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# **BIF501 - Bioinformatics II**

# **Topic - 1** Applications of Bioinformatics

# **Applications of Bioinformatics**

- 1. Drug Development
- 2. Crop Improvement
- 3. Microbial Genome
- 4. Gene Therapy
- 5. Biotechnology
- 6. Comparative Study
- 7. Evolutionary Studies
- 8. Veterinary Science
- 9. Molecular Medicine

- Personalized Medicine
   Preventive Medicine
- 12. Waste Cleanup
- 13. Antibiotic Resistance
- 14. Alternate Energy Science
- 15. Insect Resistance
- 16. Climate Change Studies
- 17. Nutritional Quality

# **Drug development**

- ✓ Drugs target only about 500 proteins
- Disease mechanisms and using computational tools identify and validate new drug targets

# **Crop improvement**

- ✓ Comparative genetics of the plant genomes
- ✓ Information obtained from the model crop systems can be used to suggest improvements to other food crops.
- ✓ At present the complete genomes of *Arabidopsis thaliana* (water cress) and *Oryza sativa* (rice) are available.

# **Microbial genome applications**

- ✓ Complete genome sequences
- ✓ Environment, health, energy and industrial applications
- Isolation of the genes that give them their unique abilities to survive under

extreme conditions.

# **Gene Therapy**

- > Gene therapy-used to treat, cure or even prevent disease
- Clinical trials

# Biotechnology

- > Archaeoglobus fulgidus and Thermotoga maritima
- > Corynebacterium glutamicum
- Xanthomonas campestris
- Lactococcus lactis

# **Evolutionary studies**

The sequencing of genomes from all three domains of life; eukaryota, bacteria and archaea

# **Topic - 2** Applications of Bioinformatics

#### **Veterinary Science**

Farm animals including cows and sheep have been sequenced

### **Molecular Medicine**

- > The human genome project
- 3000-4000 hereditary disease including Cystic Fibrosis and Huntingtons disease
- > Response to an environmental stress causes cancers, heart disease, diabetes
- Human Genome Project Data Base

#### **Personalized medicine**

- > Pharmacogenomics
- Sequence variants in DNA
- > Trial and error to find the best drug
- Patient's genetic profile
- With the specific details of the genetic mechanisms of diseases being unraveled, the development of diagnostic tests to measure a persons susceptibility to different diseases may become a distinct reality.
- Preventative actions such as change of lifestyle or having treatment at the earliest possible stages when they are more likely to be successful, could result in huge advances in our struggle to conquer disease.

#### Waste cleanup

- Deinococcus radiodurans
- Potential usefulness in cleaning up waste sites that contain radiation and toxic chemicals

# Antibiotic Resistance

- Enterococcus faecalis
- Virulence region-resistant genes
- The discovery of the region, known as a pathogenicity island

# Alternative energy sources

- Chlorobium tepidum
- > Capacity for generating energy from light

#### **Insect resistance**

- Bacillus thuringiensis
- > Control serious pests of cotton, maize and potatoes
- Insecticides can be reduced and hence the nutritional quality of the crops is increased

#### **Climate change Studies**

- > Increasing levels of carbon dioxide emission-global climate change.
- Study the genomes of microbes that use carbon dioxide as their sole carbon source.

# Improve nutritional quality

- Genes transferred into rice to increase levels of Vitamin A, iron and other micronutrients
- Reducing occurrences of blindness and anaemia

# **Topic -3 Nucleotide Sequence Databases**

# **Biological Databases:**

Biological databases in general store biological data and their main goals are

- 1. Data storage
- 2. Information retrieval
- 3. Knowledge discovery

# **Classification:**

Biological databases can be classified as

- > **Primary databases** (that stores the Primary Sequences)
- Secondary databases (the primary sequences are annotated and kept in Secondary Databases)
- Specialized Databases (they are dedicated towards some specific organism or can have some disease data)

Biological databases can also be classified on the bases of types of data which they contain, such as:

- Nucleotide databases
- Protein databases
- > RNA databases
- Genome databases
- Expression databases (Gene Expression Databases)

# **Issues:**

The issues which are present generally in other databases are also found to be in Biological databases that may be co-related with the relatively slow pace of quality assurance techniques as compared to the pace with which new data is emerging, so the issues are similar and are as follows:

# Due to limited Q/A

- Redundancy

Inconsistency

Incompatibility (format, terminology, data types, etc.)

# Nucleotide Sequence databases:

The Nucleotide Sequence Databases are one of the types of Biological Databases that contains nucleotide sequences in it, which can be DNA and cDNA or EST sequences.



Here, we have a diagram where we have a genomic DNA which has different *Exons*(we know that in Eukaryotes, we have exons and introns). So *exons* gets transcribed into mRNA and we can get cDNA from this mRNA through *reverse transcription* and then we can store this cDNA into our databases whereas the ESTs are the subsets within those cDNA's.

#### **Origin:**

The Nucleotide Sequence Databses were first assembled into Genebank (1982) at Los Alamos National Laboratory (LANL), New Maxico under the leadership of Walter Goad. GeneBankis now working under the umbrella of NCBI (National Center for Biotechnology Information).

NCBI is the central repository that stores multiple types of biological data that includes genomes, their assemblies, their sequencing data, their expression data and what not. In this diagram, we can see the page where you can search for any kind of data; a drop-down list which provides you with various options. The link to this page is http://www.ncbi.nlm.nih.gov/.

Here, is the page for GeneBank, so if you want search about nucleotides and genome sequences, this is the best resource

**NCBI** was established in United States.

#### National Center for Biotechnology Information

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Variation	II 1 2	3 4 5 6 7 8	E-Utilities users: Keep up to date with

#### GeneBank



#### Access to GenBank

There are several ways to search and retrieve data from GenBank

- Search GenBank for sequence identifiers and annotations with Entrez Nucleotide, which is divided into three divisions:
- CoreNucleotide (the main collection), <u>dbEST</u> (Expressed Sequence Tags), and <u>dbGSS</u> (Genome Survey Sequences). Search and align GenBank sequences to a query sequence using <u>BLAST</u> (Basic Local Alignment Search Tool). BLAST searches CoreNucleotide, dbEST, and dbGSS independently; see BLAST info for more information about the numerous BLAST databases

#### **EMBL and DDBJ:**

European Molecular biology Lab (EMBL established1980) was established in Europe.

• DNA databank of Japan (DDBJ) established in Mishima japan (1984).

# **INSDC:**

Genebank, DDBJ and EMBL joined together in International Nucleotide Sequence
 Database Collaboration (INSDC)



You can see in this diagram, the NCBI (National Center for Biotechnology Information), DDBJ(DNA Databank of Japan) and EBI (EuropeanBioinformaticsInstitute) /ENA (European Nucleotide Archive)forms an International collaboration known as INSDC (International Nucleotide Sequence Database Collaboration).

Where **EMBL**established**EBI**, to deal with Bioinformatics kind of stuff and within them they have established **ENA** to maintain the DNA sequence datasets

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Here, is the page of INSDC (International Nucleotide Sequence Database Collaboration), and you can observe that all three collaborators' logos are there. Similarly, if you look into the data, we can have Next Generation reads, Capillary reads and information about samples and annotated sequences all on this first page.

(We'll discuss it later).

