methodical, takes a very long time.

With the emergence of cheaper sequencing and more sophisticated computer programs, researchers have therefore relied on whole genome shotgun sequencing to tackle larger, more complex genomes.

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Shotgun sequencing was originally used by Fred Sanger and his colleagues to sequence small genomes such as those of viruses and bacteria.

Whole genome shotgun sequencing bypasses the time-consuming mapping and cloning steps that make clone-by-clone sequencing so slow.

In whole genome shotgun sequencing the entire genome is broken up into small fragments of DNA for sequencing.

These fragments are often of varying sizes, ranging from 2-20 kilobases (2,000- 20,000 base pairs) to 200-300 kilobases (200,000-300,000 base pairs).

These fragments are sequenced to determine the order of the DNA bases, A, C, G and T. The sequenced fragments are then assembled together by computer programs that find where fragments overlap.

You can imagine shotgun sequencing as being a bit like shredding multiple copies of a book (which in this case is a genome), mixing up all the fragments and then reassembling the original text (genome) by finding fragments with text that overlap and piecing the book back together again.

This method of genome sequencing was used by Craig Venter, founder of the private company Celera Genomics, to sequence the human genome.

Venter wanted to sequence the human genome faster than the publicly funded effort and felt this was the best way. To assemble the sequence Venter used the clone-by- clone publically available data from the Human Genome Project.

Now, as technologies are improving, whole genome shotgun sequencing is being used to improve the accuracy of existing genome sequences, such as the reference human genome.

It is used to remove errors, fill in gaps or correct parts of the sequence that were originally assembled incorrectly when clone-by-clone sequencing was used.

As a consequence the reference human genome is constantly being improved to ensure that the genome sequence is of the highest possible standard.

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By removing the mapping stages, whole genome shotgun sequencing is a much faster process than clone-by-clone sequencing.

Whole genome shotgun sequencing uses a fraction of the DNA that clone-by-clone sequencing needs.

Whole genome shotgun sequencing is particularly efficient if there is an existing reference sequence. It is much easier to assemble the genome sequence by aligning it to an existing reference genome.

Shotgun sequencing is much faster and less expensive than methods requiring a genetic map.

## **Disadvantages of shotgun sequencing**

Vast amounts of computing power and sophisticated software are required to assemble shotgun sequences together. To sequence the genome from a mammal (billions of bases long), you need about 60 million individual DNA sequence reads.

Errors in assembly are more likely to be made because a genetic map is not used. However these errors are generally easier to resolve than in other methods and minimised if a reference genome can be used.

Whole genome shotgun sequencing can only really be carried out if a reference genome is already available, otherwise assembly is very difficult without an existing genome to match it to. Whole genome shotgun sequencing can also lead to errors which need to be resolved by other, more labour-intensive types of sequencing, such as clone-by-clone sequencing. Repetitive genomes and sequences can be more difficult to assemble.

#### Assembly of contiguous DNA sequence

The next question to address is how the master sequence of a chromosome, possibly several tens of Mb in length, can be assembled from the multitude of short sequences generated by chain termination sequencing.

The relatively short genomes of prokaryotes can be assembled by shotgun sequencing, but that this approach might lead to errors if applied to larger eukaryotic genomes.

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The whole-genome shotgun method, which uses a map to aid assembly of the master sequence, has been used with the fruit-fly and human genomes, but it is generally accepted that a greater degree of accuracy is achieved with the clone contig approach, in which the genome is broken down into segments, each with a known position on the genome map, before sequencing is carried out. We will start by examining how shotgun sequencing has been applied to prokaryotic genomes.

Samples of each clone in row A of the first microtiter tray are mixed together and a single PCR carried out. This is repeated for every row of every tray -80 PCRs in all. Samples of each clone in column 1 of the first microtiter tray are mixed together and a single PCR carried out. This is repeated for every column of every tray -120 PCRs in all.

Clones from well A1 of each of the ten microtiter trays are mixed together and a single PCR carried out. This is repeated for every well – 96 PCRs in all.

These 296 PCRs provide enough information to identify which of the 960 clones give products and which do not. Ambiguities arise only if a substantial number of clones turn out to be positive.

#### **Genomic Library**

A genomic library is a collection of plasmid clones or phage lysates containing recombinant DNA molecules so that the sum total of DNA inserts in this collection, ideally, represents the entire genome of the concerned organism. However, inspite of all the care taken in the production of genomic libraries.

Certain DNA fragments should be expected to be under or over represented or even missing. There are several possible reasons for this, and at present they can not be taken care of.

#### **Pyrosequencing**

Pyrosequencing is a method of DNA sequencing (determining the order of nucleotides in DNA) based on the "sequencing by synthesis" principle, in which the sequencing is performed by detecting the nucleotide incorporated by a DNA polymerase. Pyrosequencing relies on light detection based on a chain reaction when pyrophosphate is released.

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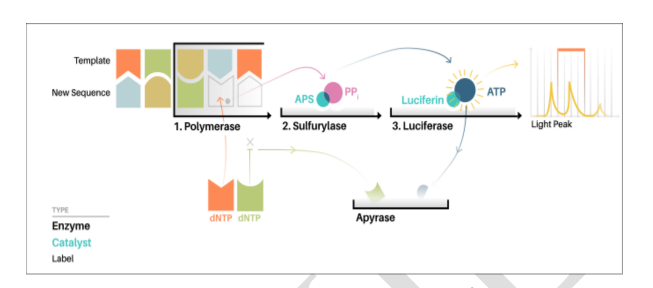
The principle of Pyrosequencing was first described in 1993 by Bertil Pettersson, Mathias Uhlen and Pål Nyren by combining the solid phase sequencing method using streptavidin coated magnetic beads with recombinant DNA polymerase lacking 3'to 5'exonuclease activity (proof-reading) and luminescence detection using the firefly luciferase enzyme. A mixture of three enzymes (DNA polymerase, ATP sulfurylase and firefly luciferase) and a nucleotide (dNTP) are added to single stranded DNA to be sequenced and the incorporation of nucleotide is followed by measuring the light emitted. The intensity of the light determines if 0, 1 or more nucleotides have been incorporated, thus showing how many complementary nucleotides are present on the template strand. The nucleotide mixture is removed before the next nucleotide mixture is added. This process is repeated with each of the four nucleotides until the DNA sequence of the single stranded template is determined.

A second solution-based method for Pyrosequencing was described in 1998 by Mostafa Ronaghi, Mathias Uhlen and Pål Nyren. In this alternative method, an additional enzyme apyrase is introduced to remove nucleotides that are not incorporated by the DNA polymerase. This enabled the enzyme mixture including the DNA polymerase, the luciferase and the apyrase to be added at the start and kept throughout the procedure, thus providing a simple set-up suitable for automation. An automated instrument based on this principle was introduced to the market the following year by the company Pyrosequencing.

A third microfluidic variant of the Pyrosequencing method was described in 2005 by Jonathan Rothberg and co-workers at the company 454 Life Sciences. This alternative approach for Pyrosequencing was based on the original principal of attaching the DNA to be sequenced to a solid support and they showed that sequencing could be performed in a highly parallel manner using a microfabricated microarray. This allowed for high-throughput DNA sequencing and an automated instrument was introduced to the market. This became the first next generation sequencing instrument starting a new era in genomics research, with rapidly falling prices for DNA sequencing allowing whole genome sequencing at affordable prices.

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"Sequencing by synthesis" involves taking a single strand of the DNA to be sequenced and then synthesizing its complementary strand enzymatically. The pyrosequencing method is based on detecting the activity of DNA polymerase (a DNA synthesizing enzyme) with another chemoluminescent enzyme. Essentially, the method allows sequencing of a single strand of DNA by synthesizing the complementary strand along it, one base pair at a time, and detecting which base was actually added at each step. The template DNA is immobile, and solutions of A, C, G, and T nucleotides are sequentially added and removed from the reaction. Light is produced only when the nucleotide solution complements the first unpaired base of the template. The sequence of solutions which produce chemiluminescent signals allows the determination of the sequence of the template.

For the solution-based version of Pyrosequencing, the single-strand DNA (ssDNA) template is hybridized to a sequencing primer and incubated with the enzymes DNA polymerase, ATP sulfurylase, luciferase and apyrase, and with the substrates adenosine 5' phosphosulfate (APS) and luciferin.

 The addition of one of the four deoxynucleotide triphosphates (dNTPs) (dATPαS, which is not a substrate for a luciferase, is added instead of dATP to avoid noise) initiates the second step. DNA polymerase incorporates the correct, complementary dNTPs onto the template. This incorporation releases pyrophosphate (PPi).

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# 2. ATP sulfurylase converts PPi to ATP in the presence of adenosine 5' phosphosulfate. This ATP acts as a substrate for the luciferase-mediated conversion of luciferin to oxyluciferin that generates visible light in amounts that are proportional to the amount of ATP. The light produced in the luciferase-catalyzed reaction is detected by a camera and

analyzed in a pyrogram.

3. Unincorporated nucleotides and ATP are degraded by the apyrase, and the reaction can restart with another nucleotide.

Currently, a limitation of the method is that the lengths of individual reads of DNA sequence are in the neighborhood of 300-500 nucleotides, shorter than the 800-1000 obtainable with chain termination methods (e.g. Sanger sequencing). This can make the process of genome assembly more difficult, particularly for sequences containing a large amount of repetitive DNA.

The company Pyrosequencing AB in Uppsala, Sweden was founded with venture capital provided by HealthCap in order to commercialize machinery and reagents for sequencing short stretches of DNA using the pyrosequencing technique. Pyrosequencing AB was listed on the Stockholm Stock Exchange in 1999. It was renamed to Biotage in 2003. The pyrosequencing business line was acquired by Qiagen in 2008. Pyrosequencing technology was further licensed to 454 Life Sciences. 454 developed an array-based pyrosequencing technology which has emerged as a platform for large-scale DNA sequencing. Most notable are the applications for genome sequencing and metagenomics. Roche announced the discontinuation of the 454 sequencing platform in 2013.

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# **Possible questions**

#### **1 Mark questions** 1. The term 'genomics' was coined by a) Tom Roderick b) Anselme Payen c) Wilhelm Johannsen d) Paulien Hogeweg 2. How many different types of chemcial treatments are required in Maxam-Gilbert method? a) 1 b) 2 c) 3 d) 4 3. The HGP was completed in a) 2002 b) 2003 c) 2004 d) 2005 4. In Maxam-gilbert method, chemical used for cytosine alteration is a) Formic acid b) hydrazine c) Dimethyl sulphate d) piperdine 5. What do you mean by 'epistasis'? a) structural genes b) functional genes c) effect of one gene on another d) all the above 6. What is pleiotropy? a) one gene affecting only one trait b) gene affecting no traits c) one gene affecting more than one trait d) two genes affecting one trait 7. The principle of sanger's method relies on a) use of chemicals for base specific cleavage dNTPs for chain b) of use termination c) use of ddNTPs for chain termination d) use of For P32 Chain termination 8. An open reading frame (ORF) is a) the sequence of a complete genome b) a plasmid vector used in genomic sequencing c) a possible gene predicted by DNA sequencing d) a fragment of a genome 9. Proteomics is a) a branch of quantum physics b) the study of algal genomes c) the study of the entire collection of proteins expressed by an organism d) study of entire set of genomes 10. Genomic libraries are useful for obtaining what product? a) periodicals on genomics research b) collections of isolated genes

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c) instructional information on how to locate the exact site of the gene of interest

d) information relating to primers and PCR

11. Protein coding genes can be identified by

a) transposon tagging b) ORF scanning c) Zoo-blotting

d) Nuclease S1 Mapping

# 2 Marks questions

- 1. What are genes?
- 2. What are pseudogenes?
- 3. Write a short note on 'Shot-gun' method.
- 4. What do you mean by 'epistasis'?
- 5. What do you mean by 'pleiotropy'?
- 6. Write any two unique applications of HGP?

# 6/8 Marks questions

1. How will you determine a short DNA sequence by 'Maxam - Gilbert' method?

- 2. Describe the 'Chain termination method' in a systematic manner with an example.
- 3. Enumerate various steps involved in a 'Genomic library construction'.

4. Discuss the merits and limitation of 'Maxam - Gilbert Method'.

5. What method will you employ to sequencing single-strand DNA fragment composed of 5 bases (5'-AGCTT-3')? Justify your answer.

- 6. Explain the role of bioinformatics tools on sequencing polynucleotide chains.
- 7. Analyze strategies, merits and imitations of 'Pyrosequencing Method' for analyzing DNA sequencing.
- 8. Describe a next-generation DNA sequencing methods in detail manner.

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#### <u>UNIT - II</u>

#### **SYLLABUS**

Managing and Distributing Genome Data: Web based servers and software for genome analysis: ENSEMBL, VISTA, UCSC Genome Browser, NCBI genome. Selected Model Organisms' Genomes and Databases.

#### Ensembl genome database project

Ensembl genome database project is a joint scientific project between the European Bioinformatics Institute and the Wellcome Trust Sanger Institute, which was launched in 1999 in response to the imminent completion of the Human Genome Project. After 10 years in existence, Ensembl's aim remains to provide a centralized resource for geneticists, molecular biologists and other researchers studying the genomes of our own species and other vertebrates and model organisms. Ensembl is one of several well-known genome browsers for the retrieval of genomic information. Similar databases and browsers are found at NCBI and the University of California, Santa Cruz (UCSC).

The human genome consists of three billion base pairs, which code for approximately 20,000–25,000 genes. However the genome alone is of little use, unless the locations and relationships of individual genes can be identified. One option is manual annotation, whereby a team of scientists tries to locate genes using experimental data from scientific journals and public databases. However this is a slow, painstaking task. The alternative, known as automated annotation, is to use the power of computers to do the complex pattern-matching of protein to DNA.

In the Ensembl project, sequence data are fed into the gene annotation system (a collection of software "pipelines" written in Perl) which creates a set of predicted gene locations and saves them in a MySQL database for subsequent analysis and display. Ensembl makes these data freely accessible to the world research community. All the data and code produced by the Ensembl project is available to download and there is also a publicly accessible database server

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allowing remote access. In addition, the Ensembl website provides computer-generated visual displays of much of the data.

Over time the project has expanded to include additional species (including key model organisms such as mouse, fruitfly and zebrafish) as well as a wider range of genomic data, including genetic variations and regulatory features. Since April 2009, a sister project, Ensembl Genomes, has extended the scope of Ensembl into invertebrate metazoa, plants, fungi, bacteria, and protists, whilst the original project continues to focus on vertebrates.

Central to the Ensembl concept is the ability to automatically generate graphical views of the alignment of genes and other genomic data against a reference genome. These are shown as data tracks, and individual tracks can be turned on and off, allowing the user to customize the display to suit their research interests. The interface also enables the user to zoom in to a region or move along the genome in either direction.

Other displays show data at varying levels of resolution, from whole karyotypes down to text-based representations of DNA and amino acid sequences, or present other types of display such as trees of similar genes (homologues) across a range of species. The graphics are complemented by tabular displays, and in many cases data can be exported directly from the page in a variety of standard file formats such as FASTA.

Externally produced data can also be added to the display, either via a DAS (Distributed Annotation System) server on the internet, or by uploading a suitable file in one of the supported formats, such as BAM, BED, or PSL.

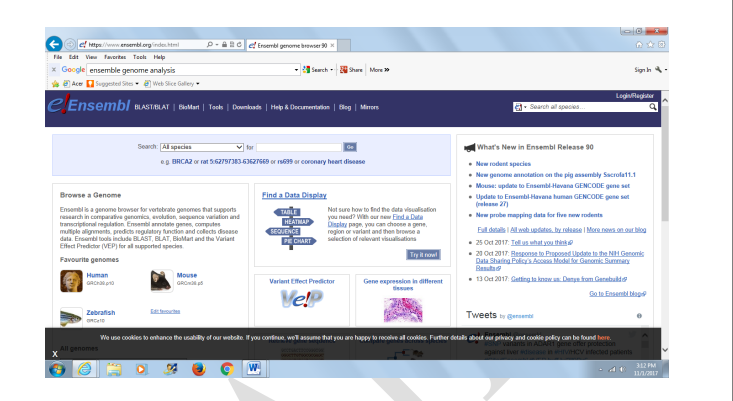
Graphics are generated using a suite of custom Perl modules based on GD, the standard Perl graphics display library.

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Alternative access method

In addition to its website, Ensembl provides a Perl API (Application Programming Interface) that models biological objects such as genes and proteins, allowing simple scripts to be written to retrieve data of interest. The same API is used internally by the web interface to display the data. It is divided in sections like the core API, the compara API (for comparative genomics data), the variation API (for accessing SNPs, SNVs, CNVs), and the functional genomics API (to access regulatory data). The Ensembl website provides extensive information on how to install and use the API.

This software can be used to access the public MySQL database, avoiding the need to download enormous datasets. The users could even choose to retrieve data from the MySQL with direct SQL queries, but this requires an extensive knowledge of the current database schema. Large datasets can be retrieved using the BioMart data-mining tool. It provides a web

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interface for downloading datasets using complex queries. Last, there is an FTP server which can be used to download entire MySQL databases as well some selected data sets in other formats.

## VISTA (comparative genomics)

VISTA is a collection of databases, tools, and servers that permit extensive comparative genomics analyses.

The VISTA family of tools is developed and hosted at Genomics Division of Lawrence Berkeley National Laboratory. This collaborative effort is supported by the Programs for Genomic Applications grant from the NHLBI/NIH and the Office of Biological and Environmental Research, Office of Science, US Department of Energy.

It was developed from modules supplied by developers at UC Berkeley, Stanford, and UC Davis, and based partly on the AVID Global Alignment program.

There are multiple VISTA servers, each allowing different types of searches.

- mVISTA can be used to align and compare your sequences to those of multiple other species
- rVISTA (regulatory VISTA) combines transcription factor binding sites database search with a comparative sequence analysis, the discovery of possible regulatory transcription factor binding sites in regions of their genes of interest. It can be used directly or through mVISTA, Genome VISTA, or VISTA Browser. A database of tissue-specific human enhancers is available through VISTA Enhancer Browser.
- GenomeVISTA allows the comparison of sequences with whole genome assemblies. It will automatically find the ortholog, obtain the alignment and VISTA plot. It allows the viewing of an alignment together with pre-computed alignments of other species in the same interval.
- Phylo-VISTA allows the analysis of multiple DNA sequence alignments of sequences from different species while considering their phylogenic relationships.
- wgVISTA allows the alignment of sequences up to 10Mb long (finished or draft) including microbial whole-genome assemblies.

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Researchers can use the VISTA Browser:

- to examine pre-computed alignments among a variety of species
- to submit sequences of their own (not limited by the species collection already in the database)

#### Genomes

There are more than 28 searchable genomes, including vertebrate, non-vertebrate, plants, fungi, algae, bacteria, and others. More are continually being added. These include:

- Human—Orangutan—Rhesus—Marmoset—Horse—Dog—Mouse—Rat—Chicken
- Drosophila spp.
- Arabidopsis—Rice—Sorghum
- E. coli—mycoplasma—nitrosomonas

Collaboration with other projects

Pre-computed full scaffold alignments for microbial genomes are available as the VISTA component of IMG (Integrated Microbial Genomes System) developed in the DOE (Department of Energy's) Joint Genome Institute.

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	VISTA Tools for Comparative Genomics About Us 💯 Cite Us 🙆 Contact Us		
	VISTA Home Custom Alignment Browser Enhancer DB Downloads Publications Help		
	Clade Genome Release Position           Vertebrate         V           Feb. 2009         Chrl9:107.543,285-107,690.4		
	VISTA-Point     VISTA Browser (Requires Java2)		
	Whole Genome Comparative Analysis of the Human Feb. 2009 Genome		
	From this page you can access the results of:		
	the multiple alignments		
	Human Feb. 2009. Chimp Mar. 2006. Callithrix jacchus v 2.0.2, Rhesus Jan. 2006 and Pongo pygmaeus abelii v 2.0.2 genomes.		
	the pairwise alignments of the		
	Human Feb. 2009 genome produced by the Genome Reference Consortium with the following genomes:		
	numan rec. 2009 genome produced by the Genome Reference Consortium with the following genomes.		

VISTA is a comprehensive suite of programs and databases for comparative analysis of genomic sequences. There are two ways of using VISTA - you can submit your own sequences and alignments for analysis (VISTA servers) or examine pre-computed whole-genome alignments of different species.

mVISTA



• mVISTA

Align and compare your sequences from multiple species

• gVISTA

Compare your sequences against whole-genome assemblies.

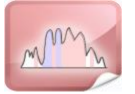
• wgVISTA

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Align pair of sequences up to 10Mb long (finished or draft) including microbial wholegenome assemblies.

Precomputed Alignments

VISTA Browser



VISTA-Point

Access complete data and visual presentation of pairwise and multiple alignments of whole genome assemblies.

VISTA Browser

Examine pre-computed pairwise and multiple alignments of whole genome assemblies.

Microbial Genomes

Access pre-computed full scaffold alignments for microbial genomes through the VISTA component of IMG.

### **UCSC Genome Browser**

The UCSC Genome Browser is an on-line genome browser hosted by the University of California, Santa Cruz (UCSC). It is an interactive website offering access to genome sequence data from a variety of vertebrate and invertebrate species and major model organisms, integrated with a large collection of aligned annotations. The Browser is a graphical viewer optimized to support fast interactive performance and is an open-source, web-based tool suite built on top of a MySQL database for rapid visualization, examination, and querying of the data at many levels. The Genome Browser Database, browsing tools, downloadable data files, and documentation can all be found on the UCSC Genome Bioinformatics website.

Initially built and still managed by Jim Kent, then a graduate student, and David Haussler, professor of Computer Science (now Biomolecular Engineering) at the University of

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California, Santa Cruz in 2000, the UCSC Genome Browser began as a resource for the distribution of the initial fruits of the Human Genome Project. Funded by the Howard Hughes Medical Institute and the National Human Genome Research Institute, NHGRI (one of the US National Institutes of Health), the browser offered a graphical display of the first full-chromosome draft assembly of human genome sequence. Today the browser is used by geneticists, molecular biologists and physicians as well as students and teachers of evolution for access to genomic information.

In the years since its inception, the UCSC Browser has expanded to accommodate genome sequences of all vertebrate species and selected invertebrates for which high-coverage genomic sequences is available, now including 46 species. High coverage is necessary to allow overlap to guide the construction of larger contiguous regions. Genomic sequences with less coverage are included in multiple-alignment tracks on some browsers, but the fragmented nature of these assemblies does not make them suitable for building full featured browsers. (more below on multiple-alignment tracks). The species hosted with full-featured genome browsers are shown in the table.

great apes

human, baboon, bonobo, chimp, gibbon, gorilla, orangutan

non-ape primates

bushbaby, marmoset, mouse lemur, rhesus macaque, squirrel monkey, tarsier, tree shrew

non-primate mammals

mouse, alpaca, armadillo, cat, Chinese hamster, cow, dog, dolphin, elephant, ferret, guinea pig, hedgehog, horse, kangaroo rat, manatee, Minke whale, naked mole-rat, opossum, panda, pig, pika, platypus, rabbit, rat, rock hyrax, sheep, shrew, sloth, squirrel, Tasmanian devil, tenrec, wallaby, white rhinoceros

non-mammal chordates

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American alligator, Atlantic cod, budgerigar, chicken, coelocanth, elephant shark, Fugu, lamprey, lizard, medaka, medium ground finch, Nile tilapia, painted turtle, stickleback, Tetraodon, turkey, Xenopus tropicalis, zebra finch, zebrafish

invertebrates

Caenorhabditis spp (5), Drosophila spp. (11), Ebola virus, honey bee, lancelet, mosquito, P. Pacificus, sea hare, sea squirt, sea urchin, yeast

The large amount of data about biological systems that is accumulating in the literature makes it necessary to collect and digest information using the tools of bioinformatics. The UCSC Genome Browser presents a diverse collection of annotation datasets (known as "tracks" and presented graphically), including mRNA alignments, mappings of DNA repeat elements, gene predictions, gene-expression data, disease-association data (representing the relationships of genes to diseases), and mappings of commercially available gene chips (e.g., Illumina and Agilent). The basic paradigm of display is to show the genome sequence in the horizontal dimension, and show graphical representations of the locations of the mRNAs, gene predictions, etc. Blocks of color along the coordinate axis show the locations of the alignments of the various data types. The ability to show this large variety of data types on a single coordinate axis makes the browser a handy tool for the vertical integration of the data.

To find a specific gene or genomic region, the user may type in the gene name, (e.g., BRCA1) an accession number for an RNA, the name of a genomic cytological band (e.g., 20p13 for band 13 on the short arm of chr20) or a chromosomal position (chr17:38,450,000-38,531,000 for the region around the gene BRCA1).

Presenting the data in the graphical format allows the browser to present link access to detailed information about any of the annotations. The gene details page of the UCSC Genes track provides a large number of links to more specific information about the gene at many other data resources, such as Online Mendelian Inheritance in Man (OMIM) and SwissProt.

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Designed for the presentation of complex and voluminous data, the UCSC Browser is optimized for speed. By pre-aligning the 55 million RNAs of GenBank to each of the 81 genome assemblies (many of the 46 species have more than one assembly), the browser allows instant access to the alignments of any RNA to any of the hosted species.

Multiple gene products of FOXP2 gene (top) and evolutionary conservation shown in multiple alignment (bottom). The juxtaposition of the many types of data allow researchers to display exactly the combination of data that will answer specific questions. A pdf/postscript output functionality allows export of a camera-ready image for publication in academic journals. One unique and useful feature that distinguishes the UCSC Browser from other genome browsers is the continuously variable nature of the display. Sequence of any size can be displayed, from a single DNA base up to the entire chromosome (human chr1 = 245 million bases, Mb) with full annotation tracks. Researchers can display a single gene, a single exon, or an entire chromosome band, showing dozens or hundreds of genes and any combination of the many annotations. A convenient drag-and-zoom feature allows the user to choose any region in the genome image and expand it to occupy the full screen.

Researchers may also use the browser to display their own data via the Custom Tracks tool. This feature allows users to upload a file of their own data and view the data in the context of the reference genome assembly. Users may also use the data hosted by UCSC, creating subsets of the data of their choosing with the Table Browser tool (such as only the SNPs that change the amino acid sequence of a protein) and display this specific subset of the data in the browser as a Custom Track.

Any browser view created by a user, including those containing Custom Tracks, may be shared with other users via the Saved Sessions tool.

#### Variation data

Many types of variation data are also displayed. For example, the entire contents of each release of the dbSNP database from NCBI are mapped to human, mouse and other genomes. This includes the fruits of the 1000 Genomes Project, as soon as they are released in dbSNP.

S

Other types of variation data include copy-number variation data (CNV) and human population allele frequencies from the HapMap project.

The Genome Browser offers a unique set of comparative-genomic data for most of the species hosted on the site. The comparative alignments give a graphical view of the evolutionary relationships among species. This makes it a useful tool both for the researcher, who can visualize regions of conservation among a group of species and make predictions about functional elements in unknown DNA regions, and in the classroom as a tool to illustrate one of the most compelling arguments for the evolution of species. The 44-way comparative track on the human assembly clearly shows that the farther one goes back in evolutionary time, the less sequence homology remains, but functionally important regions of the genome (e.g., exons and control elements, but not introns typically) are conserved much farther back in evolutionary time.

#### Analysis tools

More than simply a genome browser, the UCSC site hosts a set of genome analysis tools, including a full-featured GUI interface for mining the information in the browser database (the Table Browser), a fast sequence alignment tool (BLAT) that is also useful for simply finding sequences in the massive sequence (human genome = 2.8 billion bases, Gb) of any of the featured genomes.

A liftOver tool uses whole-genome alignments to allow conversion of sequences from one assembly to another or between species. The Genome Graphs tool allows users to view all chromosomes at once and display the results of genome-wide association studies (GWAS). The Gene Sorter displays genes grouped by parameters not linked to genome location, such as expression pattern in tissues.

#### **Creating spreadsheet links to UCSC Genome Browser views**

Many users of the Genome Browser gather data of their own in Excel spreadsheets and would like to create links to the Browser using data in the spreadsheet. For example, a clinical geneticist may have lists of regions for a patient that are duplicated or deleted, as determined by

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comparative genomic hybridization (CGH). These regions can be the source information for a browser view allowing access to each region with a single click.

			hg18	hg19		
÷ •			-	-		
* chrom	start	end	links	links		
3	12000000	15000000	<u>ucsc</u>	<u>ucsc</u>		
chr3	12000000	15000000	<u>ucsc</u>	<u>ucsc</u>		
NOTE: D	ifferent ch	romNames	require	different e	excel link	
hrN is sta	andard ucs	sc format				Click to download the spreadshe
						Cher to download the spreadshe
		gene				ucscLinks.xls
		FGFR1	<u>ucsc</u>	<u>ucsc</u>		
		EGFR	<u>ucsc</u>	<u>ucsc</u>		
		position				r i i i i i i i i i i i i i i i i i i i
		15q11	<u>ucsc</u>	<u>ucsc</u>		
chr.	3:12000000	-15000000	ucsc	ucsc		

Careful use of Excel's "copy" and "move" functions should allow the links on this sheet to be used without modification.

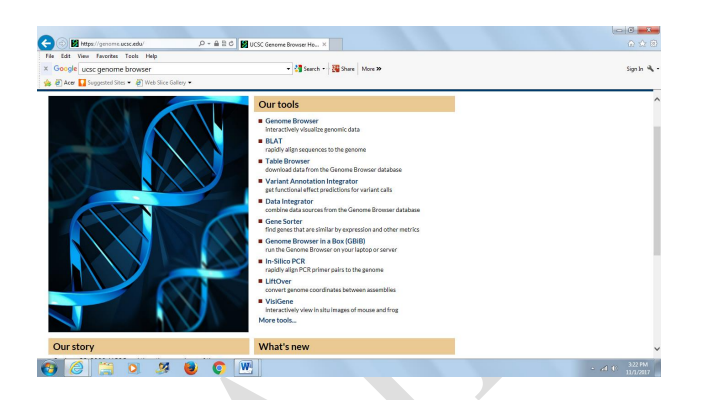
On June 22, 2000, UCSC and the other members of the International Human Genome Project consortium completed the first working draft of the human genome assembly, forever ensuring free public access to the genome and the information it contains. A few weeks later, on July 7, 2000, the newly assembled genome was released on the web at http://genome.ucsc.edu, along with the initial prototype of a graphical viewing tool, the UCSC Genome Browser. In the ensuing years, the website has grown to include a broad collection of vertebrate and model organism assemblies and annotations, along with a large suite of tools for viewing, analyzing and downloading data.

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## **National Center for Biotechnology Information**

The National Center for Biotechnology Information (NCBI) is part of the United States National Library of Medicine (NLM), a branch of the National Institutes of Health (NIH). The NCBI is located in Bethesda, Maryland and was founded in 1988 through legislation sponsored by Senator Claude Pepper. The NCBI houses a series of databases relevant to biotechnology and biomedicine and is an important resource for bioinformatics tools and services. Major databases include GenBank for DNA sequences and PubMed, a bibliographic database for the biomedical literature. Other databases include the NCBI Epigenomics database. All these databases are available online through the Entrez search engine.

NCBI was directed by David Lipman, one of the original authors of the BLAST sequence alignment program and a widely respected figure in bioinformatics. He also leads an intramural research program, including groups led by Stephen Altschul (another BLAST co-author), David

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Landsman, Eugene Koonin (a prolific author on comparative genomics), John Wilbur, Teresa Przytycka, and Zhiyong Lu. David Lipman stood down from his post in May 2017.