# Growth of Genebank:

If we look into the growth of Gene Bank as shown in the figure below (left), we can see the number of **bases** in the GeneBank which are uncountable as they are in trillions which is a huge number starting somewhere in 1982 and if we look into these curves, blue is the growth of GeneBank and red one is the whole genome sequences (which we are comparing) which is starting somewhere in 2003 or 2004 after the publication of Human genome Project.

So, if you look into the number of bases, it seems like they double after every 18 month which means the growth is huge and is exponential.

Similarly, if we look into the sequences (right figure), are also around somewhere in 1000's in 1982 but now they are more than hundred million sequences in this GeneBank.



# GeneBank

# WGS

# http://www.ncbi.nlm.nih.gov/genbank/statistics

# **Conclusions:**

In the end, we conclude some of the followings:

- ➢ Biological databases store biological data.
- > **INSDC** is joint venture of NCBI, EMBL and DDBJ.
- > Growth of bases in **GeneBank** is exponential, doubling every 18 months.

# **Topic - 4 Protein Databases**

# Introduction:

Protein databases store protein data which may include the following:

- ✓ Protein sequences
- Motif (patterns of amino acids)
- ✓ Structure
- Structure alignments (aligned structures)

# **Origin:**

First sequences to be collected were Proteins (before Nucleotide Sequences) using **Sanger and Tupy's**methods (1951) where Common protein families like cytochromes were sequenced (as in that era people were focusing on the sequences made from cytochrome molecules).

Atlas of protein sequences (mainly cytochromes) was assembled by Margret Dayhoff and her collaborators at National Biomedical Research Foundation (NBRF) in 1960s.

# PIR (Protein Information Resource):

The collection (of *Dayhoff* and co) became **PIR** (Protein Information Resource) which is now a collaboration of **NBRF**, Munich Center for Protein Sequences (**MIPS**) and Japan International Protein Information Database (**JIPID**).

# **Protein Sequences:**

**Swiss-Prot**is a Collaboration between the SIB (**S**wissInstitue of **B**ioinformatics) and EBI (**E**uropean**B**ioinformaticsInstitute) and it weekly releases from about 50 servers across the

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world, the main source beingExPASy in Geneva (i.e. it'smainlycontrolled by ExPASywhichis the main server located in Geneva).

Here, is the page for ExPASy, and you can find different structural alignments, proteomic data, genomic data.

#### **Protein Sequences:**

International partnership between PIR, EBI and SIB created **UniProt**, by unifying the PIR-PSD (**P**rotein **I**nformation **R**esource – **P**rotein **S**tructure **D**atabase i.e. they kept the PSD of PIR into UniProt known as PIR-PSD), Swiss-Prot, and TrEMBL(is where we put the translated sequences from the DNA-where DNA is translated into the protein and the protein sequences are coming from different *reading frames* of those DNAs using all 6 reading frames- will be discussed later) databases.



Here, is the page of UniProt and you can see, we have 3 main sections i.e. Protein Ontologies labelled as PRO then we have ProClass where we can have the sequences and ProLINK tells us about the literature.

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Here, we look into the PRO which is the Protein Ontologies- ontologies is where we can classify those proteins on the basis of their functions and different functions have their hierarchy so ontologies are labelled in form of different hierarchies, so there is a major function and a trend towards moving the specific function.

Here, we can see just a PRO Hierarchy Ontology in this example.

http://pir.georgetown.edu/



In this figure, as we mentioned earlier, *i*ProClass is the integration of different protein resources, so we can have sequences from here, protein expression data, and protein modifications. We can also integrate the genomic data with the proteomic data.

#### http://www.uniprot.org/



In this figure, we have *i*ProLINK which provides literature information and most of the research papers can be found here.

The link for it is: <u>http://www.uniprot.org/</u>.

Here, in this figure we have PDB and PDB stands for Protein Data Bank, basically it's a repository where we have the protein structures.

These structures are obtained by different chemistry and molecular biology techniques like Xray Crystallography in the labs and then those structures are submitted into the PDB where researchers can get those structures and can compare their predicted structures with them, so it's a good resource if you are working on structural protein bioinformatics. The link for this page is http://www.rcsb.org/pdb/home/home.do



We can see here for an example, we have the *class* in which we have all the alpha helices; these helices are formed by special arrangement of amino-acids. Basically when the protein sequences- just a linear sequence of amino-acids when it turns around on it selves, it forms those secondary structures so those structures are then recognized as alpha and beta (we are not going into the details; you can go for molecular biology course or Google about alpha or beta). The main idea to present here is that SCOP actually classify the proteins on the basis of those structures so for an example, alpha (is that class where we have all those proteins that has alpha helices in them), we can also have beta (where we have all those proteins that has beta chains in them), alpha/beta (where we have alpha helices then comes beta then comes alpha so they are present one after the other), alpha + beta ( we can have separate regions where we can have alpha helices stacked together and then we have beta chains stacked together). And the link for it is - http://scop.mrc-lmb.cam.ac.uk/scop/.

#### **Conclusions:**

We conclude that:

- ✓ First sequences to be collected were Protein sequences.
- Protein databases are classified on the basis of sequences, motifs, structures and different structural alignments.
- ✓ Growth of Sequence in Databases is exponential (just like as in Nucleotide Databases the growth of sequence is higher).

# **Topic - 5 Genome & Organism Specific Databases**

# **Origin:**

First attempt to sequence free living organism was launched in late 1990's (Blattner et al. 1997) and Viruses had already been sequenced (Fleischmann et al. 1995).

*Haemophilusinfluenzae* was the first genome that was published and the project was initiated at **The Institute of Genome Research (TIGR)** under the leadership of **Craig Ventor** (the same person who's name we'll see in the human genome project). At that time, a method which was already established known as **shotgun sequencing method** was being tested by this project to verify its reliability and efficiency. And by utilizing this method they sequenced the genome which was about 1.8 million base pairs (bp), it took 9-months and the cost was around 1 million US dollars and this project Paved the way for sequencing of many other organisms.

# Examples

• AceDB (AC. *elegans*DataBase) was the first genome database for genome sequences was developed in 1989 and was established by **Richard durbin**and**Thierry-Miegi**.

(Same Durbin whose book "*Biological Sequence Analysis*" we'll consider in the latter half of the course).



Here is the figure of AceDB webpage, you find the *sea elegans*; a worm and there are other organisms.

Link for this page is: http://www.acedb.org/.

#### **Examples:**

TAIR (The Arabidopsis Information Resource) which is a database for Arabidopsis (http://www.arabidopsis.org/) and SGB (Saccharomyces Genome Database) actually uses the system of AceDB (http://www.yeastgenome.org/).

# Human Genome Project:

Human Genome Project started initially as a Pilot Project which begun by Department OfEnergy (DOE) of USA in 1986. Two organizations, one is National Human Genome Research Initiative (NHGRI), which was federally funded organization through NIH (National Institute of Health) that started in 1988 by Francis Collin which was joined to Commercial organization named *Celera (Celera Genomics)* in 1998, a commercial under the leadership of Craig Venter.



Both of them claimed that they sequenced the full Human Genome and the issue was raised that who completed the project first, and who had a major role in it but later on it was resolved by President Bill Clinton at that time and they published it together back in year 2000. In the end, they concluded that there areTotal 3.4 billion bases which are sequenced at a cost of \$1/base.

# UCSC Genome Bioinformatics



While we have those genomes available, we want to see their graphical views where we can get the reports, get the idea about where different genes are located, so in order to do that we needed to make something which we call it as genome browsers- are the webpages where we can look into the different features within our genomes so UCSC is one of the example (shown on the left) which is University of California Santa Cruz which is the biggest genome browser. The link to this browser is http://genome.ucsc.edu/.