NCBI is listed in the Registry of Research Data Repositories re3data.org.

#### GenBank

NCBI has had responsibility for making available the GenBank DNA sequence database since 1992. GenBank coordinates with individual laboratories and other sequence databases such as those of the European Molecular Biology Laboratory (EMBL) and the DNA Data Bank of Japan (DDBJ).

Since 1992, NCBI has grown to provide other databases in addition to GenBank. NCBI provides Gene, Online Mendelian Inheritance in Man, the Molecular Modeling Database (3D protein structures), dbSNP (a database of single-nucleotide polymorphisms), the Reference Sequence Collection, a map of the human genome, and a taxonomy browser, and coordinates with the National Cancer Institute to provide the Cancer Genome Anatomy Project. The NCBI assigns a unique identifier (taxonomy ID number) to each species of organism.

The NCBI has software tools that are available by WWW browsing or by FTP. For example, BLAST is a sequence similarity searching program. BLAST can do sequence comparisons against the GenBank DNA database in less than 15 seconds.

#### **NCBI Bookshelf**

The "NCBI Bookshelf is a collection of freely accessible, downloadable, on-line versions of selected biomedical books. The Bookshelf covers a wide range of topics including molecular biology, biochemistry, cell biology, genetics, microbiology, disease states from a molecular and cellular point of view, research methods, and virology. Some of the books are online versions of previously published books, while others, such as Coffee Break, are written and edited by NCBI staff. The Bookshelf is a complement to the Entrez PubMed repository of peer-reviewed publication abstracts in that Bookshelf contents provide established perspectives on evolving areas of study and a context in which many disparate individual pieces of reported research can be organized.

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#### **Basic Local Alignment Search Tool (BLAST)**

BLAST is an algorithm used for calculating sequence similarity between biological sequences such as nucleotide sequences of DNA and amino acid sequences of proteins. BLAST is a powerful tool for finding sequences similar to the query sequence within the same organism or in different organisms. It searches the query sequence on NCBI databases and servers and post the results back to the person's browser in chosen format. Input sequences to the BLAST are mostly in FASTA or Genbank format while output could be delivered in variety of formats such as HTML, XML formatting and plain text. HTML is the default output format for NCBI's webpage. Results for NCBI-BLAST are presented in graphical format with all the hits found, a table with sequence identifiers for the hits having scoring related data, along with the alignments for the sequence of interest and the hits received with analogous BLAST scores for these.

#### Entrez

The Entrez Global Query Cross-Database Search System is used at NCBI for all the major databases such as Nucleotide and Protein Sequences, Protein Structures, PubMed, Taxonomy, Complete Genomes, OMIM, and several others. Entrez is both indexing and retrieval system having data from various sources for biomedical research. NCBI distributed the first version of Entrez in 1991, composed of nucleotide sequences from PDB and GenBank, protein sequences from SWISS-PROT, translated GenBank, PIR, PRF and PDB and associated abstracts and citations from PubMed. Entrez is specially designed to integrate the data from several different sources, databases and formats into a uniform information model and retrieval system which can efficiently retrieve that relevant references, sequences and structures.

#### Gene

Gene has been implemented at NCBI to characterize and organize the information about genes. It serves as a major node in the nexus of genomic map, expression, sequence, protein function, structure and homology data. A unique GeneID is assigned to each gene record that can be followed through revision cycles. Gene records for known or predicted genes are established here and are demarcated by map positions or nucleotide sequence. Gene has several advantages

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over its predecessor, LocusLink, including, better integration with other databases in NCBI, broader taxonomic scope, and enhanced options for query and retrieval provided by Entrez system.

#### Protein

Protein database is an important protein resource at NCBI. It maintains the text record for individual protein sequences, derived from many different resources such as NCBI Reference Sequence (RefSeq) project, GenbBank, PDB and UniProtKB/SWISS-Prot. Protein records are present in different formats including FASTA and XML and are linked to other NCBI resources. Protein provides the relevant data to the users such as genes, DNA/RNA sequences, biological pathways, expression and variation data and literature. It also provides the pre-determined sets of similar and identical proteins for each sequence as computed by the BLAST. The Structure database of NCBI contains 3D coordinate sets for experimentally-determined structures in PDB that are imported by NCBI. The Conserved Domain database (CDD) of protein contains sequence profiles that characterize highly conserved domains within protein sequences. It also has records from external resources like SMART and Pfam. There is another database in protein known as Protein Clusters database which contains sets of proteins sequences sequences as calculated by BLAST.

#### Pubchem BioAssay database

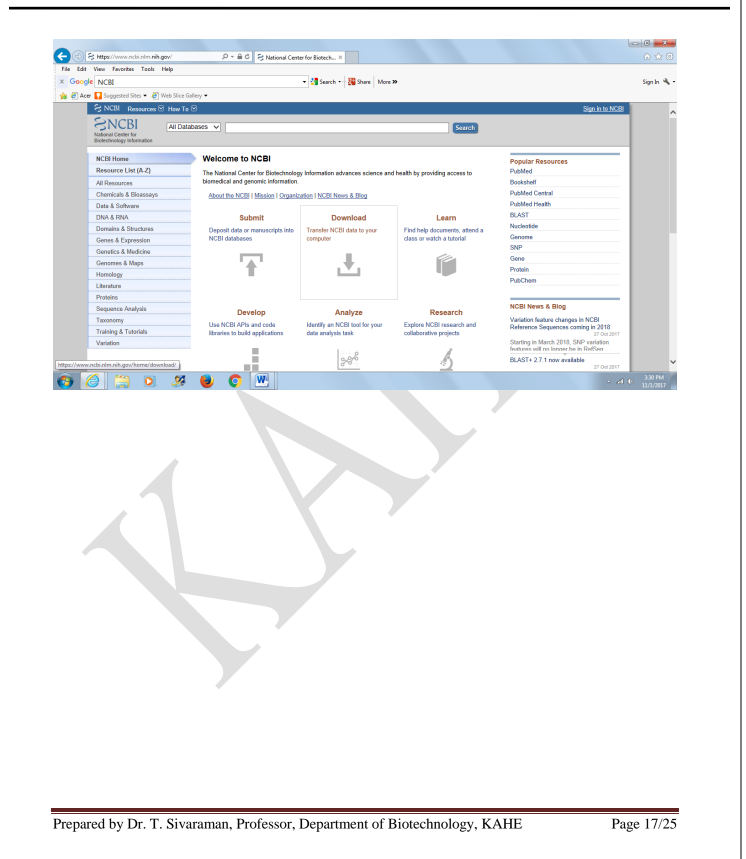
PubChem BioAssay database of NCBI is a public resource for biological tests of small molecules and siRNA reagents. The major purpose of PubChem repository is to provide easy and free of cost access to all deposited data, and to provide intuitive data analysis tools. It is structured as a set of relational databases organized on Microsoft SQL servers. PubChem's BioAssay data is searchable and accessible by Entrez information retrieval system. PubChem database provides programmatic and Web-based tools for users to search, review, and download a publications, bioactivity data for a compound, a BioAssay record, a molecular target.

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## **Genome and Organism – Specific Databases**

Table 1. A small selection of organism-specific genomic databases available on the WWW.

Organism	Database/resource	URL	
Escherichia coli	EcoGene EcoCyc (Encyclopedia of <i>E. coli</i> genes and metabolism)	http://bmb.med.miami.edu/EcoGene/EcoWeb/ http://ecocyc.pangeasystems.com/ecocyc/ecocyc.html	
	Colibri	http://genolist.pasteur.fr/Colibri/	
Bacillus subtilis	SubtiList	http://genolist.pasteur.fr/SubtiList/	
Saccharomyces cerevisiae	Saccharomyces Genome Database (SGD)	http://genome-www.stanford.edu/Saccharomyces/	
Plasmodium falciparum	PlasmoDB	http://PlasmoDB.org	
Arabidopsis thaliana	MIPS Arabidopsis thaliana Database (MAtDB)	http://mips.gsf.de/proj/thal/db	
	The Arabidopsis information resource (TAIR)	http://www.arabidopsis.org/	
Drosophila melanogaster	FlyBase	http://flybase.bio.indiana.edu/	
Caenorhabditis elegans	A C. elegans DataBase (ACeDB)	http://www.acedb.org/	
Mouse	Mouse Genome Database (MGD)	http://www.informatics.jax.org/	
Human	OnLine Mendelian Inheritance in Man (OMIM)	http://www.ncbi.nlm.nih.gov/omim	

These databases are actively curated by members of the research community working on the particular organism of interest and generally include links to organism-specific resources such as clone sets and mutant strains.

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Table 2. Useful gateway sites providing information and links to multiple, organism-specific and genomic resources.

Gateway site	URL
NCBI Genomic Biology	http://www.ncbi.nlm.nih.gov/Genomes/index.html
GOLD (Genomes OnLine Database)	http://wit.integratedgenomics.com/GOLD/
Organism-specific genome databases	http://www.unl.edu/stc-95/ResTools/biotools/biotools10.html
TIGR Microbial Database	http://www.tigr.org/tdb/mdb/mdbcomplete.html
Bacterial genomes	http://genolist.pasteur.fr/
Yeast databases	http://genome-www.stanford.edu/Saccharomyces/yeast_info.html
EnsEMBL genome database project	http://www.ensembl.org/
MIPS (Munich Information Center for Protein Sequences)	http://mips.gsf.de

Table 2	Detabase	taala far	diantautaa c	and appareting.	annomia nogunnon data
raure o.	Database i	10018 $101$	uispiaying a	ano annotating.	genomic sequence data.

Viewer format	URL for further information and tutorials
Artemis	http://www.sanger.ac.uk/Software/Artemis
ACeDB	http://www.acedb.org/Tutorial/brief-tutorial.shtml
Apollo	http://www.ensembl.org/apollo/
EnsEMBL	http://www.ensembl.org
NCBI map viewer	http://www.ncbi.nlm.nih.gov/
GoldenPath	http://genome.ucsc.edu/

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#### <u>FlyBase</u>

FlyBase is an online bioinformatics database and the primary repository of genetic and molecular data for the insect family Drosophilidae. For the most extensively studied species and model organism, Drosophila melanogaster, a wide range of data are presented in different formats. Information in FlyBase originates from a variety of sources ranging from large-scale genome projects to the primary research literature. These data types include mutant phenotypes, molecular characterization of mutant alleles and other deviations, cytological maps, wild-type expression patterns, anatomical images, transgenic constructs and insertions, sequence-level gene models and molecular classification of gene product functions. Query tools allow navigation of FlyBase through DNA or protein sequence, by gene or mutant name, or through terms from the several ontologies used to capture functional, phenotypic, and anatomical data. The database offers several different query tools in order to provide efficient access to the data available and facilitate the discovery of significant relationships within the database. Links between FlyBase and external databases, such as BDGP or modENCODE, provide opportunity for further exploration into other model organism databases and other resources of biological and molecular information.<sup>[3]</sup> The FlyBase project is carried out by a consortium of *Drosophila* researchers and computer scientists at Harvard University and Indiana University in the United States, and University of Cambridge in the United Kingdom.

*Drosophila melanogaster* has been an experimental organism since the early 1900s, and has since been placed at the forefront of many areas of research. As this field of research spread and became global, researchers working on the same problems needed a way to communicate and monitor progress in the field. This niche was initially filled community newsletters such as the Drosophila Information Service (DIS), which dates back to 1934 when the field was starting to spread from Thomas Hunt Morgan's lab. Material in these presented regular 'catalogs' of mutations bibliographies of the Drosophila literature. As computer infrastructure developed in the 80's and 90's, these newsletters gave way and merged with internet mailing lists, and these eventually became online resources and data. In 1992, data on the genetics and genomics of *D*.

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*melanogaster* and related species were electronically available over the Internet through the funded FlyBase, BDGP (Berkeley Drosophila Genome Project) and EDGP (European Drosophila Genome Project) informatics groups. These groups recognized that most genome project and community data types overlapped. They decided it would be of value to present the scientific community with an integrated view of the data. In October 1992, the National Center for Human Genome Research of the NIH funded the FlyBase project with the objective of designing, building and releasing a database of genetic and molecular information concerning *Drosophila melanogaster*. FlyBase also receives support from the Medical Research Council, London. In 1998, the FlyBase consortium integrated the information into a single Drosophila genomics server.

FlyBase contains a complete annotation of the *Drosophila melanogaster* genome that is updated several times per year. It also includes a searchable bibliography of research on *Drosophila* genetics in the last century. Information on current researchers, and a partial pedigree of relationships between current researchers, is searchable, based on registration of the participating scientist (Find a Person). The site also provides a large database of images illustrating the full genome, and several movies detailing embryogenesis (ImageBrowser).

Search Strategies - Gene reports for genes from all twelve sequenced Drosophila genomes are available in FlyBase. There are four main ways this data can be browsed: Precomputed Files, BLAST, Gbrowse, and Gene Report Pages. Gbrowse and precomputed files are for genome-wide analysis, bioinformatics, and comparative genomics. BLAST and gene report pages are for a specific gene, protein, or region across the species. When looking for cytology there are two main tools available. Use Cytosearch when looking for cytologically-mapped genes or deficiencies, that haven't been molecularly mapped to the sequence. Use Gbrowse when looking for molecularly mapped sequences, insertions, or Affymetrix probes.

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There are two main query tools in FlyBase. The first main query tool is called Jump to Gene (J2G). This is found in the top right of the blue navigation bar on every page of FlyBase. This tool is useful when you know exactly what you are looking for and want to go to the report page with that data. The second main query tool is called QuickSearch. This is located on the FlyBase homepage. This tool is most useful when you want to look up something quickly that you may only know a little about. Searching can be performed within D. melanogaster only or within all species. Data other than genes can be searched using the 'data class' menu.

### **Online Mendelian Inheritance in Man**

Online Mendelian Inheritance in Man (OMIM) is a continuously updated catalog of human genes and genetic disorders and traits, with a particular focus on the gene-phenotype relationship. As of 12 February 2017, approximately 8,425 of the over 23,000 entries in OMIM represented phenotypes; the rest represented genes, many of which were related to known phenotypes. OMIM is the online continuation of Dr. Victor McKusick's *Mendelian Inheritance in Man* (MIM), which was published in 12 editions between 1966 and 1998. Nearly all of the 1,486 entries in the first edition of MIM discussed phenotypes.

MIM/OMIM is produced and curated at the Johns Hopkins University School of Medicine (JHUSOM). OMIM became available on the internet in 1987 under the direction of the Welch Medical Library at JHUSOM with financial support from the Howard Hughes Medical Institute. From 1995 to 2010, OMIM was available on the World Wide Web with informatics and financial support from the National Center for Biotechnology Information. The current OMIM website (OMIM.org), which was developed with funding from JHUSOM, is maintained by Johns Hopkins University with financial support from the National Human Genome Research Institute. The content of MIM/OMIM is based on selection and review of the published peerreviewed biomedical literature. Updating of content is performed by a team of science writers and curators under the direction of Dr. Ada Hamosh at the McKusick-Nathans Institute of Genetic Medicine of Johns Hopkins University. While OMIM is freely available to the public, it

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is designed for use primarily by physicians and other health care professionals concerned with genetic disorders, by genetics researchers, and by advanced students in science and medicine. The database may be used as a resource for locating literature relevant to inherited conditions, and its numbering system is widely used in the medical literature to provide a unified index for genetic diseases.

#### MIM classification system - MIM numbers

Each OMIM entry is given a unique six-digit identifier as summarized below:

- 100000–299999: Autosomal loci or phenotypes (entries created before May 15, 1994)
- 300000–3999999: X-linked loci or phenotypes
- 400000–4999999: Y-linked loci or phenotypes
- 500000–5999999: Mitochondrial loci or phenotypes

600000 and above: Autosomal loci or phenotypes (entries created after May 15, 1994)

In cases of allelic heterogeneity, the MIM number of the entry is followed by a decimal point and a unique 4-digit number specifying the variant. For example, allelic variants in the HBB gene (141900) are numbered 141900.0001 through 141900.0538.

Because OMIM has responsibility for the classification and naming of genetic disorders, these numbers are stable identifiers of the disorders.

Symbols preceding MIM numbers

Symbols preceding MIM numbers indicate the entry category:

- An asterisk (\*) before an entry number indicates a gene.
- A number symbol (#) before an entry number indicates that it is a descriptive entry, usually of a phenotype, and does not represent a unique locus. The reason for the use of the number symbol is given in the first paragraph of the entry. Discussion of any gene(s) related to the phenotype resides in another entry (or entries) as described in the first paragraph.
- A plus sign (+) before an entry number indicates that the entry contains the description of a gene of known sequence and a phenotype.

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- A percent sign (%) before an entry number indicates that the entry describes a confirmed Mendelian phenotype or phenotypic locus for which the underlying molecular basis is not known.
- No symbol before an entry number generally indicates a description of a phenotype for which the Mendelian basis, although suspected, has not been clearly established or that the separateness of this phenotype from that in another entry is unclear.
- A caret (^) before an entry number means the entry no longer exists because it was removed from the database or moved to another entry as indicated.

### **Possible questions**

#### **1 Mark questions**

The human genome consists of \_\_\_\_\_ base pairs.
 a) two billion b) ten billion c) one billion d) three billion

2. Large datasets can be retrieved using the \_\_\_\_\_\_ tool. a) BioMart b) SQL c) NCBI d) FASTA

3. \_\_\_\_\_ can be used to combines transcription factor binding sites database search with a comparative sequence analysis. a) rVISTA b) mVISTA c) wgVISTA d) phylo-VISTA

4. ENSG### is \_\_\_\_

a) Ensembl Exon ID b) Ensembl Gene ID c) Ensembl Transcript ID d) Ensembl Peptide ID

6. \_\_\_\_is an algorithm used for calculating sequence similarity between biological sequences.a) VISTAb) NCBIc) BLASTd) FASTA

7. Protein-coding genes can be identified by<br/>a) Transposon taggingb) ORF scanningc) Zoo-blottingd)Nuclease S1mapping

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8. ORF scann a) to find exor d) to find prot	-	find intergenic sequences c) to find gene homologies
9. Human Ger	nome Project was firs	t initiated in the year
a) 1965	b) 1970	c) 1975 d) 1980
10 A	transplant was use	d to overcome this genetic disorder.
a) Liver	-	c) Bone marrow d) none of the above
	latabase focuses on b) Invertebra	
2 Marks ques	<u>tions</u>	
1. Write a shore	t note on 'UCSC web	server'.

- 2. Write any two applications of NCBI database.
- 3. What are databases? Give an example.
- 4. What are genome-specific databases? Give an example.

## 6/8 Marks questions

- 1. Explain various applications of the 'ENSEMBLE' database on analysing genome data.
- 2. Describe the use of 'VISTA' webserver in the analyses of genome data.
- 3. Enumerate unique features of 'NCBI' database for retrieving genome data.
- 4. Describe any two organisms specific genome database in a detailed manner.

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#### **SYLLABUS**

**Genomic mapping:** Genetic markers - VNTR, mini and micro satellites, STS, SNP, ESTs. Types of genome maps, Mapping techniques – Physical and genetic mapping, Map resources, Practical uses genome maps.

#### **Genome Mapping**

Among the main goals of the Human Genome Project (HGP) was to develop new, better and cheaper tools to identify new genes and to understand their function.

One of these tools is genetic mapping. Genetic mapping - also called linkage mapping - can offer firm evidence that a disease transmitted from parent to child is linked to one or more genes. Mapping also provides clues about which chromosome contains the gene and precisely where the gene lies on that chromosome.

Genetic maps have been used successfully to find the gene responsible for relatively rare, single-gene inherited disorders such as cystic fibrosis and Duchenne muscular dystrophy.

Genetic maps are also useful in guiding scientists to the many genes that are believed to play a role in the development of more common disorders such as asthma, heart disease, diabetes, cancer, and psychiatric conditions.

In 1911, by Thomas Hunt Morgan, gene for eye-color was located on the X chromosome of fruit fly.

Shortly after that, E.B. Wilson attributed the sex-linkedv genes responsible for color- blindness and hemophilia in human beings to be located on the X-chromosome, similar to the many X-linked factors being described by the Morgan group in flies.

It wasn't until 1968 that an autosomal assignment of v linkage was made by Donahue---"Duffy" was assigned to Chromosome #1.

"Gene mapping" refers to the mapping of genes to specific locations on chromosomes. It is a critical step in the understanding of genetic diseases.

There are two types of gene mapping:

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Genetic Mapping - using linkage analysis to determine the relative position between two genes on a chromosome.

Physical Mapping - using all available techniques or information to determine the absolute position of a gene on a chromosome.

#### **Genetic mapping**

Uses genetic techniques to construct maps showing the positions of genes and other sequence features on a genome.

Requires informative markers – polymorphic and a population with known relationships.

Best if measured between "close" markers.v

Unit of distance in genetic maps = centiMorgans, cMv

1 cM = 1% chance of recombination between markersv

Genetic techniques include crossbreeding experiments or, in the case of humans, the

examination of family histories (pedigrees).

#### Markers for genetic mapping

The first genetic maps, constructed in the organisms such as the fruit fly, used genes as markers.

The only genes that could be studied were those specifying phenotypes that were

distinguishable by visual examination.

Eg. Eye color, height.

Some organisms have very few visual characteristics so gene mapping with these organisms has to rely on biochemical phenotypes/

#### Biochemical markers in Human

In human the biochemical phenotypes that can be scored by blood typing.

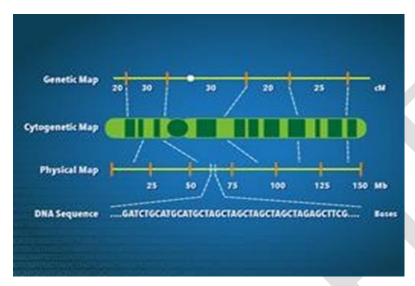
These include the standard blood groups such as the ABO series and also the human leukocyte antigens (the HLA system).

A big advantage of these markers is that many of the relevant genes have multiple alleles. For example, the gene called HLA-DRB1 has at least 290 alleles and HLA-B has over 400.

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This is relevant because if all the family members have the same allele for the gene being studied then no useful information can be obtained.



#### Drawbacks of using gene as marker

Genes are very useful markers but they are by no means ideal.

One problem, especially with larger genomes such as those of vertebrates and flowering

plants, is that a map based entirely on genes is not very detailed.

#### **DNA markers**

As with gene markers, a DNA marker must have at least two alleles to be useful.

There are three types of DNA sequence feature that satisfy this requirement:

Restriction fragment length polymorphisms (RFLPs),

Simple sequence length polymorphisms (SSLPs), and

Single nucleotide polymorphisms (SNPs).

#### **Restriction Fragment Length Polymorphism (RFLP)**

#### Introduction

Restriction Fragment Length Polymorphism (RFLP) is a difference in homologous DNA sequences that can be detected by the presence of fragments of different lengths after digestion of the DNA samples in question with specific restriction endonucleases. RFLP, as a molecular

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marker, is specific to a single clone/restriction enzyme combination.

Most RFLP markers are co-dominant (both alleles in heterozygous sample will be detected) and highly locus-specific.

An RFLP probe is a labeled DNA sequence that hybridizes with one or more fragments of the digested DNA sample after they were separated by gel electrophoresis, thus revealing a unique blotting pattern characteristic to a specific genotype at a specific locus. Short, single- or low-copy genomic DNA or cDNA clones are typically used as RFLP probes.

The RFLP probes are frequently used in genome mapping and in variation analysis (genotyping, forensics, paternity tests, hereditary disease diagnostics, etc.).

#### **Developing RFLP probes**

Total DNA is digested with a methylation-sensitive enzyme (for example, *PstI*), thereby enriching the library for single- or low-copy expressed sequences (*PstI* clones are based on the suggestion that expressed genes are not methylated).

The digested DNA is size-fractionated on a preparative agarose gel, and fragments ranging from 500 to 2000 bp are excised, eluted and cloned into a plasmid vector (for example, pUC18).

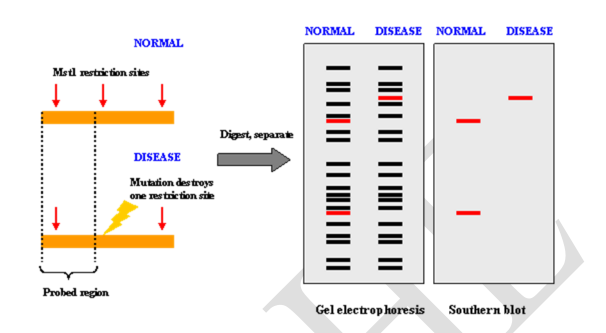
Digests of the plasmids are screened to check for inserts.

Southern blots of the inserts can be probed with total sheared DNA to select clones that hybridize to single- and low-copy sequences.

The probes are screened for RFLPs using genomic DNA of different genotypes digested with restriction endonucleases. Typically, in species with moderate to high polymorphism rates, two to four restriction endonucleases are used such as *Eco*RI, *Eco*RV, and *Hin*dIII. In species with low polymorphism rates, additional restriction endonucleases can be tested to increase the chance of finding polymorphism.

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#### Simple sequence length polymorphisms (SSLPs)

SSLPs are arrays of repeat sequences that display length variations, different alleles containing different numbers of repeat units.

Unlike RFLPs that can have only two alleles, SSLPs can be multi-allelic as each SSLP can have a number of different length variants.

There are two types of SSLP, both of which were described in Minisatellites, also known as variable number of tandem repeats (VNTRs), in which the repeat unit is up to 25 bp in length. Microsatellites or simple tandem repeats (STRs), whose repeats are shorter, usually dinucleotide or tetra-nucleotide units.

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#### Single nucleotide polymorphisms (SNPs)

Single nucleotide polymorphisms, frequently called SNPs (pronounced "snips"), are the most common type of genetic variation among people.

Each SNP represents a difference in a single DNA building block, called a nucleotide. For example, a SNP may replace the nucleotide cytosine (C) with the nucleotide thymine (T) in a certain stretch of DNA.

SNPs occur normally throughout a person"s DNA. They occur once in every 300 nucleotides on average, which means there are roughly 10 million SNPs in the human genome.

Most commonly, these variations are found in the DNA between genes. They can act as biological markers, helping scientists locate genes that are associated with disease.

When SNPs occur within a gene or in a regulatory region near a gene, they may play a more direct role in disease by affecting the gene"s function.

Most SNPs have no effect on health or development. Some of these genetic differences, however, have proven to be very important in the study of human health.

Researchers have found SNPs that may help predict an individual"s response to certain drugs, susceptibility to environmental factors such as toxins, and risk of developing particular diseases.

SNPs can also be used to track the inheritance of disease genes within families.

Future studies will work to identify SNPs associated with complex diseases such as heart disease, diabetes, and cancer.

#### Oligonucleotide hybridization

Oligonucleotide hybridization can therefore discriminate between the two alleles of an SNP. Various screening strategies have been devised including DNA chip technology and solution hybridization techniques.

#### Linkage analysis is the basis of genetic mapping

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Chromosomes are inherited as intact units, so it was reasoned that the alleles of some pairs of genes will be inherited together because they are on the same chromosome.

This is the principle of genetic linkage, Pairs of genes were either inherited independently, as expected for genes in different chromosomes, or, if they showed linkage, then it was only partial linkage: sometimes they were inherited together and sometimes they were not.

The frequency with which the genes are unlinked by crossovers will be directly proportional to how far apart they are on their chromosome. The recombination frequency is therefore a measure of the distance between two genes.

If you work out the recombination frequencies for different pairs of genes, you can construct a map of their relative positions on the chromosome.

#### The LOD score

The LOD score often used for linkage analysis in human populations, and also in animal and plant populations.

Computerized LOD score analysis is a simple way to analyze complex family pedigrees in order to determine the linkage between Mendelian traits (or between a trait and a marker, or two markers).

The method briefly, works as follows:

Establish a pedigree

Make a number of estimates of recombination frequency

Calculate a LOD score for each estimate

The estimate with the highest LOD score will be considered the best estimate

The LOD score is calculated as follows:

LOD = Z = Log10 probability of birth sequence with a given linkage probability of birth sequence with no linkage

By convention, a LOD score greater than 3.0 is considered evidence for linkage.

On the other hand, a LOD score less than -2.0 is considered evidence to exclude linkage.

#### **Physical Mapping**

A map generated by genetic techniques is rarely sufficient for directing the sequencing phase

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of a genome project.

This is for two reasons:

The resolution of a genetic map depends on the number of crossovers that have been scored.

Genetic maps have limited accuracy.

Relies upon observable experimental outcomes

hybridization

amplification

May or may not have a distance measure.

#### Physical mapping techniques

Restriction mapping, which locates the relative positions on a DNA molecule of the recognition sequences for restriction endonucleases.

Fluorescentin situhybridization (FISH), in which marker locations are mapped by hybridizing a probe containing the marker to intact chromosomes.

Sequence tagged site (STS) mapping, in which the positions of short sequences are mapped by PCR and/or hybridization analysis of genome fragments.

#### The basic methodology for restriction mapping

The simplest way to construct a restriction map is to compare the fragment sizes produced when a DNA molecule is digested with two different restriction enzymes that recognize different target sequences.

Restriction mapping is a method used to map an unknown segment of DNA by breaking it into pieces and then identifying the locations of the breakpoints.

This method relies upon the use of proteins called restriction enzymes, which can cut, or digest, DNA molecules at short, specific sequences called restriction sites.

After a DNA segment has been digested using a restriction enzyme, the resulting fragments can be examined using a laboratory method called gel electrophoresis, which is used to separate pieces of DNA according to their size.

One common method for constructing a restriction map involves digesting the unknown DNA sample in three ways.

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Here, two portions of the DNA sample are individually digested with different restriction enzymes, and a third portion of the DNA sample is double-digested with both restriction enzymes at the same time.

Next, each digestion sample is separated using gel electrophoresis, and the sizes of the DNA fragments are recorded. The total length of the fragments in each digestion will be equal.

However, because the length of each individual DNA fragment depends upon the positions of its restriction sites, each restriction site can be mapped according to the lengths of the fragments.

The information from the double-digestion is particularly useful for correctly mapping the sites. The final drawing of the DNA segment that shows the positions of the restriction sites is called a restriction map.

#### **Limitations of Restriction mapping**

Restriction mapping is more applicable to small rather than large molecules, with the upper limit for the technique depending on the frequency of the restriction sites in the molecule being mapped.

In practice, if a DNA molecule is less than 50 kb in length it is usually possible to construct an unambiguous restriction map for a selection of enzymes with six-nucleotide recognition sequences.

The limitations of restriction mapping can be eased slightly by choosing enzymes expected to have infrequent cut sites (rare cutter) in the target DNA molecule.

#### **Rare cutters**

These rare cutters' fall into two categories:

Enzymes with seven- or eight-nucleotide recognition sequences

Enzymes whose recognition sequences contain motifs that are rare in the target DNA

#### Fluorescence in situ hybridization (FISH)

It is a kind of cytogenetic technique which uses fluorescent probes binding parts of the chromosome to show a high degree of sequence complementarity. Fluorescence microscopy can be used to find out where the fluorescent probe bound to the chromosome.

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This technique provides a novel way for researchers to visualize and map the genetic material in an individual cell, including specific genes or portions of genes.

It is an important tool for understanding a variety of chromosomal abnormalities and other genetic mutations. Different from most other techniques used for chromosomes study, FISH has no need to be performed on cells that are actively dividing, which makes it a very versatile procedure.

#### Methodology

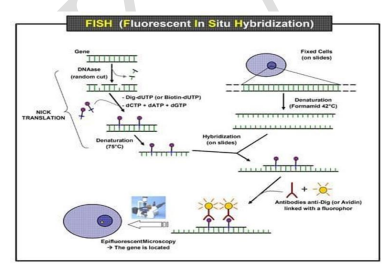
FISH is useful for example, to help a researcher identify where a particular gene falls within an individual's chromosomes. Here's how it works:

Make a probe complementary to the known sequence. When making the probe, label it with a fluorescent marker, e.g. fluorescein, by incorporating nucleotides that have the marker attached to them.

Put the chromosomes on a microscope slide and denature them.

Denature the probe and add it to the microscope slide, allowing the probe hybridize to its complementary site.

Wash off the excess probe and observe the chromosomes under a fluorescent microscope. The probe will show as one or more fluorescent signals in the microscope, depending on how many sites it can hybridize to.



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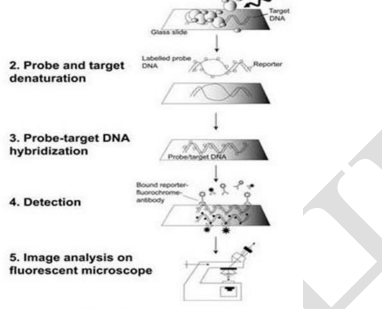


Fig. The five basic steps of FISH.

#### **Probes used in FISH**

Generally, researchers use three different types of FISH probes, each of which has a different application:

#### Locus specific probes:

It binds to a particular region of a chromosome. This type of probe is useful when researchers have isolated a small portion of a gene and want to determine on which chromosome the gene is located.

#### Alphoid or centromeric repeat probes:

They are generated from repetitive sequences found in the middle of each chromosome. Researchers use these probes to determine whether an individual has the correct number of chromosomes. These probes can also be used in combination with "locus specific probes" to determine whether an individual is missing genetic material from a particular chromosome.

#### Whole chromosome probes

They are actually collections of smaller probes, each of which binds to a different sequence

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along the length of a given chromosome. Using multiple probes labeled with a mixture of different fluorescent dyes, scientists are able to label each chromosome in its own unique color. The resulting full-color map of the chromosome is known as a spectral karyotype. Whole chromosome probes are particularly useful for examining chromosomal abnormalities, for example, when a piece of one chromosome is attached to the end of another chromosome.

#### Applications

FISH is widely used for several diagnostic applications: identification of numerical and structural abnormalities,

Characterization of marker chromosomes, monitoring the effects of therapy, detection of minimal residual disease

Ttracking the origin of cells after bone marrow transplantation, identification of regions of deletion or amplification,

Detection of chromosome abnormalities in non-dividing or terminally differentiated cells, determination of lineage involvement of clonal cells, etc.

Moreover it has many applications in research: identification of non-random chromosome rearrangements, identification of translocation molecular breakpoint, identification of commonly deleted regions, gene mapping, characterization of somatic cells hybrids, identification of amplified genes, study the mechanism of rearrangements.

FISH is also used to compare the genomes of two biological species to deduce evolutionary relationships.

#### Sequence-tagged site (STS)

It is a short region along the genome (200 to 300 bases long) whose exact sequence is found nowhere else in the genome.

The uniqueness of the sequence is established by demonstrating that it can be uniquely amplified by the PCR.

The DNA sequence of an STS may contain repetitive elements, sequences that appear elsewhere in the genome, but as long as the sequences at both ends of the site are unique, unique DNA primers complementary to those ends can be synthesized, the region amplified

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using PCR, and the specificity of the reaction demonstrated by gel electrophoresis of the amplified product.

#### **Applications of STS**

STSs are very helpful for detecting microdeletions in some genes. For example, some STSs can be used in screening by PCR to detect microdeletions in Azoospermia (AZF) genes in infertile men.

Identification of genes in elephants could provide additional information for evolutionary studies and for evaluating genetic diversity in existing elephant populations.

Sequence tagged sites (STSs) were identified in the Asian and the African elephant for the following genes: melatonin receptor 1a (MTNR1A), retinoic acid receptor beta (RARB), and leptin receptor.

Map resources

#### Assembly

A database providing information on the structure of assembled genomes, assembly names and other meta-data, statistical reports, and links to genomic sequence data.

#### BioProject (formerly Genome Project)

A collection of genomics, functional genomics, and genetics studies and links to their resulting datasets. This resource describes project scope, material, and objectives and provides a mechanism to retrieve datasets that are often difficult to find due to inconsistent annotation, multiple independent submissions, and the varied nature of diverse data types which are often stored in different databases.

#### **CloneDB** (formerly Clone Registry)

A database that integrates information about clones and libraries, including sequence data, map positions and distributor information.

#### **Database of Genome Survey Sequences (dbGSS)**

A division of GenBank that contains short single-pass reads of genomic DNA. dbGSS can be

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searched directly through the Nucleotide GSS Database.

#### Database of Genomic Structural Variation (dbVar)

The dbVar database has been developed to archive information associated with large scale genomic variation, including large insertions, deletions, translocations and inversions. Inaddition to archiving variation discovery, dbVar also stores associations of defined variants with phenotype information.

#### Genome

Contains sequence and map data from the whole genomes of over 1000 organisms. The genomes represent both completely sequenced organisms and those for which sequencing is in progress. All three main domains of life (bacteria, archaea, and eukaryota) are represented, as well as many viruses, phages, viroids, plasmids, and organelles.

#### Genome Reference Consortium (GRC)

The Genome Reference Consortium (GRC) maintains responsibility for the human and mouse reference genomes. Members consist of The Genome Center at Washington University, the Wellcome Trust Sanger Institute, the European Bioinformatics Institute (EBI) and the National Center for Biotechnology Information (NCBI). The GRC works to correct misrepresented loci and to close remaining assembly gaps. In addition, the GRC seeks to provide alternate assemblies for complex or structurally variant genomic loci. At the GRC website (http://www.genomereference.org), the public can view genomic regions currently under review, report genome-related problems and contact the GRC.

#### **HIV-1. Human Protein Interaction Database**

A database of known interactions of HIV-1 proteins with proteins from human hosts. It provides annotated bibliographies of published reports of protein interactions, with links to the corresponding PubMed records and sequence data.

#### Influenza Virus

A compilation of data from the NIAID Influenza Genome Sequencing Project and GenBank. It provides tools for flu sequence analysis, annotation and submission to GenBank. This resource also has links to other flu sequence resources, and publications and general information about flu viruses.

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#### **NCBI Pathogen Detection Project**

A project involving the collection and analysis of bacterial pathogen genomic sequences originating from food, environmental and patient isolates. Currently, an automated pipeline clusters and identifies sequences supplied primarily by public health laboratories to assist in the investigation of foodborne disease outbreaks and discover potential sources of food contamination.

#### Nucleotide Database

A collection of nucleotide sequences from several sources, including GenBank, RefSeq, the Third Party Annotation (TPA) database, and PDB. Searching the Nucleotide Database will yield available results from each of its component databases.

#### PopSet

Database of related DNA sequences that originate from comparative studies: phylogenetic, population, environmental and, to a lesser degree, mutational. Each record in the database is a set of DNA sequences. For example, a population set provides information on genetic variation within an organism, while a phylogenetic set may contain sequences, and their alignment, of a single gene obtained from several related organisms.

#### **Probe**

A public registry of nucleic acid reagents designed for use in a wide variety of biomedical research applications, together with information on reagent distributors, probe effectiveness, and computed sequence similarities.

#### **Retrovirus Resources**

A collection of resources specifically designed to support the research of retroviruses, including a genotyping tool that uses the BLAST algorithm to identify the genotype of a query sequence; an alignment tool for global alignment of multiple sequences; an HIV-1 automatic sequence annotation tool; and annotated maps of numerous retroviruses viewable in GenBank, FASTA, and graphic formats, with links to associated sequence records.

#### SARS CoV

A summary of data for the SARS coronavirus (CoV), including links to the most recent

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sequence data and publications, links to other SARS related resources, and a pre-computed alignment of genome sequences from various isolates.

#### Sequence Read Archive (SRA)

The Sequence Read Archive (SRA) stores sequencing data from the next generation of sequencing platforms including Roche 454 GS System®, Illumina Genome Analyzer®, Life Technologies AB SOLiD System®, Helicos Biosciences Heliscope®, Complete Genomics®, and Pacific Biosciences SMRT®.

#### **Trace Archive**

A repository of DNA sequence chromatograms (traces), base calls, and quality estimates for single-pass reads from various large-scale sequencing projects.

#### Viral Genomes

A wide range of resources, including a brief summary of the biology of viruses, links to viral genome sequences in Entrez Genome, and information about viral Reference Sequences, a collection of reference sequences for thousands of viral genomes.

#### Virus Variation

An extension of the Influenza Virus Resource to other organisms, providing an interface to download sequence sets of selected viruses, analysis tools, including virus-specific BLAST pages, and genome annotation pipelines.

#### FTP: Genome

This site contains genome sequence and mapping data for organisms in Entrez Genome. The data are organized in directories for single species or groups of species. Mapping data are collected in the directory MapView and are organized by species. See the README file in the root directory and the README files in the species subdirectories for detailed information.

#### **FTP: Genome Mapping Data**

Contains directories for each genome that include available mapping data for current and previous builds of that genome.

#### FTP: RefSeq

This site contains all nucleotide and protein sequence records in the Reference Sequence

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(RefSeq) collection. The ""release"" directory contains the most current release of the complete collection, while data for selected organisms (such as human, mouse and rat) are available in separate directories. Data are available in FASTA and flat file formats. See the README file for details.

#### FTP: SKY/M-Fish and CGH Data

This site contains SKY-CGH data in ASN.1, XML and EasySKYCGH formats. See the skycghreadme.txt file for more information.

#### FTP: Sequence Read Archive (SRA) Download Facility

This site contains next-generation sequencing data organized by the submitted sequencing project.

#### FTP: Trace Archive

This site contains the trace chromatogram data organized by species. Data include chromatogram, quality scores, FASTA sequences from automatic base calls, and other ancillary information in tab-delimited text as well as XML formats. See the README file for details.

#### FTP: Whole Genome Shotgun Sequences

This site contains whole genome shotgun sequence data organized by the 4-digit project code. Data include GenBank and GenPept flat files, quality scores and summary statistics. See the README.genbank.wgs file for more information.

#### A haplotype map of the human genome

The planned Haplotype Map is the next logical step in mobilizing tools for gene discovery.

The most common type of variation in the human genome is the single nucleotide polymorphism or SNP, a single-base difference at a genetic locus from person to person.

Millions of SNPs have been found, making it imperative that we find efficient and costeffective ways for using them.

The Haplotype Map is based on the recognition that the development of genetic variation from ancestral chromosomes has not proceeded uniformly across the genome.

Rather, there appear to be regions in which recombination is more likely to occur, thus

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shuffling the genetic deck at those points.

There are other regions where is it less likely to occur, leaving relatively large blocks intact. These blocks or haplotypes can be identified by a small number of SNPs.

Wise use of genetic markers will be enhanced by knowing the boundaries of these blocks. To be sure, a clear haplotype structure may not be apparent everywhere in the genome, but knowledge of the haplotype structure of the genome will speed the search for loci that confer disease risk.

The Hap Map should help us use genetic markers wisely, to speed up (and to make affordable) association studies based on candidate genes and ultimately, whole-genome association studies. Without the Hap Map, the choice of markers for association studies will remain more or less a matter of guesswork.

#### **Association Mapping**

Association mapping (genetics), also known as "linkage disequilibrium mapping", is a method of mapping quantitative trait loci (QTLs) that takes advantage of historic linkage disequilibrium to link phenotypes (observable characteristics) to genotypes (the genetic constitution of organisms), uncovering genetic associations.

Association mapping is based on the idea that traits that have entered a population only recently will still be linked to the surrounding genetic sequence of the original evolutionary ancestor, or in other words, will more often be found within a given haplotype, than outside of it.

It is most often performed by scanning the entire genome for significant associations between a panel of SNPs (which, in many cases are spotted onto glass slides to create "SNP chips") and a particular phenotype.

These associations must then be independently verified in order to show that they either (a) contribute to the trait of interest directly, or (b) are linked to/ in linkage disequilibrium with a quantitative trait locus (QTL) that contributes to the trait of interest.

The advantage of association mapping is that it can map quantitative traits with high resolution in a way that is statistically very powerful.

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Association mapping, however, also requires extensive knowledge of SNPs within the genome of the organism of interest, and is therefore difficult to perform in species that have not been well studied or do not have well-annotated genomes.

#### **Benefits of Genetic Mapping**

The techniques developed for genetic mapping have had great impact on the life sciences, and particularly in medicine. But genetic mapping technologies also have useful applications in other fields. Commercialization of the fruits of genomics research promises immense opportunities for industry. A round-up of genetic mapping applications would include (but not be limited to) the areas below.

#### Medicine

Scientists have become more proficient in genetic sequencing - the detailed genetic maps that help locate the risk genes for a host of genetic diseases. The ability to investigate the root cause of diseases may one day allow medical researchers to develop strategies to avoid the environmental conditions that serve as triggers to disease, formulate customized drugs, and techniques for gene therapy.

#### **Agricultural Applications**

Knowledge of the genetic maps of plants and animals leads to the development of agricultural crops and animal breeds that are more nutritious, productive and can better resist diseases, insects and drought. Researchers can breed special plants that help clean up wastes that are difficult to break down.

#### **Energy and the Environment**

Genetic maps of microbes enable researchers to harness the power of bacteria for producing energy from bio-fuels, reducing toxic waste, and developing environment-friendly products and industrial processes.

#### Forensics

We are already familiar with the use of genetic mapping in crime investigations, paternity tests, and identification. The technique can also be used in organ transplants to achieve better matches between recipients and donors, thus minimizing the risks of complications and

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maximizing the use of donated healthy organs, a scarce resource. For more delectable applications, genetic mapping can authenticate the origins of consumer goods like caviar, fruits, and wine or the pedigree of livestock and animal breeds.

#### **Genetic Markers**

Genetic markers are useful in identification of various genetic variations. The development of DNA-based genetic markers has had a revolutionary impact on genetic studies.

With DNA markers, it is theoretically possible to observe and exploit genetic variation in the entire genome. These markers can be used to study the evolutionary relationships among individuals.

Popular genetic markers include allozymes, mitochondrial DNA, RFLP, RAPD, AFLP, microsatellite, SNP, and EST markers.

The application of DNA markers has allowed rapid progress in investigations of genetic variability and inbreeding, parentage assignments, species and strain identification, and the construction of high-resolution genetic linkage maps for aquaculture species.

The advent of next-generation sequencing (NGS) has revolutionized genomic and transcriptomic approaches to biology.

The new sequencing tools are also valuable for the discovery, validation and assessment of genetic markers in populations. This review focuses on importance and uses of genetic markers with advent of modern technologies.

#### Minisatellite

Minisatellites have been found in association with important features of human genome biology such as gene regulation, chromosomal fragile sites, and imprinting. Our knowledge of minisatellite biology has greatly increased in the past 10 years owing to the identification and careful analysis of human hypermutable minisatellites, experimental models in yeast, and recent in vitro studies of minisatellite recombination properties.

In parallel, minisatellites have been put forward as potential biomarkers for the monitoring of genotoxic agents such as ionizing radiation.

We summarize and discuss recent observations on minisatellites. In addition we take

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advantage of recent whole chromosome sequence data releases to provide a unifying view which may facilitate the annotation of tandem repeat sequences.

Minisatellites are usually defined as the repetition in tandem of a short (6- to 100-bp) motif spanning 0.5 kb to several kilobases.

Although the first examples described 20 years ago were of human origin, (Wyman and White 1980), similar DNA structures have been found in many organisms including bacteria.

Comparisons of the repeat units in classical minisatellites led early on to the notion of consensus or core sequences, which exhibit some similarities with the  $\chi$  sequence of  $\lambda$  phage (GCTGTGG). In general, the majority of classical minisatellites are GC rich, with a strong strand asymmetry.

#### Microsatellite

Microsatellites or Single Sequence Repeats (SSRs) are extensively employed in plant genetics studies, using both low and high throughput genotyping approaches.

Motivated by the importance of these sequences over the last decades this review aims to address some theoretical aspects of SSRs, including definition, characterization and biological function.

The methodologies for the development of SSR loci, genotyping and their applications as molecular markers are also reviewed.

Finally, two data surveys are presented. The first was conducted using the main database of Web of Science, prospecting for articles published over the period from 2010 to 2015, resulting in approximately 930 records.

The second survey was focused on papers that aimed at SSR marker development, published in the American Journal of Botany's Primer Notes and Protocols in Plant Sciences (over 2013 up to 2015), resulting in a total of 87 publications.

This scenario confirms the current relevance of SSRs and indicates their continuous utilization in plant science.

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Difference between whet osatelitte and winnsatelitte	
Minisatellites	Microsatellites
a) Hypervriable family	Repeat size: 1-4 bp
Repeat size:10-60bp	Total sites: Less than 1000 bp
Total Size:1000-20,000 bp	
b) Telomeric family:	
Repeat size: 6 bp	
Total size: 1000-20000 bp	
Share a common core sequence (motif)	Repeats A and CA are the most common
GGGCAGGANG (where N is any base), dispersed,	Dispersed throughout genome.
VNTRs usually TTAGGG and repeated about a	
thousand times protects chromosome ends.	
Complexity of Array: Heterogeneous	Complexity of Array: Homogeneous

Difference between Microsatellite and Minisatellite

## Sequence-Tagged Site (STS)

It is a relatively short, easily PCR-amplified sequence (200 to 500 bp) which can be specifically amplified by PCR and detected in the presence of all other genomic sequences and whose location in the genome is mapped.

The STS concept was introduced by Olson et al (1989). In assessing the likely impact of the Polymerase Chain Reaction (PCR) on human genome research, they recognized that single-copy DNA sequences of known map location could serve as markers for genetic and physical mapping of genes along the chromosome.

The advantage of STSs over other mapping landmarks is that the means of testing for the presence of a particular STS can be completely described as information in a database: anyone who wishes to make copies of the marker would simply look up the STS in the database, synthesize the specified primers, and run the PCR under specified conditions to amplify the STS from genomic DNA.

STS-based PCR produces a simple and reproducible pattern on agarose or polyacrylamide gel.

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In most cases STS markers are co-dominant, i.e., allow heterorozygotes to be distinguished from the two homozygotes.

The DNA sequence of an STS may contain repetitive elements, sequences that appear elsewhere in the genome, but as long as the sequences at both ends of the site are unique and conserved, researches can uniquely identify this portion of genome using tools usually present in any laboratory.

Thus, in broad sense, STS include such markers as microsatellites (SSRs, STMS or SSRPs), SCARs, CAPs, and ISSRs.

### **Expressed Sequence Tag (EST)**

It is a short stretch of DNA sequence that is used to identify an expressed gene. Although EST sequences are usually only 200 to 500 nucleotides in length, this is generally sufficient to identify the full-length complementary DNA (cDNA).

ESTs are generated by sequencing a single segment of random clones from a cDNA library. A single sequencing reaction and automation of DNA isolation, sequencing, and analysis have allowed the rapid determination of many ESTs.

Now, the majority of the sequences in sequence databases are ESTs. Although most ESTs have been isolated from humans, a large number of ESTs have been isolated from model organisms, such as Caenorhabditis elegans, Drosophila, rice, and Arabidopsis.

ESTs are also being isolated from more exotic organisms, such as Entamoeba histolytica and Leishmania major promastigotes .

ESTs have numerous uses, from genetic mapping to analyzing gene expression, and the number of ESTs isolated from different organisms will continue to rise rapidly.

### Single nucleotide polymorphisms

It is frequently called SNPs (pronounced "snips"), are the most common type of genetic variation among people. Each SNP represents a difference in a single DNA building block, called a nucleotide. For example, a SNP may replace the nucleotide cytosine (C) with the nucleotide thymine

(T) in a certain stretch of DNA.

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SNPs occur normally throughout a person"s DNA. They occur once in every 300 nucleotides on average, which means there are roughly 10 million SNPs in the human genome.

Most commonly, these variations are found in the DNA between genes. They can act as biological markers, helping scientists locate genes that are associated with disease.

When SNPs occur within a gene or in a regulatory region near a gene, they may play a more direct role in disease by affecting the gene"s function.

Most SNPs have no effect on health or development. Some of these genetic differences, however, have proven to be very important in the study of human health.

Researchers have found SNPs that may help predict an individual"s response to certain drugs, susceptibility to environmental factors such as toxins, and risk of developing particular diseases.

SNPs can also be used to track the inheritance of disease genes within families. Future studies will work to identify SNPs associated with complex diseases such as heart disease, diabetes, and cancer.

### Possible Questions 1 Mark questions

1. Microsatellites are a) frequently found in bacterial genomes b) always smaller than 10 bp d) movable DNA elements c) used as DNA markers 2. Molecular markers are used to construct a) Chromosome maps b) cytogenetic maps c) physical maps d) geographic maps 3. The variation in number of tandem repeats between two or more individuals is called a) VNTRs b) RFLP c) SSRs d) AFLP 4. The variant fragment that distinguish one individual from another one is called a) variant fragment b) marking fragment c) differing fragment d) variable repeats 5. Which of these is a key characterisitic of a molecular marker? a) It is a known gene b) It is located at a known site on the chromosome c) It is only useful for linkage and physical mapping studies d) positional analysis

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6. A polymorphism is ----a) Any change in the DNA sequence b) The most common variation of a gene or marker sequence c) The least common vaariation of a gene or marker sequence d) Variation of gene or marker sequence present in > 1% of the population 7. The shotgun method ----a) is used in analyzing transcriptomes b) requires computers c) is normally used with large genomes d) is more accurate than clone contig method 8. A monomorphic DNA segment is a) A segment of DNA that exists in many forms in the population b) A segment of DNA that controls a single gene function c) A segment of DNA inherited in a dominant fashion d) A segment of DNA shared by over 99 % of the population 9. Which of these describes a contig? a) A complete genomic library including overlapping clones b) A complete mRNA library c) A chromosome specific library of overlapping clones d) An ordered genomci library 10. The variation in the restriction DNA fragment lengths between individuals of a species is called a) Retriction fragment length polymorphism b) RAPD c) AFLP d) simple sequence repeats 11. All the following statements are true regarding RFLP and RAPD except a) RAPD is a quick method compared to RFLP

- c) Species specific primers are required for RAPD
- b) RFLP is more relible than RAPD
- d) Radioactive probes are not required in RAPD

# **2 Marks question**

1. Write a short note on 'mini satellite.

- 2. What are ESTs?
- 3. What are the types of genome maps?

4. What is RFLP?

5. What is SNP?

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#### 6/8 Marks questions

- 1. What are genetic markers? Explain with an example.
- 2. Describe genetic mapping techniques and their applications in detail.
- 3. Describe physical mapping techniques and their applications in detail.
- 4. Briefly explain uses of SNP.
- 5. How do 'mini satellites' differ from 'micro satellites'?

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### UNIT - IV

### **SYLLABUS**

**Protein and structure determination:** Introduction to protein structure, Chemical properties of proteins. Physical interactions that determine the property of proteins. Short-range interactions, electrostatic forces, van der Waal interactions, hydrogen bonds, Hydrophobic interactions. Determination of sizes - Sedimentation analysis, gel filteration, Native PAGE, SDS-PAGE. Determination of covalent structures – Edman degradation.

### **Three-dimensional Structures of proteins**

The properties of a protein are largely determined by its three-dimensional structure. One might naively suppose that since proteins are all composed of the same 20 types of amino acid residues, they would be more or less alike in their properties. Indeed, denatured (unfolded) proteins have rather similar characteristics, a kind of homogeneous "average" of their randomly dangling side chains. How- ever, the three-dimensional structure of a native (physio- logically folded) protein is specified by its primary structure, so that it has a unique set of characteristics. we shall discuss the structural features of proteins, the forces that hold them together, and their hierarchical organization to form complex structures. This will form the basis for understanding the structure–func- tion relationships necessary to comprehend the biochemical roles of proteins.

### Amino acids

Amino acids contain both amino and carboxylic acid functional groups. (In biochemistry, the term amino acid is used when referring to those amino acids in which the amino and carboxylate functionalities are attached to the same carbon, plus proline which is not actually an amino acid). Modified amino acids are sometimes observed in proteins; this is usually the result of enzymatic modification after translation (protein synthesis). For example, phosphorylation of serine by kinases and dephosphorylation by phosphatases is an important control mechanism in the cell cycle. Only two amino acids other than the standard twenty are known to be incorporated into proteins during translation, in certain organisms:

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- Selenocysteine is incorporated into some proteins at a UGA codon, which is normally a stop codon.
- Pyrrolysine is incorporated into some proteins at a UAG codon. For instance, in some methanogens in enzymes that are used to produce methane.

Besides those used in protein synthesis, other biologically important amino acids include carnitine (used in lipid transport within a cell), ornithine, GABA and taurine.

### **Protein structure**

The particular series of amino acids that form a protein is known as that protein's primary structure. This sequence is determined by the genetic makeup of the individual. It specifies the order of side-chain groups along the linear polypeptide "backbone".

Proteins have two types of well-classified, frequently occurring elements of local structure defined by a particular pattern of hydrogen bonds along the backbone: alpha helix and beta sheet. Their number and arrangement is called the secondary structure of the protein. Alpha helices are regular spirals stabilized by hydrogen bonds between the backbone CO group (carbonyl) of one amino acid residue and the backbone NH group (amide) of the i+4 residue. The spiral has about 3.6 amino acids per turn, and the amino acid side chains stick out from the cylinder of the helix. Beta pleated sheets are formed by backbone hydrogen bonds between individual beta strands each of which is in an "extended", or fully stretched-out, conformation. The strands may lie parallel or antiparallel to each other, and the side-chain direction alternates above and below the sheet. Hemoglobin contains only helices, natural silk is formed of beta pleated sheets, and many enzymes have a pattern of alternating helices and beta-strands. The secondary-structure elements are connected by "loop" or "coil" regions of non-repetitive conformation, which are sometimes quite mobile or disordered but usually adopt a well-defined, stable arrangement. The overall, compact, 3D structure of a protein is termed its tertiary structure or its "fold". It is formed as result of various attractive forces like hydrogen bonding, disulfide bridges, hydrophobic interactions, hydrophilic interactions, van der Waals force etc. When two or more polypeptide chains (either of identical or of different sequence) cluster to form a protein, quaternary structure of protein is formed. Quaternary structure is an attribute of polymeric

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(same-sequence chains) or heteromeric (different-sequence chains) proteins like hemoglobin, which consists of two "alpha" and two "beta" polypeptide chains.

### Apoenzymes

An apoenzyme (or, generally, an apoprotein) is the protein without any small-molecule cofactors, substrates, or inhibitors bound. It is often important as an inactive storage, transport, or secretory form of a protein. This is required, for instance, to protect the secretory cell from the activity of that protein. Apoenzymes becomes active enzymes on addition of a cofactor. Cofactors can be either inorganic (e.g., metal ions and iron-sulfur clusters) or organic compounds, (e.g., flavin and heme). Organic cofactors can be either prosthetic groups, which are tightly bound to an enzyme, or coenzymes, which are released from the enzyme's active site during the reaction.

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Name, Three-Latter Symbol, and One-Latter Symbol	Structural I Formula"	Residue Mass (D) <sup>b</sup>	Occurrence	рК, a-COOH <sup>4</sup>	рК <sub>2</sub> α-NH <sub>3</sub> <sup>+4</sup>	pK <sub>n</sub> Side Chain <sup>d</sup>
Amino acids with non						
Glycine Gly G	соо- н-с-н мн <u>і</u>	57.D	7.1	2.35	9.78	
Alainine Ala A	COO- H-C-CH <sub>a</sub> NH <sup>+</sup> <sub>a</sub>	71.1	8.3	2.35	9.87	
Valine Val V	H-C-CH NH <sup>+</sup> <sub>h</sub> CH <sub>a</sub>	99.1	6.9	2.29	9.74	
Leucine Leu L	н-С-СH <sub>2</sub> -СH NH <sup>1</sup> <sub>3</sub> СH <sub>2</sub>	1132	9.7	2.33	9.74	
Isoleucine Ile I	$\begin{array}{c} coo^{-} cH_{3} \\ H_{-}c & c^{+} cH_{2} - cH_{3} \\ I_{-} H_{3} & H \end{array}$	1132	6.0	2.32	9.76	
Methionine Met M	COO <sup>-</sup> H-C-CH <sub>2</sub> -CH <sub>2</sub> -S-CH <sub>3</sub> NH <sup>1</sup> <sub>3</sub>	131.2	2.4	2.13	9.28	
Proline Pro P	$H_2 \\ COO - C_1 CH_2 \\ H H H_2 CH_2$	97.1	4.7	1.95	10.64	
Phenylalanine Phe F	H-C-CH2-	1472	3.9	2.20	9.31	
Tryptophan Trp W	H-C-CH2 NH <sup>1</sup> <sub>1</sub>	1862	1.1	2.46	9.41	
	Н					(continued)

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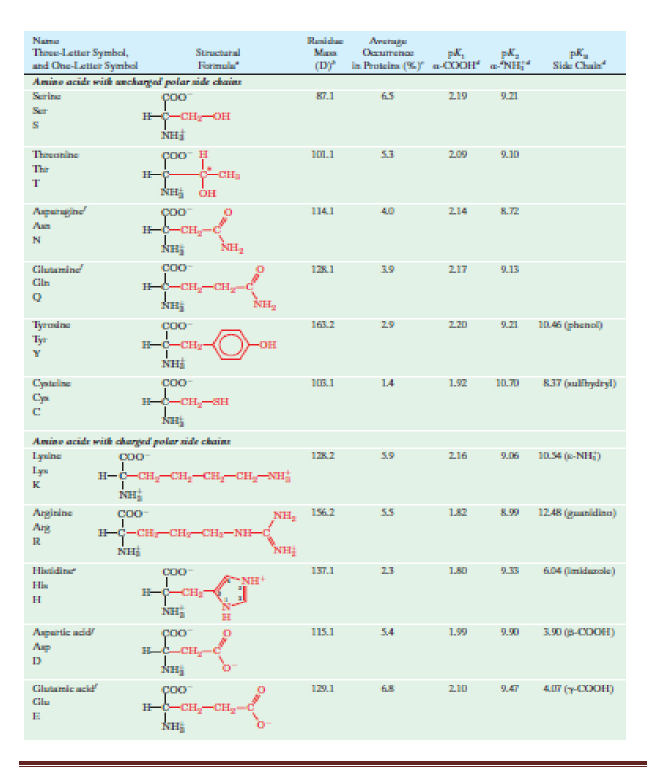
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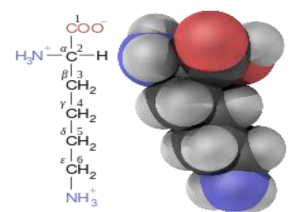
In the structure shown at the top of the page, R represents a side chain specific to each amino acid. The carbon atom next to the carboxyl group (which is therefore numbered 2 in the carbon chain starting from that functional group) is called the  $\alpha$ -carbon. Amino acids containing an amino group bonded directly to the alpha carbon are referred to as *alpha amino acids*. These include amino acids such as proline which contain secondary amines, which used to be often referred to as "imino acids".