The figure of UCSC Genome Browser, where we can have information, so on the top we see a chromosome and down below we see various lines which are known as different tracks (for snips, genes, EST's etc.) so we can look or zoom into different regions of the genome by using those genome browsers.

The link to this webpage is: http://genome.ucsc.edu/.

# **Conclusion:**

In the end, we conclude the following:

✓ Success of *Haemophilusinfluenzae* paved the way for other genome sequencing projects

- ✓ Human Genome Project was accomplished by NHGRI and Celera (they were working independently from one another).
- ✓ Genome browsers help in exploring different regions of the genome.

# **Topic -6 Gene Expression Databases**

#### Gene Expression Omnibus (GEO):

Genes are expressed into mRNA, and whenever we talk about gene expression, we generally mean the mRNA sequences so we can normally get those mRNA from techniques like microarray and another famous technique nowadays which is being established is known as RNAseq. And microarray data and RNAseq can be classified into Gene Expression Data which is stored in Gene Expression Databases.

Basically, Gene Expression Databases are the public repository for the archiving and distribution of gene expression data submitted by the scientific community.

The gene expression data are MIAME compliant data (where MIAME is Minimum Information About a Microarray Experiment and can be searched on this link: http://www.mged.org/Workgroups/MIAME/miame.html). Whenever you need to submit a microarray experiment, you need to give descriptions like how exactly the samples were prepared, normalization methods which you have used and other counts and normalized files) so while keeping in view these datasets have different records in it.

Gene Expression Omnibus is convenient for deposition of gene expression data, as required by funding agencies and journals and it's also a curated resource for gene expression data where we can do Browsing, querying, analysis and retrieval of the data.

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How to Construct a Query		Programmatic Access				
How to Download Data		FTP Site				

Here, is the webpage of GEO which is Gene Expression Omnibus running under NCBI (you can visit NCBI where you can get to the GEO Database) which are having different datasets, has expression profiles where we can see the change in expression of genes across different treatments and we can also analyze this expression data. There is a tool called as GEO2R, we can use BLAST in it. (We'll discuss later)

http://www.ncbi.nlm.nih.gov/geo/

#### **Gene Architecture:**

GEO has four kinds of records or data files (keeping in view the MIAME rules) and are as follows:

- ✓ Sample(GSM) these files stores the sample information like how the samples are prepared, how the treatments are given, how the experimental design was established.
- Platform (GPL) The idea about platforms, they are stored in GPL files so here we can see whether it's a microarray data or RNAseq data (there are different protocols coming from different agencies so we can have that information).

- ✓ Series (GSE) Sometimes different treatments are recorded as separate files so GSE are the files where we can have the similar treatment files and they are put together in a shape of series (are a set of samples and which are somehow related).
- ✓ Datasets (GDS) Whereas the actual data is stored as GDS files which are the sample data collections and are assembled by GEO.

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Series type         Expression profiling by array         Expression profiling by high throughput sequencing         Expression profiling by SAGE         Expression profiling by MPSS         Expression profiling by MPSS         Expression profiling by SNP array         Genome variation profiling by agenome tiling array         Genome variation profiling by genome tiling array         Genome variation profiling by SNP array         Genome variation profiling by SNP array	Count 36,895 612 3,446 241 20 269 12 557 978 56 728	Sories Platforms Samples	Public 51,806 51,805 13,522 1,259,985	Unreleased 7,896 438 192,394	59,70 13,90 1,455
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Series type         Expression profiling by array         Expression profiling by high throughput sequencing         Expression profiling by SAGE         Expression profiling by MPSS         Expression profiling by MPSS         Expression profiling by SNP array         Genome variation profiling by any         Genome variation profiling by some tiling array         Genome variation profiling by high throughput sequencing         Genome variation profiling by SNP array         Genome variation profiling by SNP array         Genome binding/occupancy profiling by array         Genome binding/occupancy profiling by array         Genome binding/occupancy profiling by genome tiling array         Genome binding/occupancy profiling by the throughput sequencing	Count 36,895 612 3,446 241 20 269 12 557 978 56 726 156 2,042 3,317	Sories Platforms Samples	Public 51,806 13,522 1,259,985	Unreleased 7,896 438 192,394	59,70 13,99 1,45
Series type           Expression profiling by array           Expression profiling by genome tiling array           Expression profiling by high throughput sequencing           Expression profiling by SAGE           Expression profiling by MPSS           Expression profiling by SNP array           Genome variation profiling by array           Genome variation profiling by senome tiling array           Genome variation profiling by SNP array           Genome variation profiling by SNP array           Genome binding/occupancy profiling by senome tiling array           Genome binding/occupancy profiling by SNP array	Count 36,895 612 3,446 241 20 269 12 557 978 56 726 156 2,042 3,317 11	Sories Platforms Samples	Public 51,806 13,522 1,259,985	Unreleased 7,896 438 192,394	10,7 13,9 1,45
Series type         Expression profiling by array         Expression profiling by genome tiling array         Expression profiling by high throughput sequencing         Expression profiling by SAGE         Expression profiling by MPSS         Expression profiling by SNP array         Genome variation profiling by agenome tiling array         Genome variation profiling by solve array         Genome variation profiling by solve array         Genome variation profiling by SNP array         Genome binding/occupancy profiling by shigh throughput sequencing         Genome binding/occupancy profiling by array         Genome binding/occupancy profiling by senome tiling array         Genome binding/occupancy profiling by SNP array         Genome binding/occupancy profiling by SNP array         Genome binding/occupancy profiling by SNP array         Methylation profiling by array	Count 36,895 612 3,446 241 20 289 12 557 978 56 728 156 2,042 3,317 11 476	Sories Platforms Samples	Public 51,806 13,522 1,259,985	Unreleased 7,896 438 192,394	59,7( 13,9) 1,453
Series type         Expression profiling by array         Expression profiling by high throughput sequencing         Expression profiling by SAGE         Expression profiling by MPSS         Expression profiling by SNP array         Genome variation profiling by any         Genome variation profiling by SNP array         Genome binding/occupancy profiling by high throughput sequencing         Genome binding/occupancy profiling by array         Genome binding/occupancy profiling by senome tiling array         Genome binding/occupancy profiling by SNP array         Methylation profiling by array         Methylation profiling by genome tiling array	Count           36,895           612           3,446           241           20           289           12           557           978           56           726           156           2,042           3,317           11           476           579	Series Platforms Samples	Public 51,806 13,522 1,259,985	Unreleased 7,896 438 192,394	59,7( 13,9) 1,453
Series type           Expression profiling by array           Expression profiling by high throughput sequencing           Expression profiling by SAGE           Expression profiling by MPSS           Expression profiling by RT-PCR           Expression profiling by SNP array           Genome variation profiling by enome tiling array           Genome variation profiling by SNP array           Genome variation profiling by SNP array           Genome variation profiling by SNP array           Genome binding/occupancy profiling by sename tiling array           Genome binding/occupancy profiling by array           Genome binding/occupancy profiling by SNP array           Methylation profiling by high throughput sequencing	Count           36,895           612           3,446           241           20           289           12           557           978           556           726           156           2,042           3,317           11           476           579           563	Series Platforms Samples	Public 51,806 13,522 1,259,985	Unreleased 7,896 438 192,394	59,7/ 13,9/ 1,452