### Isomerism

The alpha amino acids are the most common form found in nature, but only when occurring in the L-isomer. The alpha carbon is a chiral carbon atom, with the exception of glycine which has two indistinguishable hydrogen atoms on the alpha carbon. Therefore, all alpha amino acids but glycine can exist in either of two enantiomers, called L or D amino acids, which are mirror images of each other (see also Chirality). While L-amino acids represent all of the amino acids found in proteins during translation in the ribosome, D-amino acids are found in some proteins produced by enzyme posttranslational modifications after translation and translocation to the endoplasmic reticulum, as in exotic sea-dwelling organisms such as cone snails. They are also abundant components of the peptidoglycan cell walls of bacteria,<sup>[36]</sup> and Dserine may act as a neurotransmitter in the brain. D-amino acids are used in racemic crystallography to create centrosymmetric crystals, which (depending on the protein) may allow for easier and more robust protein structure determination. The L and D convention for amino acid configuration refers not to the optical activity of the amino acid itself but rather to the optical activity of the isomer of glyceraldehyde from which that amino acid can, in theory, be synthesized (D-glyceraldehyde is dextrorotatory; L-glyceraldehyde is levorotatory). In alternative fashion, the (S) and (R) designators are used to indicate the absolute stereochemistry. Almost all of the amino acids in proteins are (S) at the  $\alpha$  carbon, with cysteine being (R) and glycine non-chiral.<sup>[39]</sup> Cysteine has its side chain in the same geometric position as the other amino acids, but the R/S terminology is reversed because of the higher atomic number of sulfur compared to the carboxyl oxygen gives the side chain a higher priority, whereas the atoms in most other side chains give them lower priority.

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Side chains

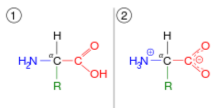


### Lysine with carbon atoms labeled by position

In amino acids that have a carbon chain attached to the  $\alpha$ -carbon (such as lysine, shown to the right) the carbons are labeled in order as  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$ , and so on. In some amino acids, the amine group is attached to the  $\beta$  or  $\gamma$ -carbon, and these are therefore referred to as *beta* or *gamma amino acids*. Amino acids are usually classified by the properties of their side chain into four groups. The side chain can make an amino acid a weak acid or a weak base, and a hydrophile if the side chain is polar or a hydrophobe if it is nonpolar. The chemical structures of the 22 standard amino acids, along with their chemical properties, are described more fully in the article on these proteinogenic amino acids. The phrase "branched-chain amino acids" or BCAA refers to the amino acids having aliphatic side chains that are non-linear; these are leucine, isoleucine, and valine. Proline is the only proteinogenic amino acid containing a secondary amine at this position.<sup>[34]</sup> In chemical terms, proline is, therefore, an imino acid, since it lacks a primary amino group,<sup>[41]</sup> although it is still classed as an amino acid in the current biochemical nomenclature,<sup>[42]</sup> and may also be called an "N-alkylated alpha-amino acid".

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### Zwitterions



The  $\alpha$ -carboxylic acid group of amino acids is a weak acid, meaning that it releases a hydron (such as a proton) at moderate pH values. In other words, carboxylic acid groups  $(-CO_2H)$  can be deprotonated to become negative carboxylates  $(-CO_2^-)$ . The negatively charged carboxylate ion predominates at pH values greater than the pKa of the carboxylic acid group (mean for the 20 common amino acids is about 2.2, see the table of amino acid structures above). In a complementary fashion, the  $\alpha$ -amine of amino acids is a weak base, meaning that it accepts a proton at moderate pH values. In other words,  $\alpha$ -amino groups (NH<sub>2</sub>-) can be protonated to become positive  $\alpha$ -ammonium groups (<sup>+</sup>NH<sub>3</sub>-). The positively charged  $\alpha$ ammonium group predominates at pH values less than the pKa of the  $\alpha$ -ammonium group (mean for the 20 common  $\alpha$ -amino acids is about 9.4). Because all amino acids contain amine and carboxylic acid functional groups, they share amphiprotic properties. Below pH 2.2, the predominant form will have a neutral carboxylic acid group and a positive α-ammonium ion (net charge +1), and above pH 9.4, a negative carboxylate and neutral  $\alpha$ -amino group (net charge -1). But at pH between 2.2 and 9.4, an amino acid usually contains both a negative carboxylate and a positive  $\alpha$ -ammonium group, as shown in structure (2) on the right, so has net zero charge. This molecular state is known as a zwitterion, from the German Zwitter meaning hermaphrodite or hybrid. The fully neutral form (structure (1) on the left) is a very minor species in aqueous solution throughout the pH range (less than 1 part in  $10^7$ ). Amino acids exist as zwitterions also in the solid phase, and crystallize with salt-like properties unlike typical organic acids or amines.

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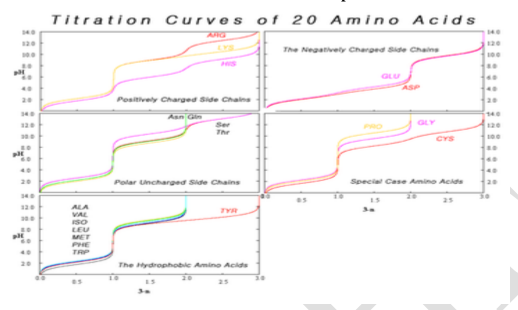
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### **Isoelectric point**



Composite of titration curves of twenty proteinogenic amino acids grouped by side chain category. The variation in titration curves when the amino acids can be grouped by category. With the exception of tyrosine, using titration to distinguish among hydrophobic amino acids is problematic. At pH values between the two pKa values, the zwitterion predominates, but coexists in dynamic equilibrium with small amounts of net negative and net positive ions. At the exact midpoint between the two pKa values, the trace amount of net negative and trace of net positive ions exactly balance, so that average net charge of all forms present is zero. This pH is known as the isoelectric point pI, so  $pI = \frac{1}{2}(pKa_1 + pKa_2)$ . The individual amino acids all have slightly different pKa values, so have different isoelectric points. For amino acids with charged side chains, the pKa of the side chain is involved. Thus for Asp, Glu with negative side chains, pI =  $\frac{1}{2}(pKa_1 + pKa_R)$ , where pKa<sub>R</sub> is the side chain pKa. Cysteine also has potentially negative side chain with  $pKa_R = 8.14$ , so pI should be calculated as for Asp and Glu, even though the side chain is not significantly charged at neutral pH. For His, Lys, and Arg with positive side chains,  $pI = \frac{1}{2}(pKa_R + pKa_2)$ . Amino acids have zero mobility in electrophoresis at their isoelectric point, although this behaviour is more usually exploited for peptides and proteins than single amino acids. Zwitterions have minimum solubility at their isoelectric point and some amino

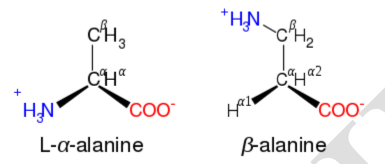
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acids (in particular, with non-polar side chains) can be isolated by precipitation from water by adjusting the pH to the required isoelectric point.

Occurrence and functions in biochemistry



 $\beta$ -alanine and its  $\alpha$ -alanine isomer

### **Proteinogenic amino acids**

Amino acids are the structural units (monomers) that make up proteins. They join together to form short polymer chains called peptides or longer chains called either polypeptides or proteins. These polymers are linear and unbranched, with each amino acid within the chain attached to two neighboring amino acids. The process of making proteins encoded by DNA/RNA genetic material is called *translation* and involves the step-by-step addition of amino acids to a growing protein chain by a ribozyme that is called a ribosome. The order in which the amino acids are added is read through the genetic code from an mRNA template, which is a RNA copy of one of the organism's genes. Twenty-two amino acids. Of these, 20 are encoded by the universal genetic code. The remaining 2, selenocysteine and pyrrolysine, are incorporated into proteins by unique synthetic mechanisms. Selenocysteine is incorporated when the mRNA being translated includes a SECIS element, which causes the UGA codon to encode selenocysteine instead of a stop codon. Pyrrolysine is used by some methanogenic archaea in enzymes that they use to produce methane. It is coded for with the codon UAG, which is normally a stop codon in other organisms.<sup>[48]</sup> This UAG codon is followed by a PYLIS downstream sequence.

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### Non-proteinogenic amino acids

Aside from the 22 proteinogenic amino acids, many *non-proteinogenic* amino acids are known. Those either are not found in proteins (for example carnitine, GABA, Levothyroxine) or are not produced directly and in isolation by standard cellular machinery (for example, hydroxyproline and selenomethionine). Non-proteinogenic amino acids that are found in proteins are formed by post-translational modification, which is modification after translation during protein synthesis. These modifications are often essential for the function or regulation of a protein. For example, the carboxylation of glutamate allows for better binding of calcium cations, and collagen contains hydroxyproline, generated by hydroxylation of proline. Another example is the formation of hypusine in the translation initiation factor EIF5A, through modification of a lysine residue. Such modifications can also determine the localization of the protein, e.g., the addition of long hydrophobic groups can cause a protein to bind to a phospholipid membrane.

### Non-standard amino acids

The 20 amino acids that are encoded directly by the codons of the universal genetic code are called *standard* or *canonical* amino acids. A modified form of methionine (*N*-formylmethionine) is often incorporated in place of methionine as the initial amino acid of proteins in bacteria, mitochondria and chloroplasts. Other amino acids are called *non-standard* or *non-canonical*. Most of the non-standard amino acids are also non-proteinogenic (i.e. they cannot be incorporated into proteins during translation), but two of them are proteinogenic, as they can be incorporated translationally into proteins by exploiting information not encoded in the universal genetic code. The two non-standard proteinogenic amino acids are selenocysteine (present in many non-eukaryotes as well as most eukaryotes, but not coded directly by DNA) and pyrrolysine (found only in some archaea and one bacterium). The incorporation of these non-standard amino acids is rare. For example, 25 human proteins include selenocysteine (Sec) in their primary structure, and the structurally characterized enzymes (selenoenzymes) employ Sec

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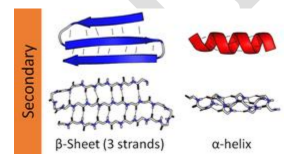
as the catalytic moiety in their active sites. Pyrrolysine and selenocysteine are encoded via variant codons. For example, selenocysteine is encoded by stop codon and SECIS element.

### **Secondary structure**

Protein secondary structure is the three dimensional form of *local segments* of proteins. The two most common secondary structural elements are alpha helices and beta sheets, though beta turns and omega loops occur as well. Secondary structure elements typically spontaneously form as an intermediate before the protein folds into its three dimensional tertiary structure. Secondary structure is formally defined by the pattern of hydrogen bonds between the amino hydrogen and carboxyl oxygen atoms in the peptide backbone. Secondary structure may alternatively be defined based on the regular pattern of backbone dihedral angles in a particular region of the Ramachandran plot regardless of whether it has the correct hydrogen bonds.

The concept of secondary structure was first introduced by Kaj Ulrik Linderstrøm-Lang at Stanford in 1952. Other types of biopolymers such as nucleic acids also possess characteristic secondary structures.

Structural features of the three major forms of protein helicesGeometry attribute $\alpha$ -helix $3_{10}$  helix $\pi$ -helixResidues per turn3.63.04.4Translation per residue1.5 Å (0.15 nm) 2.0 Å (0.20 nm) 1.1 Å (0.11 nm)Radius of helix2.3 Å (0.23 nm) 1.9 Å (0.19 nm) 2.8 Å (0.28 nm)Pitch5.4 Å (0.54 nm) 6.0 Å (0.60 nm) 4.8 Å (0.48 nm)



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The most common secondary structures are alpha helices and beta sheets. Other helices, such as the  $3_{10}$  helix and  $\pi$  helix, are calculated to have energetically favorable hydrogenbonding patterns but are rarely observed in natural proteins except at the ends of  $\alpha$  helices due to unfavorable backbone packing in the center of the helix. Other extended structures such as the polyproline helix and alpha sheet are rare in native state proteins but are often hypothesized as important protein folding intermediates. Tight turns and loose, flexible loops link the more "regular" secondary structure elements. The random coil is not a true secondary structure, but is the class of conformations that indicate an absence of regular secondary structure. Amino acids vary in their ability to form the various secondary structure elements. Proline and glycine are sometimes known as "helix breakers" because they disrupt the regularity of the  $\alpha$  helical backbone conformation; however, both have unusual conformational abilities and are commonly found in turns. Amino acids that prefer to adopt helical conformations in proteins include methionine, alanine, leucine, glutamate and lysine ("MALEK" in amino-acid 1-letter codes); by contrast, the large aromatic residues (tryptophan, tyrosine and phenylalanine) and  $C^{\beta}$ -branched amino acids (isoleucine, valine, and threenine) prefer to adopt  $\beta$ -strand conformations. However, these preferences are not strong enough to produce a reliable method of predicting secondary structure from sequence alone. Low frequency collective vibrations are thought to be sensitive to local rigidity within proteins, revealing beta structures to be generically more rigid than alpha or disordered proteins. Neutron scattering measurements have directly connected the spectral feature at ~1 THz to collective motions of the secondary structure of beta-barrel protein GFP. Hydrogen bonding patterns in secondary structures may be significantly distorted, which makes automatic determination of secondary structure difficult. There are several methods for formally defining protein secondary structure (e.g., DSSP, DEFINE, STRIDE, ScrewFit, SST).

**Protein tertiary structure** is the three dimensional shape of a protein. The tertiary structure will have a single polypeptide chain "backbone" with one or more protein secondary structures, the protein domains. Amino acid side chains may interact and bond in a number of ways. The interactions and bonds of side chains within a particular protein determine its tertiary structure. The protein tertiary structure is defined by its atomic coordinates. These coordinates

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may refer either to a protein domain or to the entire tertiary structure. A number of tertiary structures may fold into a quaternary structure.

### SECONDARY STRUCTURE

A polymer's secondary structure (2° structure) is defined as the local conformation of its backbone. For proteins, this has come to mean the specification of regular polypeptide backbone folding patterns: helices, pleated sheets, and turns. However, before we begin our discussion of these basic structural motifs, let us consider the geometrical properties of the peptide group because its understanding is prerequisite to that of any structure containing it.

### **The Peptide Group**

In the 1930s and 1940s, Linus Pauling and Robert Corey determined the X-ray structures of several amino acids and dipeptides in an effort to elucidate the structural con- straints on the conformations of a polypeptide chain. These studies indicated that *the peptide group has a rigid, planar structure, which, Pauling pointed out, is a conse- quence of resonance interactions that give the peptide bond an ~40% double-bond character.* 

This explanation is supported by the observations that a peptide's C¬N bond is 0.13 Å shorter than its N¬C<sub>a</sub> sin- gle bond and that its C<sup>•</sup> O bond is 0.02 Å longer than that of aldehydes and ketones. The peptide bond's resonance energy has its maximum value, ~85 kJ · mol<sup>-1</sup>, when the peptide group is planar because its g-bonding overlap is maximized in this conformation. This overlap, and thus the resonance energy, falls to zero as the peptide bond is twisted to 90° out of planarity, thereby accounting for the planar peptide group's rigidity. (The positive charge on the above resonance structure should be taken as a formal charge; quantum mechanical calculations indicate that the peptide N atom, in fact, has a partial negative charge aris- ing from the polarization of the C¬N bond.). *Peptide groups, with few exceptions, assume the trans conformation: that in which successive C<sub>a</sub> atoms are on op- posite sides of the peptide bond joining them. This is partly a result of steric interference, which causes the cis conformation to be ~8 kJ ·* 

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 $mol^{-1}$  less stable than the trans conformation (this energy difference is somewhat less in peptide bonds followed by a Pro residue and, in fact,~10% of the Pro residues in proteins follow a cis peptide bond, whereas cis peptides are otherwise extremely rare).

### **Polypeptide Backbone Conformations May Be Described by Their Torsion Angles**

The above considerations are important because they indicate that the backbone of a protein is a linked sequence of rigid planar peptide groups. We can therefore specify a polypeptide's backbone conformation by the torsion angles (rotation angles or dihedral angles) about the  $C_a \neg N$  bond (\$) and the  $C_a \neg C$  bond (†) of each of its amino acid residues. These angles, \$ and †, are both defined as 180° when the polypeptide chain is in its planar, fully extended (all-trans) conformation and increase for a clockwise rotation when viewed from C<sub>a</sub>. There are several steric constraints on the torsion an- gles, \$ and *†*, of a polypeptide backbone that limit its con- formational range. The electronic structure of a single (o) bond, such as a  $C\neg C$  bond, is cylindrically symmetrical about its bond axis, so that we might expect such a bond to exhibit free rotation. If this were the case, then in ethane, for example, all torsion angles about the  $C\neg C$  bond would be equally likely. Yet certain conformations in ethane are favored due to quantum mechanical effects arising from the interactions of its molecular orbitals. The staggered conformation (torsion angle =  $180^{\circ}$ ) is ethane's most stable arrangement, whereas the eclipsed conformation (torsion angle =  $0^{\circ}$ ) is least stable. The energy difference between the staggered and eclipsed con- formations in ethane is ~12 kJ  $\cdot$  mol<sup>-1</sup>, a quantity that rep- resents an energy barrier to free rotation about the  $C\neg C$  single bond. Substituents other than hydrogen exhibit greater steric interference; that is, they increase the size of this energy barrier due to their greater bulk. Indeed, with large substituents, some conformations may be sterically forbidden.

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### **Helical Structures**

Helices are the most striking elements of protein  $2^{\circ}$  struc- ture. If a polypeptide chain is twisted by the same amount about each of its C<sub>a</sub> atoms, it assumes a helical conforma- tion. As an alternative to specifying its \$ and † angles, a helix may be characterized by the number, *n*, of peptide units per helical turn and by its **pitch**, *p*, the distance the helix rises along its axis per turn. Several examples of he- lices are diagrammed in the Fig. Note that a helix has chi rality; that is, it may be either right handed or left handed (a right-handed helix turns in the direction that the fingers of a right hand curl when its thumb points along the helix axis in the direction that the helix rises). In proteins, more- over, *n* need not be an integer and, in fact, rarely is.

### The Helix

Only one helical polypeptide conformation has simulta- neously allowed conformation angles and a favorable hydrogen bonding pattern: the a **helix**, a partic- ularly rigid arrangement of the polypeptide chain. Its dis- covery through model building, by Pauling in 1951, ranks as one of the landmarks of structural biochemistry. For a polypeptide made from L-a-amino acid residues, the a helix is right handed with torsion angles  $\$ = -57^{\circ}$  and  $\dagger = -47^{\circ}$ , n = 3.6 residues per turn, and a pitch of 5.4 Å. (An a helix of p-a-amino acid residues is the mir- ror image of that made from L-amino acid residues: It is left handed with conformation angles  $\$ = +57^\circ$ ,  $\dagger = +47^\circ$ , and n = -3.6but with the same value of p.) Figure 8-11 indicates that the hydrogen bonds of an a helix are arranged such that the peptide N¬H bond of the nth residue points along the helix toward the peptide C<sup>-c</sup>O group of the (n-4)th residue. This result in a strong hydrogen bond that has the nearly optimum N P O distance of 2.8 Å. In addition, the core of the a helix is tightly packed; that is, its atoms are in van der Waals con- tact across the helix, thereby maximizing their association energies (Section 8-4A). The R groups, whose positions, as we saw, are not fully dealt with by the Ramachandran diagram, all project backward and outward from the helix so as to avoid steric interference with the polypeptide backbone and with each other. Such an arrangement can also be seen in the

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fig. Indeed, a ma- jor reason why the left-handed a helix has never been ob- served (its helical parameters are but mildly forbidden; is that its side chains contact its polypeptide backbone too closely. Note, however, that 1 to 2% of the individual non-Gly residues in proteins assume this con- formation. The a helix is a common secondary structural element of both fibrous and globular proteins. In globular proteins, a helices have an average span of ~12 residues, which cor- responds to over three helical turns and a length of 18 Å. However, a helices with as many as 53 residues have been found.

### **Beta Structures**

In 1951, the year that they proposed the a helix, Pauling and Corey also postulated the existence of a different polypeptide sec- ondary structure, the b pleated sheet. As with the a helix, the b pleated sheet's conformation has repeating \$ and † angles that fall in the allowed region of the Ramachandran diagram and utilizes the full hydrogen bonding capacity of the polypeptide backbone. In *b* pleated sheets, however, hydrogen bonding occurs between neighboring polypeptide chains rather than within one as in a helices. Pleated sheets come in two varieties: The antiparallel b pleated sheet, in which neighboring hydrogen bonded polypeptide chains run in opposite directions. The parallel b pleated sheet, in which the hydrogen bonded chains extend in the same direction. The conformations in which these b structures are opti- mally hydrogen bonded vary somewhat from that of a fully extended polypeptide ( $\$ = \dagger = \pm 180^\circ$ ), as indicated in the fig. They therefore have a rippled or pleated edge-on appearance, which accounts for the appellation "pleated sheet." In this conformation, successive side chains of a polypeptide chain extend to opposite sides of the pleated sheet with a two-residue repeat distance of 7.0 Å. b Sheets are common structural motifs in proteins. In globular proteins, they consist of from 2 to as many as 15 polypeptide strands, the average being 6 strands, which have an aggregate width of ~25 Å. The polypeptide chains in a  $\beta$  sheet are known to be up to 15 residues long, with the average being 6 residues that have a length of ~21 Å. A 6-stranded antiparallel b sheet, for example, occurs in the jack bean protein concanavalin A. Parallel b sheets of less than five strands are rare. This

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observation suggests that parallel b sheets are less stable than antiparallel b sheets, possibly because the hydrogen bonds of parallel sheets are distorted in comparison to those of the antiparallel sheets.

### **PROTEIN STABILITY**

Incredible as it may seem, thermodynamic measurements indicate that *native* proteins are only marginally stable en- tities under physiological conditions. The free energy re- quired to denature them is ~0.4 kJ  $\cdot$  mol<sup>-1</sup> of amino acid residues, so that 100-residue proteins are typically stable by only around 40 kJ  $\cdot$  mol<sup>-1</sup>. In contrast, the energy re- quired to break a typical hydrogen bond is ~20 kJ  $\cdot$  mol<sup>-1</sup>. The various noncovalent influences to which proteins are subject—electrostatic interactions (both attractive and repulsive), hydrogen bonding (both intramolecular and to water), and hydrophobic forces—each have energetic mag- nitudes that may total thousands of kilojoules per mole over an entire protein molecule. Consequently, a protein structure arises from a delicate balance among powerful countervailing forces. In this section we discuss the nature of these forces and end by considering protein denatura- tion, that is, how these forces can be disrupted.

### **Electrostatic Forces**

Molecules are collections of electrically charged particles and hence, to a reasonable degree of approximation, their interactions are determined by the laws of classical elec- trostatics (more exact calculations require the application of quantum mechanics). The energy of association, U, of two electric charges,  $q_1$  and  $q_2$ , that are separated by the distance r is found by integrating the expression for Coulomb's law, Eq. [2.1], to determine the work necessary to separate these charges by an infinite distance: Here  $k = 9.0 \times 10^9$  J  $\cdot$  m  $\cdot$  C<sup>--2</sup> and D is the dielectric constant of the medium in which the charges are immersed (recall that D = 1 for a vacuum and, for the most part, increases with the polarity of the medium; Table 2-1). The dielectric constant of a molecule-sized region is difficult to estimate. For the interior of a protein, it is usually taken to be in

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the range 3 to 5 in analogy with the measured di- electric constants of substances that have similar polarities, such as benzene and diethyl ether.

### Hydrogen Bonding Forces

Hydrogen bonds (D¬H P A), as we discussed in Section 2-1A, are predominantly electrostatic interactions (but with ~10% covalent character) between a weakly acidic donor group  $(D\neg H)$  and an acceptor (A) that bears a lone pair of electrons. In biological systems, D and A can both be the highly electronegative N and O atoms and occasionally S atoms. In addition, a relatively acidic  $C \neg H$  group (e.g., a  $C_a \neg H$  group) can act as a weak hydrogen bond donor, and the polarizable g electron system of an aromatic ring (e.g., that of Trp) can act as a weak acceptor. Hydrogen bonds have association energies that are nor- mally in the range -12 to -40 kJ  $\cdot$  mol<sup>-1</sup> (but only around —8 to —16 kJ  $\cdot$  mol<sup>-1</sup> for C¬H P A and D¬H P g hydrogen bonds and — 2 to  $-4 \text{ kJ} \cdot \text{mol}^{-1}$  for C¬H P g hydrogen bonds), values which are between those for covalent bonds and van der Waals forces. Hydrogen bonds (H bonds) are much more directional than are van der Waals forces but less so than are covalent bonds. The D P A distance is normally in the range 2.7 to 3.1 Å, al- though since H atoms are unseen in all but the very high- est resolution macromolecular X-ray structures, a possible D¬H P A interaction (where D and A are either N or O) is assumed to be a H bond if its D P A distance is sig- nificantly less than the 3.7 Å sum of a  $D\neg H$  bond length (~1.0 Å) and the van der Waals contact distance between H and A (~2.7 Å). Keep in mind, however, that there is no rigid cutoff distance beyond which H bonds cease to exist because the energy of an H bond, which is mainly electrostatic in character, varies inversely with the distance between the negative and positive centers. H bonds tend to be linear, with the D¬H bond pointing along the acceptor's lone pair orbital hydrogen bonds, roughly perpendicular to the aromatic ring and pointing at its center with the distance from the D atom to the center of the aromatic ring normally in the range 3.2 - 3.8 Å).

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### Hydrophobic effect

The hydrophobic effect is the observed tendency of nonpolar substances to aggregate in aqueous solution and exclude water molecules. The word hydrophobic literally means "waterfearing", and it describes the segregation of water and nonpolar substances, which maximizes hydrogen bonding between molecules of water and minimizes the area of contact between water and nonpolar molecules. The hydrophobic effect is responsible for the separation of a mixture of oil and water into its two components. It is also responsible for effects related to biology, including: cell membranes and vesicles formation, protein folding, insertion of membrane proteins into the nonpolar lipid environment and protein-small molecule associations. Hence the hvdrophobic effect is essential to life.<sup>[3][4][5][6]</sup> Substances for which this effect is observed are known as hydrophobes. The hydrophilic groups prevent phase separation of the molecules by maintaining the hydrophobic groups in water through formation of strong hydrogen bonds with water molecules. The driving force for this self-assembly is the hydrophobic effect. Amphiphiles are molecules that have both hydrophobic and hydrophilic domains. Detergents are composed of amphiphiles that allow hydrophobic molecules to be solubilized in water by forming micelles and bilayers (as in soap bubbles). They are also important to cell membranes composed of amphiphilic phospholipids that prevent the internal aqueous environment of a cell from mixing with external water.

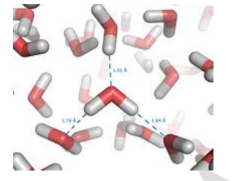
### Folding of macromolecules

In the case of protein folding, the hydrophobic effect is important to understanding the structure of proteins that have hydrophobic amino acids (such as alanine, valine, leucine, isoleucine, phenylalanine, tryptophan and methionine) clustered together within the protein. Structures of water-soluble proteins have a hydrophobic core in which side chains are buried from water, which stabilizes the folded state. Charged and polar side chains are situated on the solvent-exposed surface where they interact with surrounding water molecules. Minimizing the number of hydrophobic side chains exposed to water is the principal driving force behind the folding process, although formation of hydrogen bonds within the protein also stabilizes protein structure. The energetics of DNA tertiary structure assembly were determined to be driven by

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the hydrophobic effect, in addition to Watson-Crick base pairing, which is responsible for sequence selectivity, and stacking interactions between the aromatic bases. In biochemistry, the hydrophobic effect can be used to separate mixtures of proteins based on their hydrophobicity. Column chromatography with a hydrophobic stationary phase such as phenyl-sepharose will cause more hydrophobic proteins to travel more slowly, while less hydrophobic ones elute from the column sooner. To achieve better separation, a salt may be added (higher concentrations of salt increase the hydrophobic effect) and its concentration decreased as the separation progresses.



### Dynamic hydrogen bonds between molecules of liquid water

The origin of the hydrophobic effect is not fully understood. Some argue that the hydrophobic interaction is mostly an entropic effect originating from the disruption of highly dynamic hydrogen bonds between molecules of liquid water by the nonpolar solute. A hydrocarbon chain or a similar nonpolar region of a large molecule is incapable of forming hydrogen bonds with water. Introduction of such a non-hydrogen bonding surface into water causes disruption of the hydrogen bonding network between water molecules. The hydrogen bonds are reoriented tangentially to such surface to minimize disruption of the hydrogen bonded 3D network of water molecules, and this leads to a structured water "cage" around the nonpolar surface. The water molecules that form the "cage" (or solvation shell) have restricted mobility. In the solvation shell of small nonpolar particles, the restriction amounts to some 10%. For example, in the case of larger nonpolar molecules, the reorientational and translational motion of the water molecules in the solvation shell may be restricted by a factor of two to four; thus, at

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25 °C the reorientational correlation time of water increases from 2 to 4-8 picoseconds. Generally, this leads to significant losses in translational and rotational entropy of water molecules and makes the process unfavorable in terms of the free energy in the system. By aggregating together, nonpolar molecules reduce the surface area exposed to water and minimize their disruptive effect.

The hydrophobic effect can be quantified by measuring the partition coefficients of nonpolar molecules between water and non-polar solvents. The partition coefficients can be transformed to free energy of transfer which includes enthalpic and entropic components,  $\Delta G = \Delta H - T\Delta S$ . These components are experimentally determined by calorimetry. The hydrophobic effect was found to be entropy-driven at room temperature because of the reduced mobility of water molecules in the solvation shell of the non-polar solute; however, the enthalpic component of transfer energy was found to be favorable, meaning it strengthened water-water hydrogen bonds in the solvation shell due to the reduced mobility of water molecules. At the higher temperature, when water molecules become more mobile, this energy gain decreases along with the entropic component. The hydrophobic effect depends on the temperature, which leads to "cold denaturation" of proteins.

### Van der Waals force

In physical chemistry, the van der Waals forces, named after Dutch scientist Johannes Diderik van der Waals, are distance-dependent interactions between atoms or molecules. Unlike ionic or covalent bonds, these attractions are not a result of any chemical electronic bond, and they are comparatively weak and more susceptible to being perturbed. Van der Waals forces quickly vanish at longer distances between interacting molecules. Van der Waals forces play a fundamental role in fields as diverse as supramolecular chemistry, structural biology, polymer science, nanotechnology, surface science, and condensed matter physics. van der Waals forces also define many properties of organic compounds and molecular solids, including their solubility in polar and non-polar media.

If no other forces are present, the point at which the force becomes repulsive rather than attractive as two atoms near one another is called the van der Waals contact distance. This results

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from the electron clouds of two atoms unfavorably coming into contact. It can be shown that van der Waals forces are of the same origin as the Casimir effect, arising from quantum interactions with the zero-point field. The resulting van der Waals forces can be attractive or repulsive. It is also sometimes used loosely as a synonym for the totality of intermolecular forces. The term includes the force between permanent dipoles (Keesom force), the force between a permanent dipole and a corresponding induced dipole (Debye force), and the force between instantaneously induced dipoles (London dispersion force).

Being the weakest of the weak chemical forces, with a strength between 0.4 and 4kJ/mol they may still support an integral structural load when multitudes of such interactions are present. Such a force results from a transient shift in electron density. Specifically, as the electrons are in orbit of the protons and neutrons within an atom the electron density may tend to shift more greatly on a side. Thus, this generates a transient charge to which a nearby atom can be either attracted or repelled. When the interatomic distance of two atoms is greater than 0.6 nm the force is not strong enough to be observed. In the same vein, when the interatomic distance is below 0.4 nm the force becomes repulsive.

Intermolecular forces have four major contributions:

- 1. A repulsive component resulting from the Pauli exclusion principle that prevents the collapse of molecules.
- 2. Attractive or repulsive electrostatic interactions between permanent charges (in the case of molecular ions), dipoles (in the case of molecules without inversion center), quadrupoles (all molecules with symmetry lower than cubic), and in general between permanent multipoles. The electrostatic interaction is sometimes called the Keesom interaction or Keesom force after Willem Hendrik Keesom.
- 3. Induction (also known as polarization), which is the attractive interaction between a permanent multipole on one molecule with an induced multipole on another. This interaction is sometimes called Debye force after Peter J.W. Debye.

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4. Dispersion (usually named after Fritz London), which is the attractive interaction between any pair of molecules, including non-polar atoms, arising from the interactions of instantaneous multipoles.

Returning to nomenclature, different texts refer to different things using the term "van der Waals force". Some texts describe the van der Waals force as the totality of forces (including repulsion); others mean all the attractive forces (and then sometimes distinguish van der Waals-Keesom, van der Waals-Debye, and van der Waals-London). All intermolecular/van der Waals forces are anisotropic (except those between two noble gas atoms), which means that they depend on the relative orientation of the molecules. The induction and dispersion interactions are always attractive, irrespective of orientation, but the electrostatic interaction changes sign upon rotation of the molecules. That is, the electrostatic force can be attractive or repulsive, depending on the mutual orientation of the molecules. When molecules are in thermal motion, as they are in the gas and liquid phase, the electrostatic force is averaged out to a large extent, because the molecules thermally rotate and thus probe both repulsive and attractive parts of the electrostatic force. Sometimes this effect is expressed by the statement that "random thermal motion around room temperature can usually overcome or disrupt them" (which refers to the electrostatic component of the van der Waals force). Clearly, the thermal averaging effect is much less pronounced for the attractive induction and dispersion forces. The Lennard-Jones potential is often used as an approximate model for the isotropic part of a total (repulsion plus attraction) van der Waals force as a function of distance.

Van der Waals forces are responsible for certain cases of pressure broadening (van der Waals broadening) of spectral lines and the formation of van der Waals molecules. The Londonvan der Waals forces are related to the Casimir effect for dielectric media, the former being the microscopic description of the latter bulk property. The first detailed calculations of this were done in 1955 by E. M. Lifshitz. A more general theory of van der Waals forces has also been developed.

The main characteristics of van der Waals forces are:

• They are weaker than normal covalent and ionic bonds.

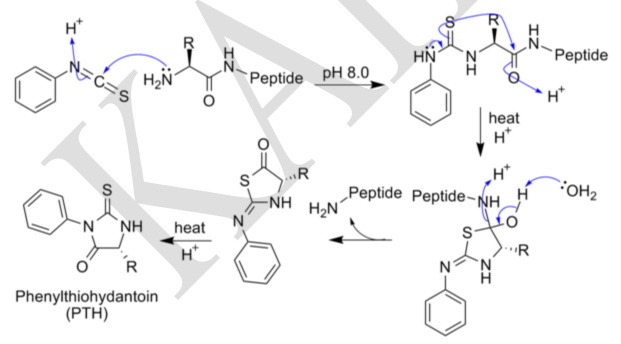
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- van der Waals forces are additive and cannot be saturated.
- They have no directional characteristic.
- They are all short-range forces and hence only interactions between the nearest particles need to be considered (instead of all the particles). Van der Waals attraction is greater if the molecules are closer.
- van der Waals forces are independent of temperature except dipole dipole interactions.

In low molecular weight alcohols, the hydrogen-bonding properties of their polar hydroxyl group dominate other weaker van der Waals interactions. In higher molecular weight alcohols, the properties of the nonpolar hydrocarbon chain(s) dominate and define the solubility.

### **Protein Sequencing**

Edman degradation, developed by Pehr Edman, is a method of sequencing amino acids in a peptide. In this method, the amino-terminal residue is labeled and cleaved from the peptide without disrupting the peptide bonds between other amino acid residues.



Phenyl isothiocyanate is reacted with an uncharged N-terminal amino group, under mildly alkaline conditions, to form a cyclical *phenylthiocarbamoyl* derivative. Then, under acidic

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conditions, this derivative of the terminal amino acid is cleaved as a thiazolinone derivative. The thiazolinone amino acid is then selectively extracted into an organic solvent and treated with acid to form the more stable phenylthiohydantoin (PTH)- amino acid derivative that can be identified by using chromatography or electrophoresis. This procedure can then be repeated again to identify the next amino acid. A major drawback to this technique is that the peptides being sequenced in this manner cannot have more than 50 to 60 residues (and in practice, under 30). The peptide length is limited due to the cyclical derivatization not always going to completion. The derivatization problem can be resolved by cleaving large peptides into smaller peptides before proceeding with the reaction. It is able to accurately sequence up to 30 amino acids with modern machines capable of over 99% efficiency per amino acid. An advantage of the Edman degradation is that it only uses 10 - 100 pico-moles of peptide for the sequencing process. The Edman degradation reaction was automated in 1967 by Edman and Beggs to speed up the process and 100 automated devices were in use worldwide by 1973.

Because the Edman degradation proceeds from the N-terminus of the protein, it will not work if the N-terminus has been chemically modified (e.g. by acetylation or formation of pyroglutamic acid). Sequencing will stop if a non- $\alpha$ -amino acid is encountered (e.g. isoaspartic acid), since the favored five-membered ring intermediate is unable to be formed. Edman degradation is generally not useful to determine the positions of disulfide bridges. It also requires peptide amounts of 1 picomole or above for discernible results.

## **Sedimentation**

Sedimentation is the tendency for particles in suspension to settle out of the fluid in which they are entrained and come to rest against a barrier. This is due to their motion through the fluid in response to the forces acting on them: these forces can be due to gravity, centrifugal acceleration, or electromagnetism. In geology, sedimentation is often used as the opposite of erosion, i.e., the terminal end of sediment transport. In that sense, it includes the termination of transport by saltation or true bedload transport. Settling is the falling of suspended particles through the liquid, whereas sedimentation is the termination of the settling process. In estuarine environments, settling can be influenced by the presence or absence of vegetation. Trees such as

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mangroves are crucial to the attenuation of waves or currents, promoting the settlement of suspended particles.

Sedimentation may pertain to objects of various sizes, ranging from large rocks in flowing water to suspensions of dust and pollen particles to cellular suspensions to solutions of single molecules such as proteins and peptides. Even small molecules supply a sufficiently strong force to produce significant sedimentation. The term is typically used in geology to describe the deposition of sediment which results in the formation of sedimentary rock, but it is also used in various chemical and environmental fields to describe the motion of often-smaller particles and molecules. This process is also used in the biotech industry to separate cells from the culture media.

### Size-exclusion chromatography

Size-exclusion chromatography (SEC), also known as molecular sieve chromatography, is a chromatographic method in which molecules in solution are separated by their size, and in some cases molecular weight. It is usually applied to large molecules or macromolecular complexes such as proteins and industrial polymers. Typically, when an aqueous solution is used to transport the sample through the column, the technique is known as gel-filtration chromatography, versus the name gel permeation chromatography, which is used when an organic solvent is used as a mobile phase. SEC is a widely used polymer characterization method because of its ability to provide good molar mass distribution (Mw) results for polymers.

The main application of gel-filtration chromatography is the fractionation of proteins and other water-soluble polymers, while gel permeation chromatography is used to analyze the molecular weight distribution of organic-soluble polymers. Either technique should not be confused with gel electrophoresis, where an electric field is used to "pull" or "push" molecules through the gel depending on their electrical charges.

The advantages of this method include good separation of large molecules from the small molecules with a minimal volume of eluate, and that various solutions can be applied without interfering with the filtration process, all while preserving the biological activity of the particles to separate. The technique is generally combined with others that further separate molecules by

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other characteristics, such as acidity, basicity, charge, and affinity for certain compounds. With size exclusion chromatography, there are short and well-defined separation times and narrow bands, which lead to good sensitivity. There is also no sample loss because solutes do not interact with the stationary phase. The other advantage to this experimental method is that in certain cases, it is feasible to determine the approximate molecular weight of a compound. The shape and size of the compound (eluent) determine how the compound interacts with the gel (stationary phase). To determine approximate molecular weight, the elution volumes of compounds with their corresponding molecular weights are obtained and then a plot of "K<sub>av</sub>" vs "log(Mw)" is made, where  $K_{av} = (V_e - V_o)/(V_t - V_o)$  and Mw is the molecular mass. This plot acts as a calibration curve, which is used to approximate the desired compound's molecular weight. The V<sub>e</sub> component represents the volume at which the intermediate molecules elute such as molecules that have partial access to the beads of the column. In addition, V<sub>t</sub> is the sum of the total volume between the beads and the volume within the beads. The  $V_0$  component represents the volume at which the larger molecules elute, which elute in the beginning. Disadvantages are, for example, that only a limited number of bands can be accommodated because the time scale of the chromatogram is short, and, in general, there must be a 10% difference in molecular mass to have a good resolution.

The technique was invented by Grant Henry Lathe and Colin R Ruthven, working at Queen Charlotte's Hospital, London. They later received the John Scott Award for this invention. While Lathe and Ruthven used starch gels as the matrix, Jerker Porath and Per Flodin later introduced dextran gels; other gels with size fractionation properties include agarose and polyacrylamide. A short review of these developments has appeared. There were also attempts to fractionate synthetic high polymers; however, it was not until 1964, when J. C. Moore of the Dow Chemical Company published his work on the preparation of gel permeation chromatography (GPC) columns based on cross-linked polystyrene with controlled pore size, that a rapid increase of research activity in this field began. It was recognized almost immediately that with proper calibration, GPC was capable to provide molar mass and molar

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mass distribution information for synthetic polymers. Because the latter information was difficult to obtain by other methods, GPC came rapidly into extensive use.

SEC is used primarily for the analysis of large molecules such as proteins or polymers. SEC works by trapping smaller molecules in the pores of the adsorbent materials adsorption ("stationary phases"). This process is usually performed with a column, which consists of a hollow tube tightly packed with extremely small porous polymer beads designed to have pores of different sizes. These pores may be depressions on the surface or channels through the bead. As the solution travels down the column some particles enter into the pores. Larger particles cannot enter into as many pores. The larger the particles, the faster the elution. The larger molecules simply pass by the pores because those molecules are too large to enter the pores. Larger molecules therefore flow through the column more quickly than smaller molecules, that is, the smaller the molecule, the longer the retention time.

One requirement for SEC is that the analyte does not interact with the surface of the stationary phases, with differences in elution time between analytes ideally being based solely on the solute volume the analytes can enter, rather than chemical or electrostatic interactions with the stationary phases. Thus, a small molecule that can penetrate every region of the stationary phase pore system can enter a total volume equal to the sum of the entire pore volume and the interparticle volume. This small molecule elutes late (after the molecule has penetrated all of the pore- and interparticle volume—approximately 80% of the column volume). At the other extreme, a very large molecule that cannot penetrate any the smaller pores can enter only the interparticle volume (~35% of the column volume) and elutes earlier when this volume of mobile phase has passed through the column. The underlying principle of SEC is that particles of different sizes elute (filter) through a stationary phase at different rates. This results in the separation of a solution of particles based on size. Provided that all the particles are loaded simultaneously or near-simultaneously, particles of the same size should elute together.

However, as there are various measures of the size of a macromolecule (for instance, the radius of gyration and the hydrodynamic radius), a fundamental problem in the theory of SEC has been the choice of a proper molecular size parameter by which molecules of different kinds

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are separated. Experimentally, Benoit and co-workers found an excellent correlation between elution volume and a dynamically based molecular size, the hydrodynamic volume, for several different chain architecture and chemical compositions. The observed correlation based on the hydrodynamic volume became accepted as the basis of universal SEC calibration. Still, the use of the hydrodynamic volume, a size based on dynamical properties, in the interpretation of SEC data is not fully understood. This is because SEC is typically run under low flow rate conditions where hydrodynamic factor should have little effect on the separation. In fact, both theory and computer simulations assume a thermodynamic separation principle: the separation process is determined by the equilibrium distribution (partitioning) of solute macromolecules between two phases --- a dilute bulk solution phase located at the interstitial space and confined solution phases within the pores of column packing material. Based on this theory, it has been shown that the relevant size parameter to the partitioning of polymers in pores is the mean span dimension (mean maximal projection onto a line). Although this issue has not been fully resolved, it is likely that the mean span dimension and the hydrodynamic volume are strongly correlated.

Each size exclusion column has a range of molecular weights that can be separated. The exclusion limit defines the molecular weight at the upper end of the column 'working' range and is where molecules are too large to get trapped in the stationary phase. The lower end of the range is defined by the permeation limit, which defines the molecular weight of a molecule that is small enough to penetrate all pores of the stationary phase. All molecules below this molecular mass are so small that they elute as a single band. The filtered solution that is collected at the end is known as the eluate. The void volume includes any particles too large to enter the medium, and the solvent volume is known as the column volume.

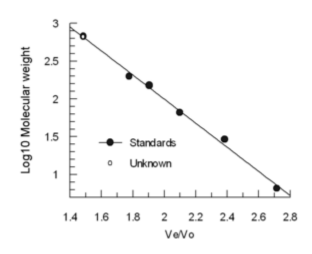
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### A cartoon illustrating the theory behind size exclusion chromatography

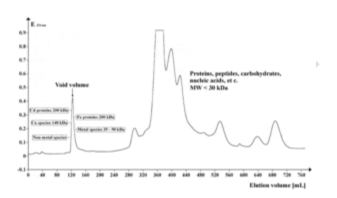
In real-life situations, particles in solution do not have a fixed size, resulting in the probability that a particle that would otherwise be hampered by a pore passing right by it. Also, the stationary-phase particles are not ideally defined; both particles and pores may vary in size. Elution curves, therefore, resemble Gaussian distributions. The stationary phase may also interact in undesirable ways with a particle and influence retention times, though great care is taken by column manufacturers to use stationary phases that are inert and minimize this issue. Like other forms of chromatography, increasing the column length enhances resolution, and increasing the column diameter increases column capacity. Proper column packing is important for maximum resolution: An over-packed column can collapse the pores in the beads, resulting in a loss of resolution. An under-packed column can reduce the relative surface area of the stationary phase accessible to smaller species, resulting in those species spending less time trapped in pores. Unlike affinity chromatography techniques, a solvent head at the top of the column can drastically diminish resolution as the sample diffuses prior to loading, broadening the downstream elution. In simple manual columns, the eluent is collected in constant volumes, known as fractions. The more similar the particles are in size the more likely they are in the same fraction and not detected separately. More advanced columns overcome this problem by constantly monitoring the eluent.

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#### Standardization of a size exclusion column.

The collected fractions are often examined by spectroscopic techniques to determine the concentration of the particles eluted. Common spectroscopy detection techniques are refractive index (RI) and ultraviolet (UV). When eluting spectroscopically similar species (such as during biological purification), other techniques may be necessary to identify the contents of each fraction. It is also possible to analyse the eluent flow continuously with RI, LALLS, Multi-Angle Laser Light Scattering MALS, UV, and/or viscosity measurements.



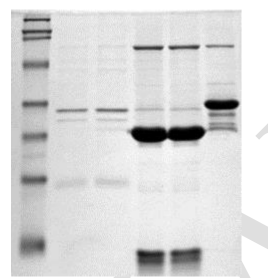
The elution volume (Ve) decreases roughly linear with the logarithm of the molecular hydrodynamic volume. Columns are often calibrated using 4-5 standard samples (e.g., folded proteins of known molecular weight), and a sample containing a very large molecule such as

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thyroglobulin to determine the void volume. (Blue dextran is not recommended for Vo determination because it is heterogeneous and may give variable results) The elution volumes of the standards are divided by the elution volume of the thyroglobulin (Ve/Vo) and plotted against the log of the standards.

# Polyacrylamide gel electrophoresis



### Picture of an SDS-PAGE. The molecular markers (ladder) are in the left lane

Polyacrylamide gel electrophoresis (PAGE), describes a technique widely used in biochemistry, forensics, genetics, molecular biology and biotechnology to separate biological macromolecules, usually proteins or nucleic acids, according to their electrophoretic mobility. Mobility is a function of the length, conformation and charge of the molecule. As with all forms of gel electrophoresis, molecules may be run in their native state, preserving the molecules' higher-order structure. This method is called native-PAGE. Alternatively, a chemical denaturant may be added to remove this structure and turn the molecule into an unstructured molecule whose mobility depends only on its length and mass-to-charge ratio. This procedure is called SDS-PAGE. Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) is a method of separating molecules based on the difference of their molecular weight. At the pH at

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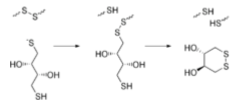
which gel electrophoresis is carried out the SDS molecules are negatively charged and bind to proteins in a set ratio, approximately one molecule of SDS for every 2 amino acids. In this way, the detergent provides all proteins with a uniform charge-to-mass ratio, independently of their original charge. By binding to the proteins the detergent destroys their secondary, tertiary and/or quaternary structure denaturing them and turning them into negatively charged linear poly peptide chains. When subjected to an electric field in PAGE, the negatively charged poly peptide chains travel toward the anode with different mobility. Their mobility, or the distance traveled by molecules, is inversely proportional to the logarithm of their molecular weight. By comparing the relative ratio of the distance traveled by each protein to the length of the gel (Rf) one can make conclusions about the relative molecular weight of the proteins, where the length of the gel is determined by the distance traveled by a small molecule like a tracking dye.

For nucleic acids, urea is the most commonly used denaturant. For proteins, sodium dodecyl sulfate (SDS) is an anionic detergent applied to protein samples to coat proteins in order to impart two negative charges (from every SDS molecule) to every two amino acids of the denatured protein. 2-Mercaptoethanol may also be used to disrupt the disulfide bonds found between the protein complexes, which helps further denature the protein. In most proteins, the binding of SDS to the polypeptide chain imparts an even distribution of charge per unit mass, thereby resulting in a fractionation by approximate size during electrophoresis. Proteins that have a greater hydrophobic content – for instance, many membrane proteins, and those that interact with surfactants in their native environment – are intrinsically harder to treat accurately using this method, due to the greater variability in the ratio of bound SDS. Procedurally, using both Native and SDS-PAGE together can be used to purify and to separate the various subunits of the protein. Native-PAGE keeps the oligomeric form intact and will show a band on the gel that is representative of the level of activity. SDS-PAGE will denature and separate the oligomeric form into its monomers, showing bands that are representative of their molecular weights. These bands can be used to assess the purity of and identify the protein.

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#### Sample preparation

Samples may be any material containing proteins or nucleic acids. These may be biologically derived, for example from prokaryotic or eukaryotic cells, tissues, viruses, environmental samples, or purified proteins. In the case of solid tissues or cells, these are often first broken down mechanically using a blender (for larger sample volumes), using a homogenizer (smaller volumes), by sonicator or by using cycling of high pressure, and a combination of biochemical and mechanical techniques – including various types of filtration and centrifugation – may be used to separate different cell compartments and organelles prior to electrophoresis. Synthetic biomolecules such as oligonucleotides may also be used as analytes.



Reduction of a typical disulfide bond by DTT via two sequential thiol-disulfide exchange reactions. The sample to analyze is optionally mixed with a chemical denaturant if so desired, usually SDS for proteins or urea for nucleic acids. SDS is an anionic detergent that denatures secondary and non-disulfide-linked tertiary structures, and additionally applies a negative charge to each protein in proportion to its mass. Urea breaks the hydrogen bonds between the base pairs of the nucleic acid, causing the constituent strands to anneal. Heating the samples to at least 60 °C further promotes denaturation.

In addition to SDS, proteins may optionally be briefly heated to near boiling in the presence of a reducing agent, such as dithiothreitol (DTT) or 2-mercaptoethanol (betamercaptoethanol/BME), which further denatures the proteins by reducing disulfide linkages, thus overcoming some forms of tertiary protein folding, and breaking up quaternary protein structure (oligometric subunits). This is known as reducing SDS-PAGE.

A tracking dye may be added to the solution. This typically has a higher electrophoretic mobility than the analytes to allow the experimenter to track the progress of the solution through the gel during the electrophoretic run.

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#### Preparing acrylamide gels

The gels typically consist of acrylamide, bisacrylamide, the optional denaturant (SDS or urea), and a buffer with an adjusted pH. The solution may be degassed under a vacuum to prevent the formation of air bubbles during polymerization. Alternatively, butanol may be added to the resolving gel (for proteins) after it is poured, as butanol removes bubbles and makes the surface smooth. A source of free radicals and a stabilizer, such as ammonium persulfate and TEMED are added to initiate polymerization. The polymerization reaction creates a gel because of the added bisacrylamide, which can form cross-links between two acrylamide molecules. The ratio of bisacrylamide to acrylamide can be varied for special purposes, but is generally about 1 part in 35. The acrylamide concentration of the gel can also be varied, generally in the range from 5% to 25%. Lower percentage gels are better for resolving very high molecular weight molecules, while much higher percentages of acrylamide are needed to resolve smaller proteins. Gels are usually polymerized between two glass plates in a gel caster, with a comb inserted at the top to create the sample wells. After the gel is polymerized the comb can be removed and the gel is ready for electrophoresis.

Various buffer systems are used in PAGE depending on the nature of the sample and the experimental objective. The buffers used at the anode and cathode may be the same or different. An electric field is applied across the gel, causing the negatively charged proteins or nucleic acids to migrate across the gel away from the negative electrode (which is the cathode being that this is an electrolytic rather than galvanic cell) and towards the positive electrode (the anode). Depending on their size, each biomolecule moves differently through the gel matrix: small molecules more easily fit through the pores in the gel, while larger ones have more difficulty. The gel is run usually for a few hours, though this depends on the voltage applied across the gel; migration occurs more quickly at higher voltages, but these results are typically less accurate than at those at lower voltages. After the set amount of time, the biomolecules have migrated different distances based on their size. Smaller biomolecules travel farther down the gel, while larger ones remain closer to the point of origin. Biomolecules may therefore be separated roughly according to size, which depends mainly on molecular weight under denaturing

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conditions, but also depends on higher-order conformation under native conditions. The gel mobility is defined as the rate of migration traveled with a voltage gradient of 1V/cm and has units of cm<sup>2</sup>/sec/V. For analytical purposes, the relative mobility of biomolecules,  $R_f$ , the ratio of the distance the molecule traveled on the gel to the total travel distance of a tracking dye is plotted versus the molecular weight of the molecule (or sometimes the log of MW, or rather the M<sub>r</sub>, molecular radius). Such typically linear plots represent the standard markers or calibration curves that are widely used for the quantitative estimation of a variety of biomolecular sizes. Certain glycoproteins, however, behave anomalously on SDS gels. Additionally, the analysis of

Certain glycoproteins, however, behave anomalously on SDS gels. Additionally, the analysis of larger proteins ranging from 250,000 to 600,000 Da is also reported to be problematic due to the fact that such polypeptides move improperly in the normally used gel systems.

**Polyacrylamide gel (PAG)** had been known as a potential embedding medium for sectioning tissues as early as 1964, and two independent groups employed PAG in electrophoresis in 1959. It possesses several electrophoretically desirable features that make it a versatile medium. It is a synthetic, thermo-stable, transparent, strong, chemically relatively inert gel, and can be prepared with a wide range of average pore sizes. The pore size of a gel and the reproducibility in gel pore size are determined by three factors, the total amount of acrylamide present (%T) (T = Total concentration of acrylamide and bisacrylamide monomer), the amount of cross-linker (%C) (C = bisacrylamide concentration), and the time of polymerization of acrylamide (cf. QPNC-PAGE). Pore size decreases with increasing %T; with cross-linking, 5%C gives the smallest pore size. Any increase or decrease in %C from 5% increases the pore size, as pore size with respect to %C is a parabolic function with vertex as 5%C. This appears to be because of non-homogeneous bundling of polymer strands within the gel. This gel material can also withstand high voltage gradients, is amenable to various staining and destaining procedures, and can be digested to extract separated fractions or dried for autoradiography and permanent recording.

#### Components

Polyacrylamide gels are composed of a stacking gel and separating gel. Stacking gels have a higher porosity relative to the separating gel, and allow for proteins to migrate in a

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concentrated area. Additionally, stacking gels usually have a pH of 6.8, since the neutral glycine molecules allow for faster protein mobility. Separating gels have a pH of 8.8, where the anionic glycine slows down the mobility of proteins. Separating gels allow for the separation of proteins and have a relatively lower porosity. Here, the proteins are separated based on size (in SDS-PAGE) and size/ charge (Native PAGE).

Chemical buffer stabilizes the pH value to the desired value within the gel itself and in the electrophoresis buffer. The choice of buffer also affects the electrophoretic mobility of the buffer counterions and thereby the resolution of the gel. The buffer should also be unreactive and not modify or react with most proteins. Different buffers may be used as cathode and anode buffers, respectively, depending on the application. Multiple pH values may be used within a single gel, for example in DISC electrophoresis. Common buffers in PAGE include Tris, Bis-Tris, or imidazole.

Counterion balance the intrinsic charge of the buffer ion and also affect the electric field strength during electrophoresis. Highly charged and mobile ions are often avoided in SDS-PAGE cathode buffers, but may be included in the gel itself, where it migrates ahead of the protein. In applications such as DISC SDS-PAGE the pH values within the gel may vary to change the average charge of the counterions during the run to improve resolution. Popular counterions are glycine and tricine. Glycine has been used as the source of trailing ion or slow ion because its pKa is 9.69 and mobility of glycinate are such that the effective mobility can be set at a value below that of the slowest known proteins of net negative charge in the pH range. The minimum pH of this range is approximately 8.0.

Acrylamide ( $C_3H_5NO$ ; mW: 71.08) when dissolved in water, slow, spontaneous autopolymerization of acrylamide takes place, joining molecules together by head on tail fashion to form long single-chain polymers. The presence of a free radical-generating system greatly accelerates polymerization. This kind of reaction is known as vinyl addition polymerisation. A solution of these polymer chains becomes viscous but does not form a gel, because the chains simply slide over one another. Gel formation requires linking various chains together.

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Acrylamide is carcinogenic, a neurotoxin, and a reproductive toxin. It is also essential to store acrylamide in a cool dark and dry place to reduce autopolymerisation and hydrolysis.

Bisacrylamide (N,N'-Methylenebisacrylamide) ( $C_7H_{10}N_2O_2$ ; mW: 154.17) is the most frequently used cross linking agent for polyacrylamide gels. Chemically it can be thought of as two acrylamide molecules coupled head to head at their non-reactive ends. Bisacrylamide can crosslink two polyacrylamide chains to one another, thereby resulting in a gel.

Sodium dodecyl sulfate (SDS) (C<sub>12</sub>H<sub>25</sub>NaO<sub>4</sub>S; mW: 288.38) (only used in denaturing protein gels) is a strong detergent agent used to denature native proteins to individual polypeptides. This denaturation, which is referred to as reconstructive denaturation, is not accomplished by the total linearization of the protein, but instead, through a conformational change to a combination of random coil and  $\alpha$  helix secondary structures. When a protein mixture is heated to 100 °C in presence of SDS, the detergent wraps around the polypeptide backbone. It binds to polypeptides in a constant weight ratio of 1.4 g SDS/g of polypeptide. In this process, the intrinsic charges of polypeptides become negligible when compared to the negative charges contributed by SDS. Thus polypeptides after treatment become rod-like structures possessing a uniform charge density, that is same net negative charge per unit weight. The electrophoretic mobilities of these proteins is a linear function of the logarithms of their molecular weights. Without SDS, different proteins with similar molecular weights would migrate differently due to differences in mass-charge ratio, as each protein has an isoelectric point and molecular weight particular to its primary structure. This is known as native PAGE. Adding SDS solves this problem, as it binds to and unfolds the protein, giving a near uniform negative charge along the length of the polypeptide.

Urea  $(CO(NH_2)_2; mW: 60.06)$  is a chaotropic agent that increases the entropy of the system by interfering with intramolecular interactions mediated by non-covalent forces such as hydrogen bonds and van der Waals forces. Macromolecular structure is dependent on the net effect of these forces, therefore it follows that an increase in chaotropic solutes denatures macromolecules,

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Ammonium persulfate (APS) ( $N_2H_8S_2O_8$ ; mW: 228.2) is a source of free radicals and is often used as an initiator for gel formation. An alternative source of free radicals is riboflavin, which generated free radicals in a photochemical reaction.

TEMED (*N*, *N*, *N'*, *N'*-tetramethylethylenediamine) ( $C_6H_{16}N_2$ ; mW: 116.21) stabilizes free radicals and improves polymerization. The rate of polymerisation and the properties of the resulting gel depend on the concentrations of free radicals. Increasing the amount of free radicals results in a decrease in the average polymer chain length, an increase in gel turbidity and a decrease in gel elasticity. Decreasing the amount shows the reverse effect. The lowest catalytic concentrations that allow polymerisation in a reasonable period of time should be used. APS and TEMED are typically used at approximately equimolar concentrations in the range of 1 to 10 mM.

The following chemicals and procedures are used for processing of the gel and the protein samples visualized in it. Tracking dye; as proteins and nucleic acids are mostly colorless, their progress through the gel during electrophoresis cannot be easily followed. Anionic dyes of a known electrophoretic mobility are therefore usually included in the PAGE sample buffer. A (BPB. 3'.3".5'.5" common .... tracking dve is Bromophenol blue very tetrabromophenolsulfonphthalein). This dye is coloured at alkali and neutral pH and is a small negatively charged molecule that moves towards the anode. Being a highly mobile molecule it moves ahead of most proteins. As it reaches the anodic end of the electrophoresis medium electrophoresis is stopped. It can weakly bind to some proteins and impart a blue colour. Other common tracking dyes are xylene cyanol, which has lower mobility, and Orange G, which has a higher mobility.

Loading aids; most PAGE systems are loaded from the top into wells within the gel. To ensure that the sample sinks to the bottom of the gel, sample buffer is supplemented with additives that increase the density of the sample. These additives should be non-ionic and nonreactive towards proteins to avoid interfering with electrophoresis. Common additives are glycerol and sucrose.

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Coomassie Brilliant Blue R-250 (CBB)( $C_{45}H_{44}N_3NaO_7S_2$ ; mW: 825.97) is the most popular protein stain. It is an anionic dye, which non-specifically binds to proteins. The structure of CBB is predominantly non-polar, and it is usually used in methanolic solution acidified with acetic acid. Proteins in the gel are fixed by acetic acid and simultaneously stained. The excess dye incorporated into the gel can be removed by destaining with the same solution without the dye. The proteins are detected as blue bands on a clear background. As SDS is also anionic, it may interfere with staining process. Therefore, large volume of staining solution is recommended, at least ten times the volume of the gel.

Ethidium bromide (EtBr) is the traditionally most popular nucleic acid stain. Silver staining is used when more sensitive method for detection is needed, as classical Coomassie Brilliant Blue staining can usually detect a 50 ng protein band, Silver staining increases the sensitivity typically 10-100 fold more. This is based on the chemistry of photographic development. The proteins are fixed to the gel with a dilute methanol solution, then incubated with an acidic silver nitrate solution. Silver ions are reduced to their metallic form by formaldehyde at alkaline pH. An acidic solution, such as acetic acid stops development.<sup>[21]</sup> Silver staining was introduced by Kerenyi and Gallyas as a sensitive procedure to detect trace amounts of proteins in gels.<sup>[22]</sup> The technique has been extended to the study of other biological macromolecules that have been separated in a variety of supports.<sup>[23]</sup> Many variables can influence the colour intensity and every protein has its own staining characteristics; clean glassware, pure reagents and water of highest purity are the key points to successful staining.<sup>[24]</sup> Silver staining was developed in the 14th century for colouring the surface of glass. It has been used extensively for this purpose since the 16th century. The colour produced by the early silver stains ranged between light yellow and an orange-red. Camillo Golgi perfected the silver staining for the study of the nervous system. Golgi's method stains a limited number of cells at random in their entirety.