Here, is the Gene Expression Omnibus page and if we look into the different types of datasets it have, we can have *Series* (on the top left side of right figure), different records for the *Platform, Samples.* If you look into the types of series, you can see there are expression profiling by array, expression profiling by high throughput sequencing (in our course we'll be getting some RNAseq data which is under the expression profiling by high throughout sequencing), similarly there are other various techniques for getting the expression which are

listed below in the Series section as can be seen and number of datasets available are also

() () www.ncbi.nim.nih.go	//gds/?term=colon+cancer+RNASeq 🔛 🖛 ce'   ( 🔝 🕶 sada hai di	aastaane haram 9 合自 4 合 Z 三
S NCBI Resources	How To 🐨	w halder My NCBI Sign Out
GEO DataSets	GEO DataSets 1   colon cancer RNASeq Save search Advanced	Search) Help
Show additional filters	Display Settings: 🕑 Summary, Sorted by Default order Send to:	Filters: Manage Filters
Entry type Series (4)	Did you mean: colon cancer ma seg (60 items)  Results: 4	Top Organisms [Tree]     Homo sapiens (4)     Mus musculus (2)
Select Study type Expression profiling by array More Author	<ul> <li>Turner cell-sevelific inhibition of MYC function using small molecule inhibitors of the HUWE1 1 ubjutilit ligase         (Submitter supplied) Deregulated expression of MYC is a driver of oblevetal cardrogenesis, necessitaling novel strategies to inhibit MYC function. The ubjutilit fligase HUWE1 (HECHN ARTEP) HUBE] second with both MYC and the MYC-associated protein M21. We show here that HUWE1 is required for growth o cleant using and inhibit required for growth o cleant using and the second secon</li></ul>	Find related data
Select Attribute name tissue strain More	cancer cells, but not in stem and normal colon epithelial cells. more Organism: Horos espiens Type: Expression profiling by high throughput sequencing; Genome binding/occupancy profiling by Platform: GPL 10999 By Horoghput sequencing Developed data (BIC (TX, VMO), BRA BR/D044/17 Berlies Accession: GBS/2232 ID : 20009223	Search details ("colonic neoplasms" [MeSH Terms) OR colon cancer[All Fields]) AND RNASeq[All Fields]
Publication dates 30 days 1 year Custom range	PubMed Similar studies     Sequential ChiP-bisuffs sequencing enables direct genome-scale investigation of chromatin an     INA methylation cross-talk     (Submitter supplied) Cross-talk between DNA methylation and histone modifications drives the establishm     (Southitter supplied) Cross-talk traditionally studied using correlative rather than direct	d Recent activity Ium.off Clear Unit Control Clear

present in the column called as *count*.

If you want to look into some dataset, you can simply type into search bar say for example, you write colon cancer RNAseq data which

leads us to the sets of records it gets and when we click onto one of them the page appears (shown below).

S NCBI		GEO
NCBI > GEO > Acce	ssion Display 🕐 Not logged in   L	ogin 😰
Scope: Self	Format: HTML     Amount: Quick     GEO accession: 05557043     Query DataSets for GSE57043	And we get this file, so here is
Title	Public on Apr 25, 2014 Dicer knockout NSCLC RNAseg and miRseg	the information of the
Organism	Mus musculus	
Experiment type	Expression profiling by high throughput sequencing Non-coding RNA profiling by high throughput sequencing	particular dataset
Summary	Dicer knockout NSCLC mRNAseq profiles the transcriptome, Dicer knockout NSCLC miRseq profiles the miRnome	particular dataset.
Overall design	DicerHet and DicerKO NSCLC, 2 biological reps each genotype for mRNAseq, 1 biological rep each for miRseq	Vou can see on the top, that
Contributor(s)	Sharp PA, Chen S	Tou can see on the top, that
Citation(s)	Chen S, Xue Y, Wu X, Le C et al. Global microRNA depletion suppresses tumor angiogenesis. <i>Genes Dev</i> 2014 May 15;28(10):1054-67. PMID: 24788094	· · · · · · · · · · · · · · · · · · ·
Submission date	Apr 24, 2014	each dataset is submitted as a
Last update date	Oct 14, 2014	
Contact name	Sidi Chen	• • • • • • • • • • • • • • • • • • • •
E-mail Rhope	273444158	series with a unique number.
Organization nam	e MIT	
Department	Biology	
Lab Data is su	ubmitted to GEO as a Series, which represents the experiment design	say for example here the
Street address	77 Mass Ave, 76-461	sug for enample here the
City	Cambridge	
State/province	MA 02129	mimber is GSE5 /043 which is
Country	USA	
		basically an id number for this

#### dataset.

So, if we look into this page, we can have the idea about the experiment, the organism from which it's coming, type of experiment, a little bit summary of the experiment, we can also see the contributors name, their publications and addresses.

It is a huge page which is portioned and the other side of it is shown in the figure below

Platforms (1) Samples (6) ≝ Less	GPL13112 Illumina HiSeq GSM1373652 DcrHet_1_r GSM1373653 DcrHet_2_r GSM1373654 DcrKO_1_m GSM1373655 DcrKO_2_m GSM1373656 DcrHet_1_r GSM1373657 DcrKO_1_m	ina HiSeq 2000 (Mus musculus) crHet_1_mRNA crHet_2_mRNA Individual crK0_1_mRNA samples crHet_1_miR in a series crK0_1_miR		5	submissions need to be associated with a platform file.		This is th of the san shown ab Here, we we need t
Relations BioProject SRA Download fan SOFT formattee MINiML formatt Series Matrix Fi	PRJNA245291 SRP041414 hily d family file(s) ted family file(s) ile(s)		F S M T	ormat OFT 2 IINIML 2 XT 2		Click ? One by one	with the platfor the platfor is can be page, and sequence sequence
S GSE57043_dice GSE57043_dice GSE57043_dice SRP/SRP041/SP Raw data provi	iupplementary file erko_fpkm.txt.gz erko_hairpin_rpm.txt.gz erko_mature_rom.txt.gz RP041414 ded as supplementary file	Size 875.5 Kb 4.1 Kb 1.6 Kb	Download (ftp)(http) (ftp)(http) (ftp)(http) (ftp)	File typ TXT TXT TXT SRA Stud	e/resour :e	Normalized counts Raw Reads	Illumina (we'll be upcoming

This is the bottom part of the same webpage shown above.

Here, we can see that we need to be associated with the platforms so the platform information is can be found on this page, and we get those sequences which are sequenced by machine Illumina HiSeq 2000 (we'll be discussed in upcoming lectures). There are total six samples in this dataset, so individual samples are put together labelled as GSM.