Autoradiography, also used for protein band detection post gel electrophoresis, uses radioactive isotopes to label proteins, which are then detected by using X-ray film. Western blotting is a process by which proteins separated in the acrylamide gel are electrophoretically

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transferred to	a stable, manipulable membrane such as a nitrocellulose, nylon, or PVDF
membrane. It	is then possible to apply immunochemical techniques to visualise the transferred
proteins, as w	ell as accurately identify relative increases or decreases of the protein of interest.
<u>Possible qu</u>	<u>lestions</u>
<u>1 Mark ques</u>	<u>tions</u>
1. Which of the a) cysteine	ne following amino acids is an alpha helix terminator? b) alanine c) proline d) glycine
2. The second a) Vander wa	ary structure is primarily maintained by als force b) Hydrogen bond c) Ionic bond d) covalent bond
3. Which of the a) Alpha helix	<ul><li>b) Beta pleated sheets</li><li>c) Beta bends</li><li>d) loops</li></ul>
4. The most c a) cystein	ommon amino acid in beta bend is b) glycine c) serine d) Aspartic acid
5. Alpha-heliz a) sheet	b) coiled spring c) linear chain d) random coil
6. The alpha l a) 0.54nm	helix rises per turn a distance of b) 1.5nm c) 3nm d) 6nm
7. Pulses cont a) Lysine	ain incomplete proteins since they lack b) Tryptophan c) Phenyl alanine d) Methinine
8. The fact th a) the hydrop forces	at the core of most globular proteins is tightly packed is due to nobic effect b) hydrogen bonding. c) electrostatic effects d)van der Waals
9. Which of the a) Phenylisothe Phenylthiocan	
a) Cleaving, s	the following is the correct order of sequencing?equencing and orderingb) Sequencing, ordering and cleavingeleaving and sequencingd) Ordering, sequencing and cleaving

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a) Phenylisot	f the following is Edman reagent?thiocyanateb) CF3 COOHc) FDNBd) Phenylthiocarbonyl	
<u>2 Marks que</u>	estions	
1. Draw the	structures of Cystine and Cysteine.	
	structures of L- $\alpha$ -Alanine and D- $\alpha$ -Alanine.	
	you differentiate peptides from proteins?	
	omomultimeric protein? Give an example.	
	eteromultimeric protein? Give an example.	
6. Define the	e quaternary structure of proteins. Give an example.	
7. What is P	rotomer? Give an example.	
8. Define the	e quaternary structure of proteins. Give an example.	
9. What is P	rotomer? Give an example.	
10. Why is electrophoresis done in solution having low salt concentration?		
11. Write a short note on 'Isotachophoresis'.		
12. Briefly write about the functions of SDS in SDS-PAGE.		
<u>6/8 Marks q</u>	uestions	
	proteins classified based on their compositions, shapes, functions, nutritional factors, ructures and folds?	
	n detail the sequencing of amino-acids by Edman method and its merits over Sanger rotein sequencing.	

3. Outline various strategies for determining primary structure and disulfide pattern(s) of a polypeptide chain.

- 4. Discuss on the following topics with suitable examples.
- a) Structural architectures of proteins.
- b) Protein classifications.

5. Write an essay on various types of regular and irregular secondary structures of proteins with suitable examples.

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6. What are essential amino acids? Draw the structures of any two essential amino acids?

7. How are amino acids classified based on their metabolic roles?

8. How will you determine the number of polypeptide chains present in a protein?

9. Discuss in detail the principles, experimental conditions and applications of agarose gel electrophoresis/SDS-PAGE/IEF/PAGE.

10. A protein analyzed by SDS-PAGE and Gel filtration techniques shows molecular weight of 100 kD and 200 kD, respectively. How will you interpret the data?

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#### **SYLLABUS**

**Proteomics:** Introduction to Proteomics, Analysis of proteomes. 2D-PAGE. Sample preparation, solubilization, reduction, resolution. Reproducibility of 2D-PAGE. Mass spectrometry based methods for protein identification. *De novo* sequencing using mass spectrometric data.

#### **Two-dimensional gel electrophoresis**

Two-dimensional gel electrophoresis, abbreviated as 2-DE or 2-D electrophoresis, is a form of gel electrophoresis commonly used to analyze proteins. Mixtures of proteins are separated by two properties in two dimensions on 2D gels. 2-DE was first independently introduced by O'Farrell and Klose in 1975.

#### **Basis for separation**

2-D <u>electrophoresis</u> begins with electrophoresis in the first dimension and then separates the molecules perpendicularly from the first to create an electropherogram in the second dimension. In electrophoresis in the first dimension, molecules are separated linearly according to their isoelectric point. In the second dimension, the molecules are then separated at 90 degrees from the first electropherogram according to molecular mass. Since it is unlikely that two molecules will be similar in two distinct properties, molecules are more effectively separated in 2-D electrophoresis than in 1-D electrophoresis.

The two dimensions that proteins are separated into using this technique can be isoelectric point, protein complex mass in the native state, and protein mass. Separation of the proteins by isoelectric point is called isoelectric focusing (IEF). Thereby, a gradient of pH is applied to a gel and an electric potential is applied across the gel, making one end more positive than the other. At all pH values other than their isoelectric point, proteins will be charged. If they are positively charged, they will be pulled towards the more negative end of the gel and if they are negatively charged they will be pulled to the more positive end of the gel. The proteins applied in the first dimension will move along the gel and will accumulate at their isoelectric point; that is, the point at which the overall charge on the protein is 0 (a neutral charge).

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For the analysis of the functioning of proteins in a cell, the knowledge of their cooperation is essential. Most often proteins act together in complexes to be fully functional. The analysis of this sub organelle organisation of the cell requires techniques conserving the native state of the protein complexes. In native polyacrylamide gel electrophoresis (native PAGE), proteins remain in their native state and are separated in the electric field following their mass and the mass of their complexes respectively. To obtain a separation by size and not by net charge, as in IEF, an additional charge is transferred to the proteins by the use of Coomassie Brilliant Blue or lithium dodecyl sulfate. After completion of the first dimension the complexes are destroyed by applying the denaturing SDS-PAGE in the second dimension, where the proteins of which the complexes are composed of are separated by their mass.

Before separating the proteins by mass, they are treated with sodium dodecyl sulfate (SDS) along with other reagents (SDS-PAGE in 1-D). This denatures the proteins (that is, it unfolds them into long, straight molecules) and binds a number of SDS molecules roughly proportional to the protein's length. Because a protein's length (when unfolded) is roughly proportional to its mass, this is equivalent to saying that it attaches a number of SDS molecules roughly proportional to the protein's mass. Since the SDS molecules are negatively charged, the result of this is that all of the proteins will have approximately the same mass-to-charge ratio as each other. In addition, proteins will not migrate when they have no charge (a result of the isoelectric focusing step) therefore the coating of the protein in SDS (negatively charged) allows migration of the proteins in the second dimension (SDS-PAGE, it is not compatible for use in the first dimension as it is charged and a nonionic or zwitterionic detergent needs to be used). In the second dimension, an electric potential is again applied, but at a 90 degree angle from the first field. The proteins will be attracted to the more positive side of the gel (because SDS is negatively charged) proportionally to their mass-to-charge ratio. As previously explained, this ratio will be nearly the same for all proteins. The proteins' progress will be slowed by frictional forces. The gel therefore acts like a molecular sieve when the current is applied, separating the proteins on the basis of their molecular weight with larger proteins being retained higher in the

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gel and smaller proteins being able to pass through the sieve and reach lower regions of the gel.

### **Detecting proteins**

The result of this is a gel with proteins spread out on its surface. These proteins can then be detected by a variety of means, but the most commonly used stains are silver and Coomassie Brilliant Blue staining. In the former case, a silver colloid is applied to the gel. The silver binds to cysteine groups within the protein. The silver is darkened by exposure to ultra-violet light. The amount of silver can be related to the darkness, and therefore the amount of protein at a given location on the gel. This measurement can only give approximate amounts, but is adequate for most purposes. Silver staining is 100x more sensitive than Coomassie Brilliant Blue with a 40fold range of linearity.

Molecules other than proteins can be separated by 2D electrophoresis. In supercoiling assays, coiled DNA is separated in the first dimension and denatured by a DNA intercalator (such as ethidium bromide or the less carcinogenic chloroquine) in the second. This is comparable to the combination of native PAGE /SDS-PAGE in protein separation.

### **Common techniques**

### **IPG-DALT**

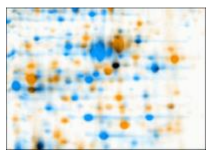
A common technique is to use an Immobilized pH gradient (IPG) in the first dimension. This technique is referred to as **IPG-DALT**. The sample is first separated onto IPG gel (which is commercially available) then the gel is cut into slices for each sample which is then equilibrated in SDS-mercaptoethanol and applied to an SDS-PAGE gel for resolution in the second dimension. Typically IPG-DALT is not used for quantification of proteins due to the loss of low molecular weight components during the transfer to the SDS-PAGE gel.

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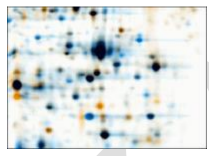
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**IEF SDS-PAGE** 

# 2D gel analysis software



Warping: Images of two 2D electrophoresis gels, overlaid with Delta2D. First image is colored in orange, second one colored in blue. Due to running differences, corresponding spots do not overlap.



Warping: Images of two 2D electrophoresis gels after warping. First image is colored in orange, second one colored in blue. Corresponding spots overlap after warping. Common spots are colored black, orange spots are only present (or much stronger) on the first image, blue spots are only present (or much stronger) on the second image.

In quantitative proteomics, these tools primarily analyze bio-markers by quantifying individual proteins, and showing the separation between one or more protein "spots" on a scanned image of a 2-DE gel. Additionally, these tools match spots between gels of similar samples to show, for example, proteomic differences between early and advanced stages of an illness. Software packages include Delta2D, ImageMaster, Melanie, PDQuest, Progenesis and REDFIN – among others.<sup>[citation needed]</sup> While this technology is widely utilized, the intelligence has not been perfected. For example, while PDQuest and Progenesis tend to agree on the

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quantification and analysis of well-defined well-separated protein spots, they deliver different results and analysis tendencies with less-defined less-separated spots. Challenges for automatic software-based analysis include incompletely separated (overlapping) spots (less-defined and/or separated), weak spots / noise (e.g., "ghost spots"), running differences between gels (e.g., protein migrates to different positions on different gels), unmatched/undetected spots, leading to missing values, mismatched spots , errors in quantification (several distinct spots may be erroneously detected as a single spot by the software and/or parts of a spot may be excluded from quantification), and differences in software algorithms and therefore analysis tendencies

#### **Protein Gel Staining Methods**

Once protein bands have been separated by electrophoresis, they can be visualized using different methods of in-gel detection, each with particular advantages and disadvantages. Over the past several decades, demand for improved sensitivity for small sample sizes and compatibility with downstream applications and detection instrumentation have driven the development of several basic staining methods. Here we discuss the general principles of protein gel staining and describe several staining methods.

#### General principles of gel staining

The first step after performing denaturing polyacrylamide gel electrophoresis (SDS-PAGE) is to disassemble the gel cassette and place the thin polyacrylamide gel in a tray filled with water or buffer. The electrophoresed proteins exist as concentrated "bands" embedded within each lane of the porous polyacrylamide gel matrix. Typically, the proteins are still bound to the anionic detergent (SDS), and the entire gel matrix is saturated in running buffer.

To make the proteins visible, a protein-specific, dye-binding or color-producing chemical reaction must be performed on the proteins within the gel. Depending on the particular chemistry of the stain, various steps are necessary to retain, or fix, the proteins in the gel matrix and to facilitate the necessary chemical reaction. All steps are done in solution, i.e., with the gel suspended in a tray filled with one liquid reagent or another.

Given the common constraints of this format, most staining methods involve some version of the same general incubation steps:

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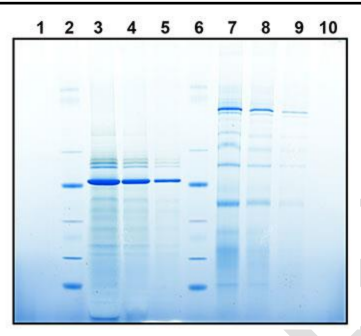
- A water wash to remove electrophoresis buffers from the gel matrix
- An acid or alcohol wash to condition or fix the gel to limit diffusion of protein bands from the matrix
- Treatment with the staining reagent to allow the dye or chemical to diffuse into the gel and bind to (or react with) the proteins
- Destaining to remove excess dye from the gel matrix background

Depending on the particular staining method, two or more of these functions can be accomplished with one step. For example, a dye reagent that is formulated in an acidic buffer can effectively fix and stain in one step. Conversely, certain functions require several steps. For example, silver staining requires both a staining reagent step and a developer step to produce the colored reaction product.

### **Coomassie dye stains**

The most common method of in-gel protein detection is staining with Coomassie dye. Several recipes for Coomassie staining reagents exist in the literature and use either the G-250 ("colloidal") or R-250 form of the dye. Colloidal Coomassie can be formulated to effectively stain proteins within 1 hour and requires only water (no methanol or acetic acid) for destaining. In acidic conditions, Coomassie dye binds to basic and hydrophobic residues of proteins, changing in color from a dull reddish-brown to intense blue (see previous images on this page). As with all staining methods, Coomassie staining detects some proteins better than others, based on the chemistry of action and differences in protein composition. Thus, Coomassie staining can detect as little as 8–10 ng per band for some proteins and 25 ng per band for most proteins.

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### Gel staining with Coomassie dye

Two-fold dilutions of protein extracts were run on an Invitrogen<sup>™</sup> NativePAGE<sup>™</sup> 3– 12% Bis-Tris Protein Gel using a Mini Gel Tank. Following electrophoresis, the gel was stained with Coomassie dye and imaged using a flatbed scanner. Lanes 1 and 10: blank; lanes 2 and 6: 5 µL Invitrogen<sup>™</sup> NativeMark<sup>™</sup> Unstained Protein Standard; lanes 3, 4 and 5: 10, 5, and 2.5 µg spinach chloroplast extract; lanes 7, 8, and 9: 10, 5, and 2.5 µg bovine mitochondrial extract. Coomassie dye staining is especially convenient because it involves a single ready-to-use reagent and does not permanently chemically modify the target proteins. An initial water wash step is necessary to remove residual SDS, which interferes with dye binding. Then the staining reagent is added, usually for about 1 hour; finally, a water or simple methanol:acetic acid destaining step is used to wash away excess unbound dye from the gel matrix. Because no chemical modification occurs, excised protein bands can be completely destained and the proteins recovered for analysis by mass spectrometry or sequencing. Coomassie staining and other traditional staining methods require several long incubation and wash steps. To expedite the

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staining process, more rapid staining protocols have been developed using powered (electrophoretic) devices such as the Thermo Scientific<sup>TM</sup> Pierce<sup>TM</sup> Power Stainer.



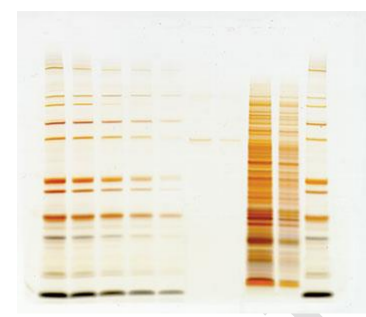
### The Thermo Scientific<sup>™</sup> Pierce<sup>™</sup> Power Stainer.

This powered device enables rapid (6–11 min) Coomassie dye staining of proteins in polyacrylamide gels, including the removal of unbound stain, in a single step. The small, easy-to-use device consists of the Pierce<sup>TM</sup> Power Station and Pierce<sup>TM</sup> Power Stain Cassette, which accommodates up to two mini gels or one midi gel at a time. The staining procedure is designed exclusively for use with Pierce<sup>TM</sup> Power Staining Kits.

### Silver stains

Silver staining is the most sensitive colorimetric method for detecting total protein. The technique involves the deposition of metallic silver onto the surface of a gel at the locations of protein bands. Silver ions (from silver nitrate in the staining reagent) interact and bind with certain protein functional groups. The strongest interactions occur with carboxylic acid groups (Asp and Glu), imidazole (His), sulfhydryls (Cys), and amines (Lys). Various sensitizer and enhancer reagents are essential for controlling the specificity and efficiency of silver ion binding to proteins and effective conversion (development) of the bound silver to metallic silver. The development process is essentially the same as for photographic film: silver ions are reduced to metallic silver, resulting in a brown-black color.

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#### Gel staining with silver stain

Samples were separated on an Invitrogen<sup>TM</sup> NuPAGE<sup>TM</sup> 4–12% Bis-Tris Protein Gel and stained with the Invitrogen<sup>TM</sup> SilverXpress<sup>TM</sup> Kit. Lanes 1–5: Invitrogen<sup>TM</sup> Mark12<sup>TM</sup> Unstained Standard (blend of 12 purified proteins), serial 2-fold dilutions ranging from 1:4 to 1:64; lane 6: 1.6 ng BSA; lane 7: 0.8 ng BSA; lane 8: *E. coli* lysate diluted 1:20; lane 9: *E. coli* lysate diluted 1:80; lane 10: replicate of lane 1. Silver staining protocols require several steps, which are affected by reagent quality as well as incubation times and thickness of the gel. An advantage of commercially available silver staining kits is that the formulations and protocols are optimized and consistently manufactured, helping to maximize consistency of results from experiment to experiment. Kits with optimized protocols are robust and easy to use, detecting less than 0.5 ng of protein in typical gels. Silver stains use either glutaraldehyde or formaldehyde as the enhancer. These reagents can cause chemical crosslinking of the proteins in the gel matrix, limiting compatibility with destaining and elution methods for analysis by mass spectrometry (MS). Therefore, optimization of sensitivity vs. protein recoverability is critical when employing silver staining as part of an MS workflow. Silver stain formulations can be made such that

Prepared by Dr. T. Sivaraman, Professor, Department of Biotechnology, KAHE

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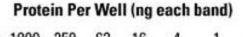
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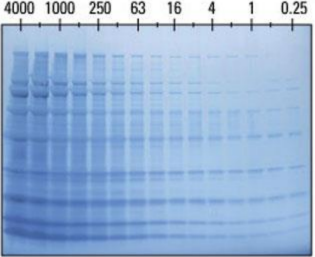
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protein bands stain black, blue-brown, red, or yellow, depending on their charge and other characteristics. This is particularly useful for differentiating overlapping spots on 2D gels.

### **Zinc stains**

Zinc staining is unlike all other staining methods. Instead of staining the proteins, this procedure stains all areas of the polyacrylamide gel in which there are no proteins. Zinc ions complex with imidazole, which precipitates in the gel matrix except where SDS-saturated proteins are located. The milky-white precipitate renders the background opaque while the protein bands remain clear. The process is short (about 15 minutes), and the gel can be photographed by viewing it over a dark background. Zinc staining is as sensitive as typical silver staining (detects less than 1 ng of protein), and no fixation steps are required. Furthermore, the stain is easily removed, making this method compatible with MS or western blotting.





# Gel staining with zinc stain

A 2-fold dilution series of a protein mixture was separated by protein gel electrophoresis using a 15-well mini gel. Subsequently the gel was stained using the Thermo Scientific<sup>™</sup> Pierce<sup>™</sup> Zinc Reversible Stain Kit, and then photographed with the gel placed over a dark blue

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background. The sensitivity on this gel is 0.25 ng, as indicated by the bands that are visible in the last lane.

### **Fluorescent dye stains**

Recent improvements in fluorescence imaging instruments and fluorescent applications have resulted in greater demand for fluorescent stains. A number of fluorescent stains for total protein have been introduced in recent years. Newer fluorescent total-protein stains provide exceptional fluorescent staining performance with fast and easy procedures. The most useful are those whose excitation and emission maxima correspond to common filter sets and laser settings of popular fluorescence imaging instruments.

### Protein gel stained with fluorescent dyes

2D gel stained with Invitrogen<sup>™</sup> SYPRO<sup>™</sup> Ruby protein gel stain and Invitrogen<sup>™</sup> Pro-Q<sup>™</sup> Emerald 300 reagent. Cohn fractions II and III from cow plasma were combined and resolved on a 2D gel. The gel was first stained with Pro-Q Emerald 300 reagent (left), followed by staining with the SYPRO Ruby stain (right). Most fluorescent stains involve simple dyebinding mechanisms rather than chemical reactions that alter protein functional groups. Therefore, most are compatible with destaining and protein recovery methods for downstream analysis by MS. Accordingly, these stains are frequently used in both 1D and 2D applications.

### **Functional group-specific stains**

Sometimes it is desirable to detect a subset of proteins rather than all of the proteins in a sample. Glycoproteins and phosphoproteins are classified as such on the basis of a particular chemical moiety (i.e., polysaccharides and phosphate groups, respectively). When a dye-binding or color-producing chemistry can be designed to detect one of these functional groups, it can be used as the basis for a specific gel stain.

### Phosphoprotein and total protein visualization in a 2D gel

Protein lysates obtained from a Jurkat T-cell lymphoma line were separated by 2D gel electrophoresis and subsequently stained with Invitrogen<sup>TM</sup> Pro-Q<sup>TM</sup> Diamond phosphoprotein

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gel stain (blue) followed by SYPRO Ruby protein gel stain (red). The gel was dried and imaged on an FLA-3000 scanner (Fuji). Shown is a digitally pseudocolored composite overlaid image.

Proteins that have been post-translationally modified by glycosylation can be detected by a procedure that involves chemical activation of the carbohydrate into a reactive group. The method works by fixing the proteins in the gel and then oxidizing the sugar residues with sodium meta-periodate. The resulting aldehyde groups can then be reacted with an amine-containing dye. In older literature, this method is known as the periodate acid–Schiff (PAS) technique. A subsequent reduction step stabilizes the dye–protein bond. Both colorimetric and fluorescent dyes have been used for this technique, and glycoprotein stain kits are available commercially.

Various protein gel staining methods, both colorimetric and fluorescent, have also been developed to detect phosphorylated proteins and His-tagged fusion proteins. For instance, certain gel stains selectively stain phosphoproteins and His-tags in acrylamide gels, without the need for blotting or phosphoprotein-specific or His-tag–specific antibodies and western blot analysis.

#### **Isoelectric focusing**

Isoelectric focusing (IEF), also known as electrofocusing, is a technique for separating different molecules by differences in their isoelectric point (pI). It is a type of zone electrophoresis, usually performed on proteins in a gel, that takes advantage of the fact that overall charge on the molecule of interest is a function of the pH of its surroundings.

IEF involves adding an ampholyte solution into immobilized pH gradient (IPG) gels. IPGs are the acrylamide gel matrix co-polymerized with the pH gradient, which result in completely stable gradients except the most alkaline (>12) pH values. The immobilized pH gradient is obtained by the continuous change in the ratio of *Immobilines*. An Immobiline is a weak acid or base defined by its pK value.

A protein that is in a pH region below its isoelectric point (pI) will be positively charged and so will migrate towards the cathode (negatively charged electrode). As it migrates through a gradient of increasing pH, however, the protein's overall charge will decrease until the protein reaches the pH region that corresponds to its pI. At this point it has no net charge and so migration ceases (as there is no electrical attraction towards either electrode). As a result, the

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proteins become focused into sharp stationary bands with each protein positioned at a point in the pH gradient corresponding to its pI. The technique is capable of extremely high resolution with proteins differing by a single charge being fractionated into separate bands.

Molecules to be focused are distributed over a medium that has a pH gradient (usually created by aliphatic ampholytes). An electric current is passed through the medium, creating a "positive" anode and "negative" cathode end. Negatively charged molecules migrate through the pH gradient in the medium toward the "positive" end while positively charged molecules move toward the "negative" end. As a particle moves towards the pole opposite of its charge it moves through the changing pH gradient until it reaches a point in which the pH of that molecules isoelectric point is reached. At this point the molecule no longer has a net electric charge (due to the protonation or deprotonation of the associated functional groups) and as such will not proceed any further within the gel. The gradient is established before adding the particles of interest by first subjecting a solution of small molecules such as polyampholytes with varying pI values to electrophoresis.

The method is applied particularly often in the study of proteins, which separate based on their relative content of acidic and basic residues, whose value is represented by the pI. Proteins are introduced into an Immobilized pH gradient gel composed of polyacrylamide, starch, or agarose where a pH gradient has been established. Gels with large pores are usually used in this process to eliminate any "sieving" effects, or artifacts in the pI caused by differing migration rates for proteins of differing sizes. Isoelectric focusing can resolve proteins that differ in pI value by as little as 0.01. Isoelectric focusing is the first step in two-dimensional gel electrophoresis, in which proteins are first separated by their pI and then further separated by molecular weight through SDS-PAGE.

#### Living cells

According to some opinions, living eukaryotic cells perform isoelectric focusing of proteins in their interior to overcome a limitation of the rate of metabolic reaction by diffusion of enzymes and their reactants, and to regulate the rate of particular biochemical processes. By concentrating the enzymes of particular metabolic pathways into distinct and small regions of its

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interior, the cell can increase the rate of particular biochemical pathways by several orders of magnitude. By modification of the isoelectric point (pI) of molecules of an enzyme by, e.g., phosphorylation or dephosphorylation, the cell can transfer molecules of the enzyme between different parts of its interior, to switch on or switch off particular biochemical processes.

### Microfluidic chip based

Microchip based electrophoresis is a promising alternative to capillary electrophoresis since it has the potential to provide rapid protein analysis, straightforward integration with other microfluidic unit operations, whole channel detection, nitrocellulose films, smaller sample sizes and lower fabrication costs.

### **Multi-junction**

The increased demand for faster and easy-to-use protein separation tools has accelerate the evolution of IEF towards in-solution separations. In this context, a multi-junction IEF system was developed to perform fast and gel-free IEF separations. The multi-junction IEF system utilizes a series of vessels with a capillary passing through each vessel. Part of the capillary in each vessel is replaced by a semipermeable membrane. The vessels contain buffer solutions with different pH values, so that a pH gradient is effectively established inside the capillary. The buffer solution in each vessel has an electrical contact with a voltage divider connected to a highvoltage power supply, which established electrical field along the capillary. When a sample (a mixture of peptides or proteins) is injected in the capillary, the presence of the electrical field and the pH gradient separates these molecules according to their isoelectric points. The multijunction IEF system has been used to separate tryptic peptide mixtures for two-dimensional proteomics and blood plasma proteins from Alzheimer's disease patients for biomarker discovery.

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An ampholyte pH 9solution is incorporated into a gel. Decreasing pH pH3A stable pH Protein After staining, gradient is solution is proteins are established in added and shown to be the gel after electric field distributed along application of is reapplied. pH gradient an electric according to field. their pI values.

### **Mass Spectrometry**

Mass spectrometry (MS) is an analytical technique that measures the mass-to-charge ratio of charged particles. It is used for determining masses of particles, for determining the elemental composition of a sample or molecule. The MS principle consists of ionizing chemical compounds to generate charged molecules or molecule fragments and measurement of their mass-to-charge ratios by using the one of a variety of techniques (e.g EI/CI/ESI/APCI/MALDI).

#### **Electron Ionization (EI) technique**

EI is the most appropriate technique for relatively small (m.w.<600) neutral organic molecules which can easily be promoted to the gas phase without decomposition, i.e. volatile. Since EI samples are thermally desorbed to the gas phase and subjected to the high energy of EI, analytes must be thermally stable. The gas phase molecules enter into the ion source where they are bombarded with free electrons emitted from a filament (Figure 1). The electrons bombard the molecules causing a hard ionization that fragments the molecule, and turn into positively charged

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particles called ions. This is important because the particles must be charged to pass through the analyser. As the ions continue from the source, they travel through an analyser (electromagnetic/quadrupole/the ion trap) that filters the ions based on mass to charge ratio. The filter continuously scans through the range of masses as the stream of ions come from the ion source. A detector counts the number of ions with a specific mass. This information is sent to a computer and a mass spectrum is created. The mass spectrum is a graph of the number of ions with different masses that traveled through the analyser.

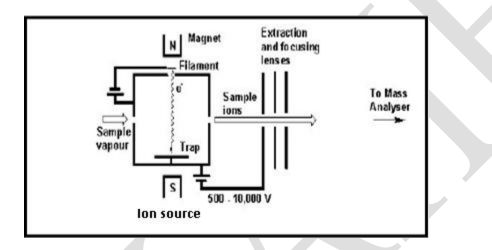


Figure 1. Schematic diagram of EIMS

### **Chemical Ionization (CI) technique**

CI technique is especially useful when no molecular ion is observed in EI mass spectrum of a compound, and also in the case of confirming the molecular weight of a compound. CI technique uses nearly the same ion source device as in EI, except, CI uses tight ion source, and a reagent gas (Figure 2).

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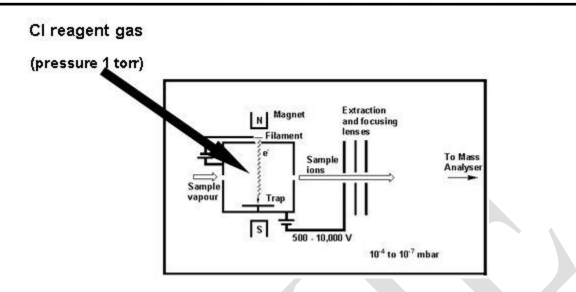


Figure 2. Schematic diagram of CI interface

Reagent gas (e.g. methane, iso-butane and ammonia) is first subjected to electron impact to yield reagent gas ions. These initial reagent gas ions further undergo ion-molecule reactions with neutral reagent molecules (G) to yield reagent selective ions (reagent plasma, e.g., GH+). When sample is introduced, the sample molecules (M) undergo ion-molecule reactions with reagent plasma to produce sample ions. In general, reagent gas molecules are present in the ratio of about 100:1 with respect to sample molecules. Pseudo-molecular ions, [M+H]+ (positive ion mode) or [M-H]- (negative ion mode) are often observed. Unlike in EI method, the CI process is soft ionization and yields abundant quasi-molecular ions, with less fragment ions.

Positive ion mode: GH+ Μ MH+ G +----> Negative mode: [G-H]-+Μ [M-H]-G ion ----> +The fragmentation pattern of protonated molecules obtained under CI conditions may be different from that of the molecular ions observed under EI conditions. In CI mass spectrometry the molecules of a vaporized sample are ionized by a set of reagent ions (reagent plasma) in a series of ion-molecule reactions. The energy transferred by these reactions is lower than the energy imparted by electrons in EI source, and therefore fragmentation of the sample molecules is greatly decreased. For this reason CI mass spectrometry has been finding increasing use as a

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tool for the molecular weight confirmation and for elucidation of structure of variety of organic compounds including differentiation of isomeric compounds. Generally hydrogen (H2), methane (CH4), isobutane (iso-C4H10) and ammonia (NH3) are used as reagent gases in CI mass spectrometry; with all these CI gases the compounds form protonated molecule ion in their CI spectra.

### **Electrospray Ionization (ESI)**

ESI technique involves spraying of a solution of the sample through a highly charged needle so-called capillary which is at atmospheric pressure (Figure 3). The spraying process can be streamlined by using a nebulizing gas. The charged droplets are produced in which the positive or negative ions are solvated with solvent molecules. Heat gas or a dry gas, usually called as desolvation gas is applied to the charged droplets to cause solvent evaporation. The desolvation process decreases the droplet size, leads to the columbic repulsion between the charges present in the droplet and further the droplet fission leads to the formation of individual gas phase analyte ions (that critical point known as the Rayleigh limit), that are guided through ion optics into the mass analyzer. ESI can produce singly or multiply charged ions. The number of charges retained by a particular analyte depends on several factors such as the size, chemical composition, and higher order structure of the analyte molecule, the solvent composition, the presence of co-solutes, and the instrument parameters. For small molecules (< 2000 Da) ESI typically generates singly, doubly, or triply charged ions, while for large molecules (> 2000 Da) ESI can produce a series of multiply charged ions.

CLASS **COURSE NAME: GENOMICS AND PROTEOMICS** : II B. Sc., BT BATCH : 2018 - 2021 **COURSE CODE** :18BTU403 : V (Introduction to Proteomics) UNIT Cone (counterelectrode) Taylor multiply Spray needle tip cone charged droplet the 'Ravleigh' limit is reached analy

ESI is very suitable for a wide range of biochemical compounds including peptides and proteins, lipids, oligosaccharides, oligonucleotides, bio-inorganic compounds, synthetic polymers, and intact non-covalent complexes.

ions

midh

charged droplet

### Atmospheric pressure chemical ionization (APCI) technique

analyt

molecule

Power Supply

►ve

APCI has also become an important ionization source because it generates ions directly from solution and it is capable of analyzing relatively non-polar compounds. Similar to electrospray, the liquid effluent of APCI (Figure 4) is introduced directly into the ionization source. The droplets are not charged and the APCI source contains a heated vaporizer, which facilitates rapid desolvation/vaporization of the droplets. Vaporized sample molecules are carried through an ion-molecule reaction region at atmospheric pressure. APCI ionization originates from the solvent being excited or ionized from the corona discharge. Because the solvent ions are present at atmospheric pressure conditions, chemical ionization of analyte molecules is very efficient; at atmospheric pressure analyte molecules collide with the reagent ions frequently.

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Skimmers lons Spray Needle/Capillary Transfer Analyte/Solvent capillary droplets (spray) Analyte/Eluent To Mass Spectrometer flow .......... syringe pump or LC) N<sub>2</sub>(g) flow Cone Heater Nebulizer gas Corona dsicharge Pumping ATMOSPHERIC PRESSURE

In general, proton transfer occurs in the positive mode to yield [M+H]+ ions. In the negative ion mode, either electron transfer or proton loss occurs to yield M- . or [M-H]- ions, respectively. The moderating influence of the solvent clusters on the reagent ions, and of the high gas pressure, reduces fragmentation during ionization and results in primarily intact quasi-molecular ions. Multiple charging is typically not observed presumably because the ionization process is more energetic than ESI.

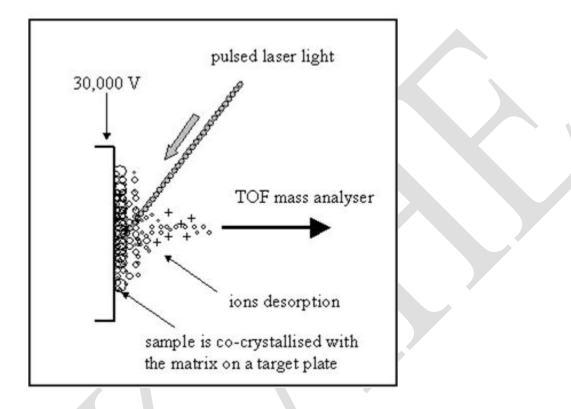
### MALDI technique

Matrix-assisted laser desorption/ionization (MALDI) is a technique to allows the high molecular weight compounds such as organic macro molecules and labile bimolecular into the gas phase as intact ions. MALDI is one of the recent developments of soft ionization techniques in the field of mass spectrometry. It can desorb intact analyte molecular ions with relative masses up to 300KDa. In MALDI-MS analysis, the analyte is first co-crystallized with a larger excess of a matrix compound (CHCA, DBA, Sinapic acid etc), after which, on laser radiation of this matrix-analyte preparation results in desorption of the matrix as a plume, which carries the analyte along with it into gas phase (Figure 5). Thus the matrix plays a key role by strongly absorbing the laser light energy and causing, indirectly, the analyte to vaporize. The matrix also

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serves as a proton donor and acceptor, acting to ionize analyte in both positive and negative ionization modes, respectively. The TOF analyzers are typically used with the MALDI ionization source.



Schematic diagram of MALDI source

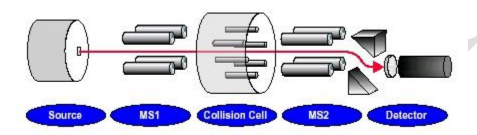
# Tandem Mass spectrometry/Collision induced dissociation (CID)

The general approach of CID mass spectrometry (MS/MS), in modern terms known as product ion scanning using a triple quadrupole mass spectrometer is shown in Figure 6, and it is routinely used for primary structure determination. The first mass analyzer (MS1) is used to select ion of interest specifically from those transmitted by the ionization source. The precursor ion is passed into the collision cell where it undergoes low-energy collisions with an inert gas (argon or nitrogen) to induce fragmentation. The second mass analyser (MS2) acquires the m/z ratio for all the ions exiting the collision cell. For an MS/MS experiment performed using a

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triple-quadrupole equipped with an ESI source, a quadrupole/hexapole mass analyzer acts as the collision cell. Using the second quadrupole as a collision cell enables the re-focusing of scattered ions following fragmentation. Similarly, for MS/MS experiments using a quadrupole-time of flight (TOF) mass spectrometer, the precursor is selected using a quadrupole analyzer and the product ions that are formed in the collision cell are analyzed by TOF (Figure 7). Precursor ion scanning is used to confirm the identity of compounds from a mixture that result in a common daughter ion. For precursor ion scanning, the second mass analyzer (MS2) is fixed to only monitor and transmit product ions of a specific m/z ratio. The first mass analyzer (MS1) is set to scan the whole mass range that includes all the precursor ions whose fragmentation would result in the selected daughter ion.

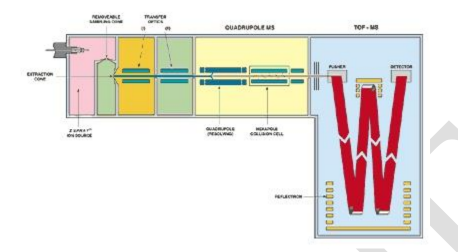


Schematic of a triple quadrupole mass spectrometer

Prepared by Dr. T. Sivaraman, Professor, Department of Biotechnology, KAHE

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### Schematic of a quadrupole-TOF mass spectrometer

### Possible questions <u>1 Mark questions</u>

- 1. Electrophoretic mobility is directly proportional to
- (a) Field strength
- (b) Molecular weight
- (c) Molecular structures
- (d) Solvent viscosity

2. Electrophoretic mobility is inversely proportional to

- (a) Electrostatic potential
- (b) Overall charge of proteins
- (c) Molecular structures
- (d) Field strength

3. The pH at which a protein assume zero net charge is its

- (a) pK<sub>1</sub>
- (b) pK<sub>2</sub>
- $(c) Pk_R$
- (d) pI

4. Separation of protein molecules in SDS-PAGE is on the basis of

- (a) Shapes.
- (b) Size

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(c) Charge

(d) Shape, size and Charge.

- 5. Separation of protein molecules in IEF is on the basis of
- (a) Shapes.
- (b) Size
- (c) Charge
- (d) Shape, size and Charge.
- 6. How does acrylamide affect pore sizes of SDS-PAGE?
- (a) When acrylamide is less, pore sizes are bigger.
- (b) When acrylamide is less, pore sizes are smaller.
- (c) When acrylamide is more, pore sizes are bigger.
- (d) When acrylamide is more, pore sizes are smaller.

7. What are ampholytes?

- (a) Poly amino and poly carboxylic compounds.
- (b) Poly amino compounds.
- (c) Poly carboxylic compounds.
- (d) None of the above.
- 8. The SDS-PAGE is very useful to determine
- (a) Approximate molecular weight of a protein.
- (b) Structural stability of a protein.
- (c) Overall shape of a protein.
- (d) Overall net charge of a protein.

9. In MALDI-TOF, molecules are separated by subjecting to

(A) Electric field only (B) Magnetic field only (C) Both electric and magnetic field

(D) pH gradient

- 10. Isocratic elution in which the concentration of mobile phase is
  - (A) Constant (B) Variable (C) Neither constant nor variable
  - (D) Either constant or variable
- 11. The essential feature for a compound to be used as a matrix in MALDI is(A) Highly volatile (B) Highly stable (C) Highly soluble (D) Highly reactive

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## 2/6/8/10 Marks Questions

- 1. Write any two unique features of ampholytes.
- 2. What are the components in the SDS-PAGE 'loading buffer'?
- 3. What is isotachophoresis?
- 4. How will you optimize the pore size in PAG?
- 5. Write the unique feature of MALDI-TOF.

## 6/8 Marks Questions

1. Explain an electrophoresis method to identify proteins from a mixture containing five molecules which are similar in their size and shape but differing from each other in their pI values (3.5, 5.5, 7.0, 9.4 and 11.2).

2. Write an essay on principles, experimental set-up and applications of Isoelectric focussing electrophoresis technique.

3. Write an essay on principles, experimental set-up and applications of SDS-PAGE

4. Describe the principles of MALDI-MS/MALDI-MS-TOF and its applications on analyzing protein-protein interactions in a detailed manner.

Questions	Option 1	Option 2	Option 3	Option 4	Answers
Unit I					
The term 'genomics' was coined by	Tom Roderick	Anselme Payen	Wilhelm Johannsen	Paulien Hogeweg	Tom Roderick
Pseudogenes	Are nonfunctional genes	Are expressed genes	Are incomplete genes	Are complete genes	Are nonfunctional genes
What do you mean by 'epistasis'?	structural genes	effect of one gene on another	functional genes	all the above	effect of one gene on another
What is pleiotropy?	one gene affecting only one trait	a gene affecting no trait	one gene affecting more than one trait	two genes affecting one trait	one gene affecting more than one trait
The HGP was completed in	2002	2003	2004	2005	2003
Nucleic acds are made up	Nucleotides that are linked by peptded bonds	Nucleotides that are linked by glycosidic bonds	Nucleotides that are linked by ester bond	Nucleotides that are linked by phosphodiester bond	Nucleotides that are linked by phosphodiester bond
	technique used to determine the sugar sequence in a DNA molecule	technique used to determine the phosphate sequence in a DNA molecule	technique used to determine the baase sequence in a DNA molecule	technique used to determine the amino acid sequence in a protein	technique used to determine the phosphate sequence in a DNA molecule
DNA sequencing refers to chemical nucelotide sequencing method?	Sanger method	Maxam-Gilbert Method	Edmans method	Automated sequencing method	Maxam- Gilbert method
Gilbert method for P32 labelling of the DNA at 3' end is	polynucelotide kinase	alkaline phosphatase	exonuclease	terminal nucelotidyl transferase	polynucelotide kinase
chemcial treatments are required in Maxam-Gilbert method?	1	2	3	4	4
Maxam-Gilbert method	formic acid	hydrazine	Dimethyl sulphate	piperdine	Dimethyl sulphate

The principle of sanger's method	use of chemicals for	Use of dNTPs for	Use of ddNTPs for	use of For P32 Chain	Use of ddNTPs for chain
relies on	base specific cleavage	chain termination	chain termination	termination	termination
The sample in sanger's method	AGE	PAGE	PFGE	2-D gel	PAGE
after reaction are separated				electrophoresis	
Which of the following sequencing methods uses PCR	Sanger's method	Sanger's method	LMPCR	LMPCR and automated DNA	LMPCR
Automated DNA sequencing is	ddNTPs are used for	PCR is used for	Fluorescently labelled	Fluorescently labelled	Fluorescently labelled
an improvement of Sanger's	chain termination	making sequencing	ddNTPs are used for	ddNTPs are used for	ddNTPs are used for chain
method where		templates	chain termination	chain termination	termination
Which of the following is not a DNA sequencing method?	LMPCRR	Edmans method	Sanger's method	Maxam-Gilbert method	Edman's method
GFP protein was originally	A.thaliana	A.victoria	C.elegans	D.melanogaster	A.victoria
The first crop plant genome	Maize	A.thaliana	Rice	Barley	A.thaliana
The year of launching of Human	1982-2003	1990-2000	1990-2001	1991-2001	1982-2003
Genome project and completion		DOD		D 4 DD	DCD
The name Kary Mullis is associated with	RFLP	PCR	Chain termination method	RAPD	PCR
In Maxam-gilbert method, chemical used for cytosine	Formic acid	hydrazine	Dimethyl sulphate	piperdine	hydrazine
The chemical used for strand cleavage in Maxam-Gilbert	Piperdine	hydrazine	formic acid	DMSO	Piperdine
ddNTP is different from dNTP	H in place of OH in 3	OH in place of H in 3	OH in place of H 2	CH3 in place of OH in	H in place of OH in 3
in having	position of dNTP	position of dNTP	position of dNTP	3 position of dNTP	position of dNTP
Sanger sequenced the first	T4 phage	C. elegans	TMV	ΦX 174	oX 174
Pseudogenes are important to	they are silent genes	they did not exist until	they occur entirely	they tend to be	they are silent genes that
evolution because	that have been	recently	within multigene	transcribed more	have been activated by
	activated by mutations		families	frequently	mutations
Tandem clusters occur most	satellite DNA	centromere	the nuclear organizer	plasmids	the nuclear organizer
commonly in			regions		regions

A recombinant vector in a yeast genomic library may contain more than one gene. This is not the case wit the recombinant vectors in a yeast cDNA library unlike a genomic DNA library particular care must be taken to ensure that the cDNA library represents every gene in the genome. Why is this the case?	every cDNA fragment cloned into each vector carries only one gene some genes are transcribed only in certain cell types or at low levels	different restriction enzymes are used Host cells are more likely to reject vectors with cDNAs as compared to genomic DNA	the host cells cannot be transformed with a vector carrying more than one gene It is harder to clone cDNA fragments as compared to genomic fragments in to the vectors	the vectors used in making a cDNA library are different a cDNA library lacks introns	every cDNA fragment cloned into each vector carries only one gene some genes are transcribed only in certain cell types or at low levels
Many cDNA libraries are used as expression libraries. The vector chosen for use in an expression library must have additional DNA sequence that is	an origin of replication	a second selectable marker	additional restriction enzyme sites	a promoter sepecific for the host organism	a promoter sepecific for the host organism
Whole-genome shotgun sequencing requires	Construction of a genomic library of plasmid vectors	Random sequencing of DNA fragments	Fragment alignment and gap closure	cloning in to vectors and sequencing using shot-gun	Fragment alignment and gap closure
what information is absent from the sequence of a cDNA clone?	intron sequence	promoter sequence	amino acid sequence of the encoded polypeptides	intron and promoter sequences	intron and promoter sequences
What is the function of the ampr gene in a vector?	To allow resistant transformants to grow in selective medium	To distinguish introns from exons	To allow viral replication	To screen for vectors with inserts	To allow resistant transformants to grow in selective medium
The lacz gene is sometimes included in a cloning vector. What is its purpose?	allow resistant transformants to grow in selective medium	distinguish introns from exons	allow viral replication	screen for vectors with inserts	screen for vectors with inserts
Which vector is appropriate for subcloning an insert of 3 Kb? The goal ofis to determine the location of specific genes	ampr plasmid cloning	YAC annotaion	cDNA proteomics	cosmid genomics	ampr plasmid annotaion

An open reading frame (ORF) is	the sequence of a complete genome	a plasmid vector used in genomic sequencing		a fragment of a genome	a possible gene predicted by DNA sequencing
The whole genome shot gun sequencing approach depends priamrily on	rapidly sequencing thousands of small randomly cloned fragments	methodical sequencing a few large cloned fragments of DNA	sequencing the bacterial chromosome while it is still intact	fragmenting the genome and cloning	rapidly sequencing thousands of small randomly cloned fragments
How are the four different bases distinguished in automated sequencing systems?	each base has a different radioactive tag	each base has a distinctive fluorescent tag	each base has a unique antibody bound to it	each base tagged with antigen to detect by electrophoresis	each base has a distinctive fluorescent tag
proteomics is	a branch of quantum physics	the study of algal genomes	the study of the entire collection of proteins expressed by an organism	study of entire set of genomes	the study of the entire collection of proteins expressed by an organism
The enzyme reverse transcriptase enables scientists	Restriction endonucleases	cDNA molecules	Restriction fragment length polymorphisms	mRNA transcripts	cDNA molecules
Oligo nucelotide gene pobes are defined as what?	Enzymes that recognize and subsequently degrade foreign DNA	The pieces of DNA produced by restriction endonucleases	A short stretch of DNA of a known sequence that will bnase-pair with a complementary		A short stretch of DNA of a known sequence that will base-pair with a complementary sequence
The techniques that utilizes probes to detect specific DNA	Southern blot	Northern blot	Northwestern blot	eastern blot	Southern blot
Which of the following statements regarding the polymerase chain reaction is intrue?	It can increase the amount of DNA in a sample	It has the potential of diagnosing an infection from a single copy of a gene	It utilizes DNA polymerases from psychrophilic organisms	It can amplify DNA of only a few base pairs up to a whole genome	It utilizes DNA polymerases from psychrophilic organisms
When a gene has to be studied in detail, selected gene is removed from an animal, plant and	A primer	an oligonucleotide	a palindrome	a vector	Vector

Genomic libraries are useful for	Periodicals on	collections of isolated	Instructional	Information relating	collections of isolated
obtaining what product?	genomics research	genes	information on how to locate the exact site of the gene of interest	to primers and PCR	genes
Good cloning vectors must possessall but which of the following qualitites?	They should possess their won origin of replication	They should be readily accepted by the cloning host	They should be easily manipulated	They should be resistant to restriction endonucleases	They should be resistant to restriction endonucleases
The insertion of a cloning vector into a cloning host typcially	Transduction	polymerase chain reaction	Tranformation	Conjugation	Tranformation
Which of the following is NOT a step in whole genome shotgun sequencing?	library construction	sequencing of randomly produced fragments	fragment alignment and closure	library construction, sequencing of randomly produced fragments alignment	library construction, sequencing of randomly produced fragments alignment and editing
DNA was first isolated in the	17th century	18th century	19th century	20th century	19th century
Sizes of genomes of free-living organisms have been found to	2-200 Mbp	0.5-200,000 Mbp	100-200000 Mbp	0.5-1000 Mbp	0.5-200,000 Mbp
The size of the human genome is	0.5 Mbp	3,000 Mbp	10 Mbp	500,000 Mbp	3000 Mbp
Most sequences in the human	Genes	Pseudogenes	Interspersed repeats	Tandem repeats	Interspersed repeats
Which genome has the fewest introns?	Drosophila genome	human genome	Maize genome	E.coli genome	E.coli genome
Gene density can be high	In telomeres	anywhere on the chromosomes	in centromeres	in anaphase chromosomes	anywhere on the chromosomes
This vector can be used to construct genomic libraries	phasmid	phagmid	cos site	BAC	BAC
mRNA can be readily isolated from lysed eukaryotic cells by adding magnetic beads which	Oligo (dT)	Oligo (dG)	Oligo(dC)	Oligo (dA)	Oligo (dT)
This enzyme is used to dephsophorylate the vector	Terminal transferase	Alkaline phosphatase	Klenow enzyme	DNA ligase	Alkaline phosphatase
usually nucleic acid sequences are added to create sticky ends	linkers	sequencers	promoters	enhancers	linkers

Minisatellites are	10-40 bp sized short	Short coding	short non-coding	Are regions of	short non-coding repetitive
	sequences with in	repetitive regions on	repetitive sequences	chromosomes after	sequences present through
	genes	the eukaryotic genome	1 1		out the chromosome
	0.1		chromosome	, , , , , , , , , , , , , , , , , , ,	
Each individual has a unique	number of	location, size and	number of fragmented	fragemented number	location, size and number
DNA fingerprinting as	minisatellites or	number of the	DNA	of clones	of the minisatellites on
individuals differ in	chromosomes	minisatellites on			chromsomes
		chromsomes			
Protein coding genes can be identified by	transposon tagging	ORF scanning	Zoo-blotting	Nuclease S1 Mapping	ORF scanning
The functions of the genes can	Gene inactivation	Exon trapping	Zoo-blotting	Northern analysis	Gene inactivation
Reporter genes	Indicate the presence	can often be detected	are all of bacterial	are used to	can often be detected by
	of stress conditions	by histochemical	origin	characterize proteomes	histochemical assays
		assays			
Chromosoem walking	Is used in genetic	Occurs in mitosis	requires a genomic	can be sued to close	requires a genomic library
	mapping		library and done by	physical sequence	and done by PCR
			PCR	maps	
Unit II					
What are amphiphilic				Having both polar and	Having both polar and non-
molecules?	Highly polar	Highly non-polar	Neutral	non-polar groups	polar groups
The shape of a water molecule is	Linear	Trigonal	Tetrahedron	Distorted Tetrahedron	Distorted Tetrahedron
In taxonomy, classifications of					
organisms are on the basis of	Overall morphology	Proteomics	Genomics	Evolution	Overall morphology
In phylogeny, classifications of					
organisms are on the basis of	Overall morphology	Proteomics	Genomics	Evolution	Evolution
The human genome consists of					
base pairs.	two billion	ten billion	one billion	three billion	three billion
Large datasets can be retrieved					
using the tool.	BioMart	SQL	NCBI	FASTA	BioMart
combines transcription factor					
binding sites database search					
with a comparative sequence	rVISTA	mVISTA	wgVISTA	phylo-VISTA	rVISTA

wgVISTA allows the alignment					
of sequences up to 10Mb long	100Mb	10Mb	50Mb	25Mb	10Mb
ENSG### is	Ensembl Exon ID	Ensembl Gene ID	Ensembl Transcript ID	Ensembl Peptide ID	Ensembl Gene ID
ENSE### is	Ensembl Exon ID	Ensembl Gene ID	Ensembl Transcript ID	Ensembl Peptide ID	Ensembl Exon ID
ENST### is	Ensembl Exon ID	Ensembl Gene ID	Ensembl Transcript ID	Ensembl Peptide ID	Ensembl Transcript ID
ENSP### is	Ensembl Exon ID	Ensembl Gene ID	Ensembl Transcript ID	Ensembl Peptide ID	Ensembl Peptide ID
tool in Ensembl	BioMart	SQL	NCBI	FASTA	BioMart
interface for downloading					
datasets using complex queries.	BioMart	SQL	NCBI	FASTA	BioMart
FTP server which can be used to					
download entire	BioMart	MySQL	NCBI	FASTA	MySQL
human enhancers is available					
through Enhancer	VISTA	SQL	NCBI	FASTA	VISTA
calculating sequence similarity					
between biological sequences.	VISTA	NCBI	BLAST	FASTA	BLAST
identified by	Transposon tagging	ORF scanning	Zoo-blotting	Nuclease S1 mapping	ORF scanning
	F	8	8	A northern	
	RNA interference is	Homologous	Transposons can be	hybridization	Transposons can be
Which of the following	not possible in	recombination can be	directed to disrupt	identifies genes that	directed to disrupt specific
statements are not true	prokaryotes	used to disrupt genes	specific genes	are transcribed	genes
determined by	Gene inactivation	Exon trapping	Zoo-blotting	Northern analysis	Gene inactivation
Dideoxynucleotides are used in	Transformation	Cloning	Zoo-blotting	DNA sequencing	DNA sequencing
			Can have a tertiary	Can fold into a double	Can have a tertiary
Polypeptides	Consist of nucleotides	Can contain phosphate	5	helix	structure
			Are used to delineate		Are used to delineate
	Indicate the presence	Are used to	regulatory sequence	Are all of bacterial	regulatory sequence
Reporter genes	of stress conditions	characterize proteomes	<b>C I</b>	origin	elements
		r			
	Are used for analysis		Are smaller than DNA	Contain DNA	
Microarrays	of transcriptomes	Are made of glass	chips	sequences	Contain DNA sequences
	r r r	to find intergenic	to find gene	to find protein-coding	to find protein-coding
ORF scanning is used	to find exons	sequences	homologies	genes	genes
6		1			
		Is used to find	Is used to find gene		
Chromosome walking	Occurs in mitosis	intergenic sequences	homologies	Can be done by PCR	Can be done by PCR

	Is used in genome	Is used to find	Is used to identify	Is found in intergenic	
A codon bias	mapping	intergenic sequences	genes	regions	Is used to identify genes
Human Genome Project was	11 0		0		
first initiated in the year	1982	1972	1970	1980	1970
Human genome was completed					
by the year	2002	2000	2003	2005	2003
A physical map is a		nucleotides between			
representation of	physical distance	genes	genetic markers	all the above	all the above
genome makes it easier for		nucleotides between			
researchers to study	physical distance	genes	genetic markers	individual genes	individual genes
collective genomes of multiple					
species that grow and interact in					
an environmental niche.	Metagenomics	Proteomics	Genomics	Evolution	Metagenomics
Ais the entire set of					
proteins produced by a cell type.	Metagenome	Paragenome	proteome	genome	proteome
What is the most challenging issue facing genome sequencing?	the inability to develop fast and accurate sequencing techniques	genomes at the	the availability and stability of DNA	all of the above	the ethics of using information from genomes at the individual
Genomics can be used in	generate new hybrid	improve disease			
agriculture to:	strains	resistance	improve yield	all of the above	all of the above
studied using genome-wide association studies?	viral diseases	single-gene inherited diseases	diseases caused by multiple genes	diseases caused by environmental factors	diseases caused by multiple genes
Drosophila melanogaster is a	viral diseases spreader	bacterial diseases spreader	model organism	all the above	model organism
to overcome this genetic					
disorder.	kidney	liver	bone marrow	none of the above	bone marrow
·	Human	Invertebrates	vertebrates	all the above	vertebrates
Comparative genomics helps us to better understand	Vertebrate evolution	Differences between species at the genome level	Identification of highly conserved regions	all the above	all the above

effort to identify and catalogue					
genetic similarities and					
differences in populations	VISTA	NCBI	НарМар	FASTA	НарМар
	Lymphoblastoid cell			Human Umbilical	
CD4 is which binds to the SNPs as it is	line	Embryonic Stem Cells	T cells	Vein Endothelial Cells	T cells
which binds to the SNPs as it is					
specific.	Tab Assay	Tap-Man Assay	Taq-Man Assay	none of them	Taq-Man Assay
Genomics is the study of	entire genomes	cheomosome	cell	DNA	entire genomes
of the physical distance between					
genes or genetic markers	physical map	physical distance	physical signature	none	physical map
with the genome is called as					
	pharmacogenomics	genomics	model gemomics	proteomics	pharmacogenomics
location on a chromosome is					
called as	physical map	genetic map	protemics	toxicogenomics	genetic map
traits of organisms	physical map	genetic map	Genome mapping	toxicogenomics	genome mapping
Ensembl Gene ID is	ENSG###	ENSE###	ENST###	ENSP###	ENSG###
Ensembl Exon ID is	ENSG###	ENSE###	ENST###	ENSP###	ENSE###
Ensembl Transcript ID is	ENSG###	ENSE###	ENST###	ENSP###	ENST###
Ensembl Peptide ID is	ENSG###	ENSE###	ENST###	ENSP###	ENSP###
with the genome is termed as					
	toxicogenomics	genomics	model gemomics	proteomics	toxicogenomics
Large datasets can be retrieved					
using the tool.	BioMart	SQL	NCBI	FASTA	BioMart
effort to identify and catalogue					
genetic similarities and					
differences in populations	VISTA	NCBI	НарМар	FASTA	НарМар
		the ethics of using			
What is the most challenging	the inability to develop	0			the ethics of using
issue facing genome	fast and accurate	genomes at the	the availability and		information from genomes
sequencing?	sequencing techniques	individual	stability of DNA	all of the above	at the individual
genome makes it easier for		nucleotides between	-		
researchers to study	physical distance	genes	genetic markers	individual genes	individual genes
Polypeptides can have	nucleotides	phosphate	tertiary structure	double helix	tertiary structure

Having a complete map of the		nucleotides between			
genome makes it easier to study	physical distance	genes	genetic markers	individual genes	individual genes
<u> </u>	r Jana a ann a s	find intergenic	6	0	
A codon bias is used to	genome mapping	sequences	identify genes	intergenic regions	identify genes
	8FF8			8	
Unit III					
most ideal for understanding the					
nature of genes?	Linkage maps	Physical maps	Sequence Maps	fingerprint maps	Sequence maps
Which of the following	Humans share	Genetic screening of	Human chromosome	A large amount of	Human chromosome are
statements regarding the	approximately 80 % of	families for inheritable	are capable of walking	human genome	capable of walking
findings of the human genome	their gene sequence	diseases may become		contains DNA	
project is incorrect?	with mice	possible		sequences that do not	
				code for cell protein	
	frequently found in	always smaller than		movable DNA	
Microsatellites are	bacterial genomes	10 bp	used as DNA markers	elements	used as DNA markers
show easily detectable					
difference among different					
strains of a species of among	DNA fingerprinting	molecular markers	moelcular scissors	RFLP	molecular markers
Molecular markers include	RFLP	PCR	alkaline phosphatase	phosphatase	RFLP
Molecular markers are used to					
construct	Chromosome maps	cytogenetic maps	physical maps	geographic maps	physical maps
The variation in the restriction					
DNA fragment lengths between	Retriction fragment			simple sequence	<b>Retriction fragment length</b>
individuals of a species is called	length polymorphism	RAPD	AFLP	repeats	polymorphism
	used to identify a	used to identify a	used to identify a	used to identify both	used to identify a sepcific
RFLP involves	specific protein	sepcific DNA	sepcific RNA	DNA and RNA	RNA
Locations of quantitative genes				construct or position	
on chromosomes are called	Qualitative trait loci	Quantitative trait loci	maps	of gene	Quantitative trait loci
	DNA sequencing	Restriction digestion			
RAPD is a	based method	based method	PCR based method	enzyme based method	PCR based method
using random primers in a PCR					
reaction is called	RAPD	RFLP	AFLP	Insitu hybridization	RAPD

All the following statements are	RAPD is a quick		Species specific	Radioactive probes	
true regarding RFLP and RAPD	method compared to	RFLP is more relible	primers are required	are not required in	Species specific primers
except	RFLP	than RAPD	for RAPD	RAPD	are required for RAPD
has several repeating units of					
short sequences called	random repeats	tandem repeats	mini satellites	microsatellites	Tandem repeats
tandem repeats between two or					
more individuals is called	VNTRs	RFLP	SSRs	AFLP	VNTRs
	1-6 bp long ssequences distributed				1-6 bp long ssequences
Simple sequence repeats SSRs	along the	also called as satellite	not specific in number		distributed along the
are	chromosomes	repeats	and position	not used as markers	chromosomes
	emoniosonies	repetition		not used us markers	
		it is not associated	method that detects		method that detects the
	not a PCR based	with polymorphism of	the presence or	not specific in	presence or absence of a
AFLP is a	methods	DNA	absence of a fragment	position and number	fragment
distinguish one individual from					
another one is called	variant fragment	marking fragment	differing fragment	variable repeats	Marking fragment
		Karyotyping analysis	Analysis of the		
		in human chromosome			
	hybridization studies		transgenic strain using		
	in Drosophila		transgenic mice		
	determined the				hybridization studies in
	location of the yellow				Drosophila determined the
	gene, which influences				location of the yellow gene,
	body color, to be near				which influences body
Which of these is an example of	the tip of the X			banding pattern	color, to be near the tip of
cytopgenetic mapping?	chromosome			techniques	the X chromosome
Cytologistics can use which of					
the following to describe		protein expression		compariosn to	Den l'accestione of
locations of a gene at a specific	DNA sequence of a	from a chromosomal	Banding pattern of	markers located with	Banding pattern of
palce on the chromosome?	chromosomal region	region	stained chromosomes	in a few thoudsand bp	stained chromosomes