We can also download the sequence expression counts or values in different formats and there



An expression signature for p53 status in human breast cancer predicts mutation status, transcriptional effects, and patient survival

- Lance D. Miller\*+, Johanna Smeds<sup>4</sup>, Joshy George\*, Vinsensius B. Vega\*, Liza Vergara\*, Alexander Ploner Yudi Pawitan<sup>5</sup>, Per Hall<sup>5</sup>, Sigrid Klaar<sup>8</sup>, Edison T. Liu\*+, and Jonas Bergh<sup>8</sup>
- \*Genome Institute of Singapore, 60 Biopolis Street, #02-01, Singapore 138672; \*Department of Oncology and Pathology, Radiumhermat, Karolinska institute and Hospital, 5-17176 Stockholm, Sweden: and \*Department of Medical Epidemiology and Biostatistics, Karolinska Instit

GENETICS. For the article "An expression signature for p53 status in human breast cancer predicts mutation status, transcriptional effects, and patient survival," by Lance D. Miller, Johanna Smeds, Joshy George, Vinsensius B. Vega, Liza Vergara, Alexander Ploner, Yudi Pawitan, Per Hall, Sigrid Klaar, Edison T. Liu, and Jonas Bergh, which appeared in issue 38, September 20, 2005, of *Proc. Natl. Acad. Sci. USA* (102, 13550-13555; first published September 2, 2005; 10.1073/pnas.0506230102), the breast cancer microarray data discussed in this publication have been deposited in the National Center for Biotechnology Information's Gene Expression Omnibus database (GEO, www.ncbi.nlm.nih.gov/geo/) and are accessible through GEO Series accession no. GSE3494 [NCBI GEO].

are also some normalized counts as shown in the figure, these are compressed files. There are also raw reads data are present in the format, which we call as SRP or Sequencing Read Archive so that stores the raw read data.

Since, funding and the publication agencies demands that your data should be submitted and shared with the community so here is an example (in the figure shown) where we can see a publication and they have put GSE into their publication which helps other scientists to get access to this data using this ID number as highlighted (in the figure). If you are submitting your paper, you need to provide this information to the publication agencies which is an essential consideration.

# **Conclusion:**

So we sum-up that **GEO** is a public repository for the archiving and distribution of gene expression data and is the Best resource to get microarray and Next Generation Sequencing (RNASeq) data.

# **Topic -7 Medical Databases**

# Introduction

Informatics in health care may be called as health informatics

• It deals with the resources, devices, and methods required to optimize the acquisition, storage, retrieval, and use of information in health and biomedicine.

#### (wiki)

Medical databases store and provide medical information

The premier database for biomedical literature is the National Library of Medicine (NLM)'s MEDLINE, accessible through PubMed

# PUBMED

- Comprises of more than 24 million citations for biomedical literature from MEDLINE, life science journals, and online books
- Citations may include links to full-text content from PubMed Central and publisher web sites



# MEDLINE

- MEDLINE is the primary resource for biomedical journal articles
- Millions of citations to articles in biomedical journals
- MEDLINE uses the MeSH vocabulary

#### **Other Databases**

MEDLINE is the primary resource, but other databases may also be helpful

- Academic OneFile
- CINAHL (Cumulated Index of Nursing and Allied Health Literature)
- PsycINFO
- Web of Knowledge

#### **Academic OneFile**

- Academic OneFile lists articles from journals covering a broad range of subjects
- While it does not primarily focus on medical topics, useful articles can still be found here



# PsycINFO

- PsycINFO searches the psychological literature
- While it does not primarily focus on medical topics, useful articles can still be found here

http://www.apa.org/pubs/databases/psycinfo/coverage.aspx



#### Web of Science

- Major source for articles in a wide range of fields, including the sciences, social sciences, and humanities.
- Excellent place to find articles from scientific journals that may not be included in MEDLINE

#### Conclusions

Informatics in health care may be called as health informatics

Medical databases deal with the acquisition, storage, retrieval, and use of information in health and biomedicine.

# **Topic -8 Sequence Submission Introduction**

# Introduction

Sequences are submitted to the databases in order to share them with the scientific community (sometimes they are also required by the Publication and funding agencies to submit them). Generally sequences are submitted at the time of publication and are reviewed by peers.

#### Caution

It is important to ensure that sequence files do not contain any special characters because sometimes the control characters can also be incorporated into or normal sequences, which can then mess-up the down-stream analysis or data-transfer.

Table 2.1.	Base-nucleic acid code	5
Symbol	Meaning	Explanation
G	G	Guanine
A	A	Adenine
Т	Т	Thymine
C	C	Cytosine
R	A or G	puRine
Y	C or T	pYrimidine
M	A or C	aMino
K	G or T	Keto
S	C or G	Strong interactions 3 h bonds
W	A or T	Weak interactions 2 h bonds
Н	A, C or T not G	H follows G in alphabet
В	C, G or T not A	B follows A in alphabet
v	A, C or G not T (not U)	V follows U in alphabet
D	A, G or T not C	D follows C in alphabet
N	A,C,G or T	Any base

So, there is an issue of how to put the ambiguous nucleotides or amino acids in the sequences (because at some places you are not sure whether it is 'A' or 'T' or 'G' or 'C' and you are restricted to put a single letter). So, there is an organization known as International Union of Biochemistry (IUB), it has established some standard codes to represent those ambiguous bases or amino acids.

For example, here we see that G, A, T or C are just Guanine, Adenine, Thymine and Cytosine respectively. If we see R, it can be either A or G and the word is derived from the group they are coming from i.e. the puRines. We see Y that is the pYrimidine, it can be C or

T.

M stands for if they are having some amine group / amino group in them, K is if they have Keto group i.e. G or T.

S is if they have strong interactions (3 hydrogen bonds) like C and G, who forms triple bonds.

W is for weak interactions, A or T.

Mount, pg 28

Since H follows G in Alphabet so it's everything except G, it can be A, C or T and similar procedure is followed for B,V, and D whereas N can be any base.

1-letter code	3-letter code	Amino acid	
Aª	Ala	alanine	
C	Cys	cysteine	
D	Asp	aspartic acid	
E	Glu	glutamic acid	
F	Phe	phenylalanine	
G	Gly	glycine	
н	His	histidine	
I	Ile	isoleucine	
K	Lys	lysine	
L	Leu	leucine	8
М	Met	methionine	5
N	Asn	asparagine	6
Р	Pro	proline	ţ,
Q	Gln	glutamine	5
R	Arg	arginine	ē
S	Ser	serine	2
Т	Thr	threonine	
v	Val	valine	
W	Trp	tryptophan	
X	Xxx	undetermined amino acid	
Y	Tyr	tyrosine	
Zb	Glx	either glutamic acid or glutamine	

<sup>b</sup> Note that sometimes when computer programs translate DNA sequences, they will put a Z" at the end to indicate the termination codon. This character should be deleted from the

**NCBI:** 

NCBI has two options for sequence submission

**BANKIt** - for simple sequences (not related with down-stream analysis) and annotations and can be submitted through web (if the datasets are small) which does not requires any advanced tools.

Sequin - For Complex sequences and annotations and is also good if we want to do some off-line submissions normally where we have our datasets which are huge ones and can be used in future with some advanced tools (for analysis) and graphical reports.

Similarly, for amino acids, we have single letter codes i.e. from A to Z. And we can see in the figure on the left that some letters are missing.

There are four amino acids that are starting with G, but we gave that G letter to Glycine and for rest of them, we might use some other letters like Glutamic acid is represented as E.

Y stands for Tyrosine (down below) and X can be any amino acid like N (in case of the nucleotide sequences).



In the figures above, are the glances BankIt and Sequin webpages.

# **UniProt:**

# SPIN

Welcome to the new SPIN website! Please use this new website for any new submissions you would like to make. To finish and submit any submissions you may have started using the old website please go here cr. Please note that submissions created using the new website will not be seen in the old one and vice versa.					
is the web-based tool for submitting directly sequenced ledgebase. The information required to create a database	protein sequences and their biological annotations to the UniProt e entry will be collected during the submission process.				
Please sign in	New user?				