		Is a high resolution	Can be used to	banding pattern	
	Requires that gene	method of gene	determine relative	techniques	
	have been cloned if in situ hybridization is to	localization	order of genes located very close to each	1	Requires that gene have been cloned if in situ
Cytogenetic mapping:	be used		other		hybridization is to be used
Which of the following questions could be easily answered by FISH?	Recombination occurs between species	Is the mutations due to a deletion of the entire gene?	How large DNA would be need to clone	position of genes	Is the mutations due to a deletion of the entire gene?
Which of these is a key characterisitic of a molecular marker?	It is a known gene	It is located at a known site on the chromosome	It is only useful for linkage and physical mapping studies	positional analysis	It is located at a known site on the chromosome
amplified by PCR and is polymorphic by length is	RFLP	VNTRs	AFLP	SNP	AFLP
A polymorphism is	Any change in the DNA sequence	The most common variation of a gene or marker sequence	The least common vaariation of a gene or marker sequence	Variation of gene or marker sequence present in > 1% of the population	Variation of gene or marker sequence present in > 1% of the population
Which of the these statements regarding RFLP analysis is correct?	RFLP analysis requires southern blotting for detection of fragments	RFLPs can identify single base pair changes at any site in the chromosomes	An RFLP typically rpoduces several different alleles	positional analysis of the gene	RFLP analysis requires southern blotting for detection of fragments
A monomorphic DNA segment	A segment of DNA that exists in many forms in the population	A segment of DNA that controls a single gene function	A segment of DNA inherited in a dominant fashion	A segment of DNA shared by over 99 % of the population	A segment of DNA shared by over 99 % of the population
the distance between which of the following pairs of DNA sequences?	AFLPs and RFLPs	Two AFLPs	Two known genes	A known gene and any type of molecualr marker, AFLP, RFLPs	A known gene and any type of molecualr marker, AFLP, RFLPs
Which of the following would be a reasonable use of an RFLP map?	Identification of the exact location of an unknown gene along the chromosome	Description of the size of the DNA fragment required to clone the region containing an unknown gene	Identification of the region in which an unknown gene is located	positional analysis of genes	Identification of the region in which an unknown gene is located

is most likely to be highly				All other forms of	
polymorphic	An RFLP	A Microsatellite	An SNP	markers	A Microsatellite
Why might use of microsatellite		Microsatellites are		Easy to detect, more	Easy to detect, more
in genetic mapping studies be an	A microsatellite are	more abundant than	Microsatellites with	abundant and have	abundant and have more
advantage over RFLPs?	easier to detect	RFLPs	out potential alleles	more potential alleles	potential alleles
	Relative level of				
how are individual	fluorescence when				Relative level of
chromosomes identified in	stained with a dye	Level of charge			fluorescence when stained
chromosome sorting techniques?	mixture	relativ eto size	Size of the molecule	shape and orientation	with a dye mixture
		A complete mRNA			
	A complete genomci	library	A chromosome		A chromosome specific
Which of these describes a	library including		specific library of	An ordered genomci	library of overlapping
contig?	overlapping clones		overlapping clones	library	clones
suited for creating a contig of					
bovine chromosome 10?	λ phage	A plasmid	YAC	Cos site	YAC
which of the following would		A gene encoding a			A gene encoding a
not be a critical characteristic of		required structural	AN origin of		required structural
a YAC vector?	Telomeric sequences	proteins	replication	A centromere	proteins
mapping ,After identifed small		The DNA sequence of			
area of chromosomes which	The clone will also	the lone will contain	Cytological		
contains the gene of interset, by	contain the nearest	an ORF	hybridization of the		
contig maps the positional	marker mapped by		clone produces a		The DNA sequence of the
cloning of the gene is identified	linkage analysis		different pattern	Fragmental analysis	lone will contain an ORF
		Identification of all			
	Correlation of	the known alleles of a			
Which of these is a reasonable	physicla and linkage	single disease-			
use for the contigs and clones	map distances to	containing gene	Insertion in to cells		Correlation of physical
generated in the process of	determine		affected with genetic		and linkage map distances
obtaining a physical map of the	recombination		disease in the process		to determine
human genome?	frequency		of gene therapy	Insertional cloning	recombination frequency

Sequencing of genomes other than humans is potentially valuable because	Disorders of cellualr function can be studies in relatively simple model systems kilobases	Comparison of gene sequences between different species can allow prediction of desase causing mutations	genes may have a similar function in other species, giving us a palce to start with fucntional analysis of similar genes	positional cloning of evolutionary genes	genes may have a similar function in other species, giving us a palce to start with fucntional analysis of similar genes Kilobases
physical maps is A haplotype is	the set of polymorphic nucelotides dound together on a single chromosome	centomorgans a genotype that is unique to non-african populations	cytological bands a genotype that is only found in a single individual in a population	a set of diploid genotypes at two or more loci in an individual	the set of polymorphic nucelotides dound together on a single chromosome
Alternative splicing refers to	a differnec in the number of exons in two or more species	the production of two or more mRNAs from a single gene	regulation of two different genes by a single regulatory element	post-translational modification of the cleavage site of receptor proteins	the production of two or more mRNAs from a single gene
Tagging SNP are designed	to capture the majority of genetic variation in haplotype	reducing the number of sites that must be tested in a genomic scan and the genetic variation in haplotype	positional cloning	functional cloning	reducing the number of sites that must be tested in a genomic scan and the genetic variation in haplotype
genes A and B is 10 map units and the map distance between genes B and C is 25 map units, what is the map distance	15 map units	35 map units	Either 15 map units or 35 map units depending on the order of the genes.	The map distance between A and C cannot be predicted from the data	Either 15 map units or 35 map units depending on the order of the genes.
four linked genes are as follwos: A-B=22 m.u., B-C= 7 m.u., C- D= 9 m.u., A-D=20 m.u., A-C= 29 m.u. What is the order of	ABCD	ADBC	ABDC	BADC	ADBC

genes D and E in a two point test	D and E are on	D and E are linked	D and E are linked		
cross is 50 map units. What does	different pairs of	and exactly 50 map	and at least 50 map		
this mean in physical terms?	chromosomes	units apart	units aprat	Either a or c.	Either a or c.
		grow with in bacteria,			grow with in bacteria, and
	can generally	and are present in			are present in bacterial
	accommodate larger	bacterial colonies on	can accommodate	burst bacteria and	colonies on an agar plate
	inserts than pahge	an agar plate	inserts of over 100	form plaques on a	
plasmid vectors for cloning	vectors can		kilobases	"lawn" bacteria	
		reconstructing the		estimating	
Simple tandem repeat		relationships of		relationships of	
polymorphisms in humans are	solving criminal and	humans and chimps	estimating matches	humans and	solving criminal and
most useful for	paternity cases		for blood transfusions	neanderthals	paternity cases
		proceeded much more			proceeded much more
	has been restricted to	successfully as large	has shown that there	has demonstrated that	
	the sex chromosomes	numbers of DNA	are more genes on the	almost all of the DNA	numbers of DNA markers
Mapping of human	because of small	markers became	Y than on the X	is invloved in coding	became available
chromosomes	family sizes	available	chromosomes	of genes	
		is used to determine			
	uses hybridization to	whether a gene is			uses hybridization to
	detect specific DNA	transcribed in specific	measures the transfer	is used to amplify	detect specific DNA
RFLP analysis is a technique	restriction fragments	cells	frequency of genes	genes for producing	restriction fragments in
that	in genomic DNA		during conjugation	useful products	genomic DNA
		easy identification of		Pieces of DNA from	
		plasmids which carry		different sources to	Pieces of DNA from
	Selection for plasmids	an insert	-	hybridize to each other	
The "sticky ends" generated by	lacking antibiotic		RNA with in bacterial	and to be joined	hybridize to each other
restriction enzymes allow:	resistance		cells	together	and to be joined together
~		are typed by		mapping and a type of	are used in genetic
Simple sequence length		oligonucleotide		mini and micro	mapping and a type of
polymorphisms	can be satellite DNA	hybridization	are not mini satellite	satellite	mini and micro satellite
		requires computers			requires computers
	is used in analyzing		is normally used with	is more accurate than	
The shotgun method	transcriptomes		large genomes	clone contig method	

		are not very common		are normally found at	
	are tandemly repeated	•	are usually longer	the end of	are tandemly repeated
Microsatellites	sequences	C	than 200 bp	chromosomes	sequences
Transcriptomes	Consist of RNA	consist of DNA	do not change	consist of proteins	consist of RNA
_		are used in physical			
	are used to determine	mapping			
	the position of			usually occurs as	
	restriction sites in a		are used in genetic	multiple alleles in a	are used in genetic
RFLPs	genome		mapping	genome	mapping
		web lab			
The term used to refer					
something performed on					
computer or computer			· ·	,.	
simultions	dry lab predicting disease risk		invitro comparative analysis	insilico predicting disease risk	insilico predicting disease risk at
what are the applicatiosn of which of the following is used	predicting disease fisk	Genome wide annotations	comparative analysis	predicting disease fisk	predicting disease fisk at
for determining the location of		annotations			
secific genes with in a genome	Genomcis		cloning	proteomics	Annotations
transmitted togethr exemplies	chromosomal	genetic linkage and			
and a unit of gentic map	interference and chi	chi square	recombination and	genetic linakge and	genetic linakge and
distance is given by	square		centimorgan	centimorgan	centimorgan
known asand	centromere,	centromere and map	chaisma and	centiniorgan	
recombination between genes	chromosomal	units	chromosomal	chiasma and chi	chaisma and chromosomal
can be modulated by	interference		interference	square	interference
linkage can be given byand	chromosomal	null hypothesis and	null hypothesis and	square	null hypothesis and chi
a statistical test of linkage is	interference and chi	chi square	chromosomal	chi square and null	square
called	square	om square	interference	hypothesis	Square
	-Jame			-JP 0410010	
T I \$4 TN7					
Unit IV					
The primary structure is	peptide bond	Hydrogen bond	Ionic bond	Hydrophobic bonds	peptide bonD
acids is an alpha helix					
terminator?	cysteine	alanine	proline	glycien	proline
The secondary structure is					
primarily maintained by	Vander waals force	Hydrogen bond	Ionic bond	covalent bond	Hydrogen bond

most common and stable					
conformation for a poly peptide	Alpha helix	Beta pleated sheets	Beta bends	loops	Alpha helix
	· ·				
Which of the following			Primary structure of	peptide bonds	
statements are true regarding	3D structure of a	it is the biologically	protein deermines the	determines the	
tertiary structure of proteins	protein	active conformation	tertiary structure	tertiary structure	3 D structure of a protein
beta bend is	cystein	glycine	serine	Aspartic acid	glycine
associated with Alzhemiers					
diease is composed of	Alpha helix	Beta pleated sheets	Beta bends	loops	Beta pleated sheets
but recover their activity upon					
folding are	denatured	folded	synthesized	unfolded	folded
resembles	sheet	coiled spring	linear chain	random coil	coiled spring
Chiral centers of polypeptide					
chains can undergo	formylation	amidation	racemization	none of the above	racemization
a distance of	0.54nm	1.5nm	3nm	6nm	0.54nm
the following	Blomycin	Penicillin	Microcystine	none of the above	Penicillin
Which of the given protein					
has a length :widh ratio >10?	Hemoglobin	Myidlobin	Fibrinogen	Lysozyme	Fibrinogen
acids is abundantly found in					
collgen?	Glycine	Serine	Alanine	Tyrosine	Glycine
a confirmatory test for					
trytophan	Sakaguchi test	Ninhydrin test	Hopkin cole test	none of the above	Hopkin cole test
Pulses contain incomplete					
proteins since they lack	Lysine	Tryptophan	Phenyl alanine	Methinine	methinine
Which out of the following is a					
standard amino acid ?	Hydroxy proline	Beta Alanine	Orthnine	Orthnine	Arginine
acid in the struture of	Gluthamine -Glycine-	Glutamic Acid-	Glutamic acid -Glycine-	Glutamic acid -Glycine-	Glutamic Acid-Cysteine-
Glutathione	Cystine	Cysteine-Glycine	Cysteine	Cysteine	Glycine
Which out of the following is		Low density			
not a conjugated protein?	Albumin	lipoprotein	Glycopprotein	Glycopprotein	Albumin
Which out of the following		Minimum			
features is not observred at		Electrohorectic	Maximum	Maximum	
isoelectric pH of a protein?	Maximum solubility	migratio	percipitation	percipitation	Maximum solubility
amino acids acid is a precursor					
of niacin	Tyrosine	Threonine	Tryptophan	Phenyl alanine	Tyrosine

			1		
Which of the following			The cis configuration is		
satetments is true about a		It is capable of forming		nitrogen and the	It is capable of forming
peptide bond(RCONH)	It is non planar	hrdrogen bondings	trans configuration	carbonyl group	hrdrogen bondings
terms refers to the arrangment					
of different protein subunits in a					
multiprotein complex	primary structure	secondary structure	tertiry structure	quaternary structure	quaternary structure
Native state of a protein can				Presence of	Presence of hydrophilic
be disrupted by	Temperature	рН	Removal of water	hydrophilic surfaces	surfaces
	Hemoglobin is a	Hemoglobin is a	Fibrous proteins are	Collagen is a fibrous	Hemoglobin is a fibrous
Identify the wrong statement	globular protein	fibrous protein	insoluble in water	protein	protein
In 3° structure of proteins,	a) Hydrophobic				
folding and shaping is done	interactions	b)Polar interactions		d)None of the	
by			c)Hydrogen bonding	mentioned	Hydrophobic interactions
The binding of 2,3-	an increase in O2	a decrease in	an increase in		
bisphosphoglycerate (BPG) to	affinity for	O2 affinity for	O2 binding	a decrease in O2	a decrease in O2 affinity for
hemoglobin causes	hemoglobin.	hemoglobin.	stoichiometry.	binding stoichiometry.	hemoglobin.
theory, a decrease in the free					
energy of activation (DG‡) will	a decrease in enzyme	a decrease in reaction	an increase in enzyme	an increase in reaction	
result in	binding specificity	rate.	binding specificity	rate.	an increase in reaction rate.
		are very specific and	drive reactions to		
		can prevent the	completion while		
	are consumed in the	conversion of	other catalysts drive	lower the activation	lower the activation energy
Enzymes are potent catalysts	reactions they	products back to	reactions to	energy for the	for the reactions they
because they:	catalyze.	substrates	equilibrium	reactions they catalyze	catalyze
Glycosidic bonds are found in	DNA	RNA	Sucrose	All of above	All of above
A stabilizing interaction found in					
nucleic acids, but not found in		favorable electrostatic	stacking of aromatic	sulfhydryl cross	
proteins is:	C=O < HN H-bonds	interactions	bases.	linking.	stacking of aromatic bases.
membrane proteins that is in			less hydrophobic than	more hydrophobic	more hydrophobic than the
contact with the phospholipid	charged	hydrated	the protein core.	than the protein core.	protein core.
protein tertiary structure are					
the same as those that stabilize				Van der Waals	
ligand binding except for:	disulfide bonds	electrostatics	hydrogen bonding	interactions	disulfide bonds

individual noncovalent forces					
that stabilize protein structure	aromatic-aromatic				aromatic-aromatic
is:	interactions	electrostatics	hydrogen bonding	Van der Waals forces	interactions
responsible for the UV					
absorption of proteins at 280	His	Tyr	Trp	lle	Trp
A hydrogen bond to water that	has no effect on the	decreases the stability	decreases the stability	decreases the stability	decreases the stability of
is broken during folding and not	stability of the folded	of the folded state by	of the unfolded state	of the folded state by	the folded state by 20
reformed	state	2 kJ/mol	by 2 kJ/mol	20 kJ/mol	kJ/mol
globular proteins is tightly					
packed is due to	the hydrophobic effect	hydrogen bonding.	electrostatic effects.	van der Waals forces.	van der Waals forces.
not among the six					
internationally accepted classes	Hydrolases	Ligases	Oxidoreductases	Polymerases	Polymerases
The portion of proteins having					
the highest mobility are	a-helices	b-sheets	peptide bonds	d)surface side chains.	surface side chains.
stabilizes sub units in 4°	hydrophilic		hydrophobic		
structure of proteins are	interactions	hydrogen bonding	interactions	none of above	hydrophobic interactions
		decreases surface area			
Benefit of hydrophobic		to allow interaction			
interaction includes	protein aggregation	with water	both A and B	none of above	both A and B
You find that protein samples		uses a protease	Perform each step as		
losses activity during storage	Add an additional	inhibitor during	quickly as possible in a		uses a protease inhibitor
what can you do about this	purification step	purification steps	cold room	All of them	during purification steps
often considered a suitable			Hydrophobic		
polishing step in a protein	Affinity	Ion - exchenga	interaction	Size exclusion	Size exclusion
purification strategy	chromatography (AC)	chromatography	chromatography	chromatography	chromatography
types are suitable as a capture					
step in the purification of non-				Ammonium suiphate	
tagged proteins	SEC	Dialysis	IEX and HIC	precipitation	Dialysis
could be used to check the					
molecular weight of your		Mass spectometry			
purified protein	SDS PAGE only	only	Analytical SEC only	All of the above	Mass spectometry only

				relative content of	
In isoelectric focusing	relative content of	relative content o		positively and	relative content of
separation of the proteins are	positively charged	negative charged		negatively charged	positively and negatively
basd on	groups	groups	Size	residue	charged residue
listed below which should elute				RNA	
second in size-exclusion		immunoglobulin	ribonucleasae A Mr	polymeraseMr=45000	immunoglobulin
chromatography	cytochrome cMr=1300	GMr=145000	=137000	0	GMr=145000
By adding SDS during the			dertmine the amino	exclusively on the	separate proteins
electrophoresis of proteins it is	determine a proteins	determine an enzymes	acid composistion of	basis of molecular	exclusively on the basis of
possible to :	isoelectric poin	specific activity	the protein	weight	molecular weight
What properties of a protein					
does hydrophophic interaction		Hydrophobic amino			
chromatography expoit for		acids on the protein			Hydrophobic amino acids
purification	charged amino acids	surface	Molecular weight	Enzyme activity	on the protein surface
Which of the following statements is in correct	In affinity chromatography lectins are uesde to purify a glycoprotein	The separation in gel filtration chromatography is based on size shape and net charge of the protein	In ion exchange chromatography the bound priteins are eluted using NaCl solution	All of them	The separation in gel filtration chromatography is based on size shape and net charge of the protein
Which of the following is true? Hemoglobin is a	The disulfide bridges formed by reduction of the sulfhydryl groups on cysteine stabilizes protein tertiary structure monomer	The disulfide bridges formed by oxidation of the sulfhydryl groups on cysteine destabilizes protein tertiary structure Dimer	The disulfide bridges formed by oxidation of the sulfhydryl groups on cysteine stabilizes protein tertiary structure Timer	The disulfide bridges formed by reduction of the sulfhydryl groups on cysteine destabilizes protein tertiary structure Tetramer	The disulfide bridges formed by oxidation of the sulfhydryl groups on cysteine stabilizes protein tertiary structure Tetramer
	monomer	Dimer			retrainer
insulin molecule are held	culfido bridgos	Disulfido bridgo	Dantida hand	Covolont linkogo	Disulfida huidaa
together by	sulfide bridges	Disulfide bridge	Peptide bond	Covalent linkage	Disulfide bridge

Disulfied bridge formed	cysteine residues that	cystine residues that	proline residues that	histidine residues that	cystine residues that are
between	are close together	are close together	are close together	are close together	close together
	Nuclear magnetic				
The 3-D structure of protein can	resonance				Nuclear magnetic
be determined by	Spectroscopy	Raman Spectroscopy	IR spectroscopy	UV Spectroscopy	resonance Spectroscopy
compound is not involved in					
Edman degradation?	Phenylisothiocyanate	CF3 COOH	FDNB	Phenylthiocarbonyl	СҒЗ СООН
thousands of different proteins					
from many species have been					
determined using principles first	Edman	Sanger	Mende	Watson and Crick	Edman
	Identifying N-terminal	Identifying C-terminal		Identifying	Identifying N-terminal
Edman degradation is used for	amino acids	amino acids	Identifying amino acid	carbohydrates	amino acids
Which of the following is the	Cleaving, sequencing	Sequencing, ordering	Ordering, cleaving and	Ordering, sequencing	Cleaving, sequencing and
correct order of sequencing?	and ordering	and cleaving	sequencing	and cleaving	ordering
Which of the following is Edman					
reagent?	Phenylisothiocyanate	CF3 COOH	FDNB	Phenylthiocarbonyl	Phenylisothiocyanate
Unit V					
directly proportional to	Field strength	Molecular weight	Molecular structures	Solvent viscosity	Field strength
Electrophoretic mobility is		Overall charge of			
inversely proportional to	Electrostatic potential	proteins	Molecular structures	Field strength	Molecular structures
The pH at which a protein					
assume zero net charge is its	pK1	pK2	PkR	pI	pI
Separation of protein molecules				Shape, size and	
in SDS-PAGE is on the basis of	Shapes	Size	Charge	Charge.	Size
Separation of protein molecules				Shape, size and	
in IEF is on the basis of	Shapes	Size	Charge	Charge.	Charge
	When acrylamide is	When acrylamide is	When acrylamide is	When acrylamide is	
How does acrylamide affect pore	less, pore sizes are	less, pore sizes are	more, pore sizes are	more, pore sizes are	When acrylamide is less,
sizes of SDS-PAGE?	bigger.	smaller.	bigger.	smaller	pore sizes are bigger

	Poly amino and poly	Poly amino	Poly carboxylic		Poly amino and poly
What are ampholytes?	carboxylic compounds	compounds	compounds	None of the above	carboxylic compounds
	Approximate				
The SDS-PAGE is very useful to		Structural stability of a	Overall shape of a	Overall net charge of a	Approximate molecular
determine	protein	protein	protein	protein	weight of a protein
	Approximate				
The IEF is very useful to	molecular weight of a	Structural stability of a	Overall shape of a	Isoelectric point of a	Isoelectric point of a
determine	protein	protein	protein	protein	protein
Why is sucrose added in the	To denature the	To increase density of	To stabilize the protein	-	To increase density of the
loading buffer of SDS-PAGE?	protein molecules	the protein molecules	molecules	molecules	protein molecules
Why is urea added in the loading		To increase density of	To stabilize the protein	•	To denature the protein
buffer of SDS-PAGE?	protein molecules	the protein molecules	molecules	molecules	molecules
Why is mercaptoethanol added					
in the loading buffer of SDS-	To denature the	To increase density of	To stabilize the protein	1	To reduce the protein
PAGE?	protein molecules	the protein molecules	molecules	molecules	molecules
What are the limitations of 2D		Reproducibility of the			
gel electrophoresis?	Detection procedures	data	Dynamic ranges	All the above factors	All the above factors
What are the limitations of 2D		Reproducibility of the			
gel electrophoresis?	Detection procedures	data	Dynamic ranges	All the above factors	All the above factors
Two-dimensions of the 2D gel		TGGE and SDS-			
electrophoresis are	PAGE and SDS-PAGE		IEF and SDS-PAGE	IEF and PAGE	IEF and SDS-PAGE
gel electrophoresis is	Expensive	Time consuming	Reproducibility	Data analysis	Reproducibility
molecules would be	Fragmented	Condensed	Multiple charged	Uncharged	Multiple charged
In MALDI-TOF, molecules are			Both electric and		
separated by subjecting to	Electric field only	Magnetic field only	magnetic field	pH gradient	Electric field only
compound to be used as a matrix					
in MALDI is	Highly volatile	Highly stable	Highly soluble	Highly reactive	Highly volatile
Name the surfactant used in 2D		7M urea			
gel electrophoresis process	thiourea		100 mM DTT	sulfobetaine	sulfobetaine

		congo red			
Staining used in 2D gel	comassie brilliant	-			comassie brilliant blue
electrophoresis	blue R250 staining		methylene blue	crystal violet	R250 staining
The first dimension separation		gradient pH			
of proteins is based on	isoelectric focussing		amphipathic	hydrophobic	isoelectric focussing
sepration for two dimensional		gradient pH			
electrophoresis is based on	molecular mass		zwitter ion	amphipathic form	molecular mass
		Limited by pH range	Analysis and		
Advantages of two Dimensional	Good resolution of		quantification are	not for hydrophobic	Good resolution of
gel electrophoresis	proteins		difficult	proteins	proteins
		detection of post	Analysis and		
Disadvantages of 2D gel	Good resolution of	translational	Quantification are	Not limited by pH	Analysis and
electrophoresis	proteins	modifications	difficult	range	Quantification are difficult
	Proteins are denatured	not for hydrophobic	smaller proteins		Proteins are denatured by
In an SDS-PAGE	by the SDS	proteins	migrate more slowly	limited pH range	the SDS
Proteins can be visualized	Staining them with	using electron	measuring their		
directly in gels by	the dye	microscope only	molecular weight	limited pH range	Staining them with the dye
	treated with a	fractioned by	treated with an	separated by their	
	reducing agent and	electrophoresisthen	oxidizing agent and	difference in pH	treated with a reducing
	then anionic detergent	treated with an	then with anionic		agent and then anionic
	followed by	oxidising agent	detergent followed by		detergent follwoed by
In SDS-PAGE, the protein	fractionation by	follwoed by anionic	fractionation by		fractionation by
sample is first	electrophoresis	detergent	electrophoresis		eelctrophoresis
		relative content of		relative content of	
In isoelectric focussing, proteins	Relative content of	negatively charged		positively and	relative content of
are separated on the basis of	positively charged	residue		negatively charged	positively and negatively
their	residue		Size of the molecule	residue	charged residue
The subunit molecular weight		Gel filtration	information from SDS		1
as well as the number of		chromatography	PAGE and gel		combining information
subunits in the quaternary	SDS-PAGE		filtration		from SDS PAGE and gel
structure can be determined	electrophoresis	1 1 1 1	chromatography	Isoelectric focussing	filtration chromatography
PAGE experiment on the basis	positively charged	molecular weight	negatively charged	different isoelectric	molecular weight
of their	side chains		side chains	points	

sample are of similar chemical		Adsorption			
type; choose the correct	Partition	chromatography	Chiral	Ion pair	
chromatographic technique for	chromatography		Chromatography	chromatography	Partition chromatography
which of the follwoing statements about the use of mass spectrometry in protein investigation is correct?	Mass spectrometry can be used to investigate post translational modifications	It requires the crystallization of proteins	Mass spectrometry involves ionized molecules in the liquid phase	It separates the proteins on the basis of the size	Mass spectrometry can be used to investigate post translational modifications
charged molecules under the					
influence of electric current is	Colony hybridization	electrophoresis	dot blot technique	western blotting	electrophoresis
electrophoresis, most commonly used probe is	antibody	lectin	antigens	interferons	antibody
In isoelectric focussing proteins are separated on the basis of their	relative content of positively charged residue only	relative content of negatively charged residue only	size	relative content of positively and negatively charged residues	relative content of positively and negatively charged residues
method for separation of proteins. In isoelectric focussing proteins are separated in the	A pH gradient	A salt gradient	A temperature gradient	A density gradient	A pH gradient
mass of the subunits and the number of each subunit present, the follwoing can be utilized.	SDS-PAGE	Isoelectri focussing	Gel filtration Chromatography	A combination of SDS-PAGE and isoelectric focussing	A combination of SDS- PAGE and isoelectric focussing
features is utilized in the first dimension of 2D gel	Molecular weight	Isoelectric point	Protein conformation	The number of sulphur bridges	Isoelectric point
will migrate to	greater than Rf	lesser than Rf	equal to RF	none of the above	greater than Rf
Impurities present in paper are removed by washin with	1 N HCl	0.1 N HCl	0.01 N HCl	0.001N HCl	0.001N HCl
column chromatography is known as	filter paper	gradient	gel matrix	none of the above	gel matrix
Protein-coding genes can be identified by	Transposon tagging	ORF scanning	Zoo-blotting	Nuclease S1 mapping	ORF scanning
determined by	Gene inactivation	Homology search	Exon trapping	Zoo-blotting	Homology search
analyzed by	Microarrays	Northern analysis	Southern analysis	Protein profiling	Northern analysis

Polypeptides	Can fold into a double helix	Can have a tertiary structure	Can contain phosphate	Are synthesized in the nucleus	Can have a tertiary structure
Reporter genes	Indicate the presence of stress conditions	Are used to characterize proteomes	Are all of bacterial origin	Are used to delineate regulatory sequence elements	Are used to delineate regulatory sequence elements
Microarrays	Are used for analysis of transcriptomes	Are made of glass	Contain RNA sequences	Are smaller than DNA chips	Are used for analysis of transcriptomes
ORF scanning		Is used to find intergenic sequences	Is used to find gene homologies	Is used to find protein- coding genes	Is used to find protein- coding genes
Mass spectrometry is used in be identified by	Transcriptome analysis Phage display	Proteome analysis Microarrays	Protein seperation Hierarchical clustering	Protein identification	Proteome analysis Phage display
which anno actu can torm	Glycine	Proline	Glutamate	Cysteine	Cysteine
wmbh of nyardgêns a-a m me following molecule gives a triplet signal in a normal <sup>1</sup> U	hydrogen a	hydrogen b	hydrogen c	hydrogen d	hydrogen c
Which of the following statements regarding NMR spectroscopy is wrong?	the left of the spectral chart correspond to	Chemical shifts are larger when the frequencies of the radiation which induces the nuclear transitions are higher.	Chemical shifts are larger when shielding effects are greater.	A hydrogen signal splits into $n + 1$ peaks by spin-spin coupling when the number of equivalent hydrogen atoms on adjacent atom(s) is $n$ , and no other neighbouring	Chemical shifts are larger when shielding effects are greater.
Which of the following statements regarding mass spectrometry is wrong?	In a normal mass spectrometer, electron impact causes a molecule to lose an electron and become a molecular radical cation which decomposes into fragment cations and radicals.	Only cations can be detected by a normal mass spectrometer.	A compound whose molecules contain just one bromine atom shows two molecular ion peaks of similar intensity, one at +1 and one at -1 of the average $m/z$ value.	Molecular ion peaks always have even- numbered values of <i>m</i> / <i>z</i> .	Molecular ion peaks always have even- numbered values of <i>m/z</i> .

Which of the following statements is wrong?	A conventional mass spectrometer employs high energy UV radiation.	A conventional mass spectrometer does not employ a spectrophotometric detector.	Conventional mass spectrometry does not always require samples of high purity.	not show signals due	A conventional mass spectrometer employs high energy UV radiation.
Which of the following statements is wrong.	The wavenumber of a band in an IR spectrum is proportional to the frequency of the associated molecular vibration.	Water is a good solvent for recording UV spectra of water- soluble compounds.	Water is a good solvent for recording	leads to broadening of	Water is a good solvent for recording IR spectra of water-soluble compounds.
Northern blots probes are	DNA and RNA	DNA	RNA and protein	RNA	RNA