For protein sequences, just like NCBI tools, we have UniProt and the similar tool is called as **SPIN** which is a web-based tool for submitting directly sequenced protein sequences and biological annotations to the knowledgebase.

Shown in this figure, is the webpage of SPIN.

We can register here and then we can submit our data.

https://www.ebi.ac.uk/swissprot/Submissions/spin

# **Conclusion**:

We conclude that sequences are stored in databases in specific format and when we want to submit them into a database then we need to follow the guidelines provided by those databases.

# **Topic #9 DNA Sequence Retrieval**

# Introduction

Databases not merely collect and organize data (i.e. not only stores it) but allow intelligent data retrieval (we can do some down-stream analysis on those data sets). Let's see how we can get the DNA data from the NCBI.



So, here is the webpage of NCBI, for example you want to search for say p53 gene; tumour suppressor gene. We write p53 on the search bar, then we get then results, so here we can find many ID entries like 9000 entries are there, we are just looking into the first page in this we choose the first two. So let's click the first one, the p53 where the ID is 2768677, there is a description that what sort of gene is it, and its actually coming from *Drosophila melanogaster*, the location is Chromosome number 3 and we see some Aliases; the alternative names of this gene. The link to NCBI is <u>http://www.ncbi.nlm.nih.gov/</u>.

#### p53 [Drosophila melanogaster (fruit fly)]

Gene ID: 2768677, update	d on 4-Jan-2015	
Summary		2
Official Symbol	p53 provided by FlyBase	
Primary source	FLYBASE:FBgn0039044	
Locus tag	Dmel CG33336	
Gene type	protein coding	
RefSeq status	REVIEWED	
Organism	Drosophila melanogaster (old-lineage: Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota; Neoptera	i.
	Endopterygota; Diptera; Brachycera; Muscomorpha; Ephydroidea; Drosophilidae; Drosophila; Sophophora)	
Lineage	Eukaryota; Metazoa; Ecdysozoa; Arthropoda; Hexapoda; Insecta; Pterygota; Neoptera; Endopterygota; Diptera;	
	Brachycera; Muscomorpha; Ephydroidea; Drosophilidae; Drosophila; Sophophora	
Also known as	CG10873; CG31325; CG33336; D-p53; Dm-P53; Dmel\CG33336; dmp53; Dmp53; DmP53; DMP53; dp53; Dp53;	prac

When we clicked on the first gene as shown in the figure above, we now come to this webpage which is a huge page that is portioned into different figures.

In this figure (on the left), we can see the summary of this gene.

The official symbol is p53 provided by FlyBase which is also written in the *Primary source* (FlyBase is the databases that stores the genome of this fruit-fly *Drosophila*), then the *locus tag*, *gene type* is protein coding, RefSeq says reviewed (sometimes the genes are submitted and reviewed by some other scientist so it means that this gene has been REVIEWED). In the

organism section, we see the classification of that organism and the Aliases are written beneath

Genomic contex	xt				2			
Location: 94D10-	94D10			See p53 in Epigenomics,	MapViewer			
Exon count: 10								
Annotation release	Status	Assembly	Chr	Location				
Release 6.01	current	Release 6 plus ISO1 MT ( <u>GCF_000001215.4</u> )	3R	NT_033777.3 (2304965723054082, complement)				
Release 5.57	previous assembly	Release 5 (GCF_000001215.2)	3R	NT_033777.2 (1887537918879804, complement)				
Chromosome 3R - NT_033777.3								
	C6171	21 Gr94a Ublep C617119 C63	1					

#### it.

In this figure, we can look into the structure of this gene and its coordinates (genomic coordinates), where we can see the location from where it is coming from, we can also see the orientations- the directions in which it is going (down below).

<ul> <li>Genomic regions, transcripts, and products</li> </ul>	
	Go to reference sequence details
Genomic Sequence: NT_033777.3 Chromosome 3R Reference Release 6 plus ISO1 M	T Primary Assembly 💲
	Go to nucleotide: Graphics FASTA GenBank
😉 NT_033777.3: 23M23M (5.8Kbp) C -   🔍   🛟 🖒   - 🥌 🔟 + 🐺	🄀 Tools - 🔮 🕸 Configure ಿ 🤋 -
K. 23,854,500 23,854 K. 23,853,500 23,853 K. 23,852,500 23,852 K. 23,851,50	0 23,051 K 23,050,500 23,050 K 23,049,500
NCBI Genes	×
6r94a	
P53	
NH_206544.2	
NH_2065452	>         >         NP_9962681           >         >         >         NP_901247252.2

In this figure, we can see the genomic region, the transcripts and products tabs, we can look into the products of this gene (the gene when is expressed, the DNA is converted into the RNA). Since it's a eukaryotic genome where there is alternative splicing, so we can find different alternative splice

variants of this gene.

On the upper right side of the figure, it is written as *Go to nucleotide, Graphics, FASTA and GeneBank*, so these are the different views with which we can get access to data files associated with this gene. When we click GeneBank, we are guided to another page, shown in the next figure.

#### Drosophila melanogaster chromosome 3R

NCBI Referen	ce Sequence: NT_033777.3
FASTA Grap	hics
LOCUS	NT_033777 4426 bp DNA linear INV 05-AUG-2014
DEFINITION	Drosophila melanogaster chromosome 3R.
ACCESSION	<u>NT_033777</u> REGION: complement(2304965723054082)
VERSION	NT_033777.3 GI:671162122
DBLINK	BioProject: PRJNA164
	BioSample: SAMN02803731
KEYWORDS	RefSeq.
SOURCE	Drosophila melanogaster (fruit fly)
ORGANISM	Drosophila melanogaster
	Eukaryota; Metazoa; Ecdysozoa; Arthropoda; Hexapoda; Insecta;
	Pterygota; Neoptera; Endopterygota; Diptera; Brachycera;
	Muscomorpha; Ephydroidea; Drosophilidae; Drosophila; Sophophora.
REFERENCE	1 (bases 1 to 4426)
AUTHORS	Hoskins,R.A., Carlson,J.W., Kennedy,C., Acevedo,D., Evans-Holm,M.,
	<pre>Frise,E., Wan,K.H., Park,S., Mendez-Lago,M., Rossi,F.,</pre>
	Villasante, A., Dimitri, P., Karpen, G.H. and Celniker, S.E.
TITLE	Sequence finishing and mapping of Drosophila melanogaster
	heterochromatin
JOURNAL	Science 316 (5831), 1625-1628 (2007)
PUBMED	17569867

We can see the entry in GeneBank and how does it look.

Here, again we see the *name* of the gene, *locus* (where it's ID is written), length of the gene (it is 4426 BP), DNA, it is a linear type of DNA then we have the submission date.

Then the *definition* which is describing the organism's name, chromosome from which it is coming, then it has *accession* (the regions of the genome

from which it is coming from), then we have the *version* (which is NT\_033777.3, so there should have been .1 and .2 and since this is the third review, we can see .3 version here), we also see the reference (down below) and the authors from which this gene is coming and then their publications (it was seen to be published in Science).

	FEATURES source	Location/Qualifiers 14426 /organism="Drosophila melanogaster" /mol_type="genomic DNA" /db_xref="taxon: <u>7227</u> "	Who we d	en we scrolled can see in the find	down, igure on ceatures
	gene	<pre>/chromosome="3R" /genotype="y[1]; Gr22b[1] Gr22d[1] cn[1] CG33964[R4.2] bw[1] sp[1]; LysC[1] MstProx[1] GstD5[1] Rh6[1]" 14426 /gene="p53"</pre>	of the leng	that there are in the so the gene at the solution of the gene at the solution of the solution	total is 4426.
		<pre>/locus_tag="Dmel_CG33336" /gene_synonym="CG10873; CG31325; CG33336; D-p53; Dm-P53; Dmel\CG33336; dmp53; Dmp53; DmP53; DMP53; dp53; Dp53; prac" /map="94D10-94D10" /db_xref="FLYBASE:FBgn0039044"</pre>	We belo	can see mRNA w), and since i	(down t's a
	mRNA	<pre>/db_xref="GeneID:2768677" join(1118,178501,884964,10351071,11351161, 29593268,33333579,36424036,40964426) /gene="p53" /locus_tag="Dmel_CG33336" /gene_synonym="CG10873; CG31325; CG33336; D-p53; Dm-P53; Dmel\CG33336; dmp53; Dmp53; DmP53; dp53; dp53; Dp53; prac" /product="p53, transcript variant B" /note="p53-RB; Dmel\p53-RB; CG33336-RB; Dmel\CG33336-RB" /transcript_id="NM_206544.2" /db xref="C1.281262333"</pre>	is co and it ca with	the regions from the regions from the are shown the word <i>join</i> .	mkinA exons m which below
		/db_xref="GI:281362333 /db_xref="FLYBASE:FB±r0084360" /db_xref="FLYBASE:FBgn0039044" /db_xref="GeneID:2768677"	<b>C</b>	Q,	
	CDS	join(75118,178501,884964,10351071,11351161,	T	hen, within this	mRNA
		29593268,33333579,36424036,40964118) /gene="p53"	w	e find the codin	ng
		/locus_tag="Dmel_CG33336"	se	equences (show	n in the
		/gene_synonym="CG10873; CG31325; CG33336; D-p53; Dm-P53;	fi	gure on the left	), where
		Dmel\CG33336; dmp53; Dmp53; DmP53; DMP53; dp53; Dp53; prac"	cc	oding sequence	s are the
		<pre>/note="CG33336 gene product from transcript CG33336-RB;</pre>	pa	arts of the mRN	JA
		CG33336-PB; p53-PB; p53-like regulator of apoptosis and	W	hich are transla	ited into
		cell cycle; Dmp53; protein 53; drosophila p53"	th	e proteins so th	nere are
		/codon_start=1 /product="p53isoform_B"	fu	rther sub-sets v	within
		/protein_id="NP_996267.1"	th	ose mRNA reg	gions.
		/db_xref="GI:45553461" /db_xref="FIVPAGE.EDDD00002752"	D	own helow we	see the
		/db_xref="FLYBASE:FBgn0039044"	tr	anslated version	n where
		/db_xref="GeneID:2768677"	u (	e can see the w	vord
		/translation="MSLHKSASFSLTFNQNTSIVSRSNSRTIFEAFKEFLDFWDIGN	IE VV	ritton og trangl	ation
		VSAESAVRVSSNGAFNLPQSFGNESNEYAHLATPVDPAYGGNNTNNMQFTNNLEIL	A W	d have we see	the
		LNSGNLMOFSOOSVLREMMLODIOIOANTLPKLENHNIGGYCFSMVLDEPPKSLWMY	s al	id here we see	uie
		IPLNKLYIRMNKAFNVDVQFKSKMPIQPLNLRVFLCFSNDVSAPVVRCQNHLSVEPL	T ai	inno acid seque	ences
		ANNAKMRESLLRSENPNSVYCGNAQGKGISERFSVVVPLNMSRSVTRSGLTRQTLAF	rk co	oming from this	s gene.
I	ORIGIN	סמעשעה זוספסר גופע אראר אנועראנגראניגע אראספרא אפעעראיניאס אופעעסע אופעעסע	C		
	61	cctggagcac ggaagattet tgeggacaca aate ttgagtgeac agecatgagt etteacaagt eege	gcaact gtcgtt	gctaaataaa tagcttgact	atttattat tttaaccagt
	121	gagcggagat attttattcg gtcttaccca acaaa cacttcgatt gtttcgcgta gcaatagtcg cacaa	attaat	gttgcgcctt gaagctttca	aggagtteet
	241	ggatttttgg gatatcggca acgaagtttc tgcag cggagctttc aacttgccgc agagttttgg caace	gagtca	gcagttcggg aacgaatatg	tetecageaa
	361	tacgcctgtg gatccagcct acggaggcaa caaca caatctggaa attttggcca acaataattc cgato	gcaat	aacatgatgc aacaaaatta	agttcacgaa
1	481	caaattegte tgecacaagg ggtgagcaaa tteaa	aacac	gcgctccaat	cgataaacat
	601	ctacgattet gtagttttt gttagegat tttta	atatt	tagectectt	ccccaacaag
	721	accycttgat cagatatagc cgactaagat gtata agaaaggtac agtgcggcaa caaattgatg atcga	acagt	agaaaccttg	catgtagcaa

4261 ggcatgttcg atggccgaaa agaaaacatt tttatatttt tgatagtata ctgttgttaa 4321 ctgcagttct atgtgactac gtaacttttg totaccacaa caaacatact ctgtacaaaa 4381 aagccaaaag tgaatttatt aaagagttgt catattttgc aaacat

sequences which are present starting from 1 until the last nucleotide and the sequence ends with a double slash sign (//).

#### **Conclusions:**

So, we conclude that DNA Sequences are stored in DNA sequence databases in specified formats and Genebank format is a standard format.

# **TOPIC no.10 Protein Sequence Retrieval**

### **Protein Sequences:**

Now we talk about the data retrieval and first we'll talk about the protein sequence retrievals and structures. Protein data is of the following types:

- Actual sequences (from the proteomic data or some other experimental techniques) or translated sequences (sometimes, we go to nucleotides databases, we get those nucleotides and then we translate them by using some softwares, so these are kind of predicted protein sequences).
- Structures (we can also make structures from those proteins that maybe predicted or the real structures coming from various X-ray Crystallography Techniques).
- Annotations (sometimes, we are interested in the functions of the proteins so those are stored as annotations).

#### UniProt (It is an international partnership between PIR, EBI and SIB):

Now as far as the resources are concerned, we have multiple resources for protein sequences but **UniProt**claims to be the biggest and integrated resource whereas for the structures **PDB** seems like a good resource.

As shown in his figure, is the webpage for data retrieval from UniProt, so we

IniProt			Unit	rotHB	p53		36 Adva	nced + Q
LAST Align Retrieve/I	D Mappi	ng		1	-			Help Con
esults							Show	help for UniP
er by'	10	olumns	S BLAST	ligin	± Download 🐵 Add to be		◄ 1 to 25 of 18,363 ►	Show 25
Reviewed (1,868)		Entry	Entry name 🖨		Protein names 🖨 🛛 💹	Gene names 🗘	Organism 🗢	Length 🗘
Unreviewed	0	P04637	P53_HUMAN	-	Cellular tumor antigen p53	<b>TP53</b> , P53	Homo sapiens (Human)	393
,495) 18L	0 1	P02340	P53_MOUSE	-	Cellular tumor antigen p53	<b>Tp53</b> , P53, Trp53	Mus musculus (Mouse)	387
oular organisms nan (1,085)	0	P10361	P53_RAT	5	Cellular tumor antigen p53	<b>Tp53</b> , P53	Rattus norvegicus (Rat)	391
(317)	0	Q42578	PER53_ARATH	2	Peroxidase 53	PER53, P53, At5g06720, MPH15.8	Arabidopsis thaliana (Mouse-ear cress)	335
ine (276) rafish (248)	0	P25035	P53_ONCMY	5	Cellular tumor antigen p53	<b>tp53</b> , p53	Oncorhynchus mykiss (Rainbow trout) (Salmo gairdneri)	396
	0 1	P79820	P53_ORYLA		Cellular tumor antigen p53	<b>tp53</b> , p53	Oryzias latipes (Medaka fish) (Japanese ricefish)	352
r organisms								

want to search a protein, say

p53, where we put it into the search box and press enter which gives us the output. And we see that there are 18,000 different records and it is showing us the first 25 out of them.

We can have different columns for the output on this webpage so we can have *entry; it's ID, entry name* (the Suffix Human is written so it's coming from Human, it can be from mouse, rat and Arabidopsis), the *protein name* is Cellular tumour antigen, then *gene name* which is TP53 (where TP stands for Tumour Protein), the *organism* is obviously the human (here) and in the end we have it's length i.e. 393 bp (base pairs).

The link to this webpage is http://www.uniprot.org/uniprot/.

So, let's check the first one and here we reach on the record for this protein

P04637	- P53	HUMAN				🔀 Basket 👻
	Protein	Cellular tumor a	ntigen p53			
	Gene	TP53				
On	ganism	Homo sapiens (Hum	ian)			
	Status	Reviewed - 👀	••• Exper	imental evidence at	protein level <sup>i</sup>	
Display	None	SBLAST EAlign	Format	Here Add to basket	() History	📌 Feedback 🗖 Help video
FUNCTION		Function				
NAMES & TAXONO	MY	Acts as a tumor suppre	essor in many	tumor types; indu	es growth arrest	or apoptosis depending on the physiological circumstances and cell type.
SUBCELLULAR LO	CATION	Involved in cell cycle n	egulation as a	trans-activator that	t acts to negative	ely regulate cell division by controlling a set of genes required for this
PATHOLOGY & BIO	тесн	of BAX and FAS antige	n expression,	or by repression of	Bcl-2 expression	In cooperation with mitochondrial PPIF is involved in activating
	G	oxidative stress-induce RNA p21 (lincRNA-p21	ed necrosis; ti ) and lincRNA	he function is largel MkIn1. LincRNA-p2	y independent of 1 participates in	transcription. Induces the transcription of long intergenic non-coding TP53-dependent transcriptional repression leading to apoptosis and
EXPRESSION		seem to have to effect	on cell-cycle	regulation. Implicat	ed in Notch signa	aling cross-over. Prevents CDK7 kinase activity when associated to CAK
		some but not all TP53-	inducible pro	e, thus stopping cei moters. Isoform 4 s	uppresses transa	ctivation activity and impairs growth suppression mediated by isoform 1.
		Isoform 7 inhibits isofo	orm 1-mediate	ed apoptosis. 🎸 11 P	ublications 👻	

(shown in the figure on the left) which is Cellular Tumour Antigen p53 protein, commonly known as TP53.

We can have different tabs, showing us the outputs. We can look into the *functions*, its *taxonomy*, and lot many other characteristics so if we look into the function so it gives us some description about what it's doing.

SEQUENCES (9)	Feature key	Position(s	) Length	Description	Graphical vi	ew Feature identifier	Actions
CROSS-REFERENCES	Chal	120 12		Teterantian with Chia			
PUBLICATIONS	Sice	120 - 1.		- Interaction with Diva			
ENTRY INFORMATION	Metal binding	176 - 1.	6 1	Zinc			
	Metal binding	179 - 11	19 1	Zinc			
	Metal binding	238 - 23	18 1	Zinc			
SIMILAR PHOTEINS	Metal binding	242 - 24	12 1	Zinc			
Тор							
Regions							
Feature key	Position(s)	Length	Description		Graphical view	Feature identifier	Action
DNA binding <sup>1</sup>	102 - 292	191					
GO - Molecular fur ATP binding & Sour chromatin binding damaged DNA bin	Ce: UniProtKB - Source: UniProtK Ging & Source: Ref	B 🔶 Genome		<ul> <li>Chaperone bindin</li> <li>Copper ion bindin</li> <li>DNA binding </li> </ul>	g & Source: UniProtKB ~ g & Source: UniProtKB ~ urce: UniProtKB ~		
GO - Molecular fur ATP binding & Sour chromatin binding damaged DNA bin enzyme binding &	Ce: UniProtKB Source: UniProtKB Source: UniProtKB Source: UniProtKB	B		<ul> <li>&gt;chaperone bindin</li> <li>&gt;copper ion bindin</li> <li>&gt;DNA binding </li> <li>✓ So</li> <li>&gt;histone acetyltrat</li> </ul>	g & Source: UniProtKB ~ g & Source: UniProtKB ~ nurce: UniProtKB ~ nsferase binding & Sou	rce: UniProtKB 👻	
GO - Molecular fur ATP binding & Sour I chromatin binding I damaged DNA bin I enzyme binding & histone deacetylas	Ce: UniProtKB ~ Ø Source: UniProtK ding Ø Source: Ref Source: UniProtKB ie regulator active	B 🛩 Genome V ity & Source: Ens	embi	<ul> <li>chaperone bindin</li> <li>copper ion bindin</li> <li>DNA binding  So</li> <li>histone acetyltrat</li> <li>identical protein binding</li> </ul>	g & Source: UniProtKB ~ g & Source: UniProtKB ~ urce: UniProtKB ~ nsferase binding & Sou binding & Source: IntAct	rce: UniProtKB 👻	
GO - Molecular fur ATP binding & Sour chromatin binding damaged DNA bin enzyme binding & histone deacetylas >p53 binding & Sour	Action Action	B ← Genome ← ilty ∲ Source: Ens	embl	<ul> <li>chaperone bindin</li> <li>copper ion bindin</li> <li>DNA binding </li> <li>So</li> <li>histone acetyltran</li> <li>identical protein i</li> <li>protease binding</li> </ul>	g & Source: UniProtKB ~ g & Source: UniProtKB ~ nurce: UniProtKB ~ nsferase binding & Source: IntAct & Source: UniProtKB ~	rce: UniProtKB 👻	
GO - Molecular fur >ATP binding & Sour >chromatin binding damaged DNA bin >enzyme binding & histone deacetylas p53 binding & Sour >protein heterodimu	rec: UniProtKB ~ # Source: UniProtK Source: UniProtKB ie regulator activity ce: RefGenome erization activity	B - Genome ity & Source: Ens	embl	<ul> <li>chaperone bindin</li> <li>copper ion bindin</li> <li>DNA binding # 55</li> <li>histone acetyltrat</li> <li>identical protein i</li> <li>protease binding</li> <li>protein kinase bin</li> </ul>	g & Source: UniProtKB - g & Source: UniProtKB - urce: UniProtKB - source: IniProtKB - Source: UniProtKB - nding & Source: UniProtK	rce: UniProtKB -	

After scrolling the same webpage (shown in the figure on the left), we can see the *feature key* and in some *site* written (there are unique sites in different proteins which are having some specific properties in them so this is just one amino-acid present in this protein that *interacts with the DNA*). Similarly, there are different *metal binding sites* and we can see that it's mainly binding to the *Zinc*metal.The number of amino-acids is shown here so these are the regions where it interacts with the metal.

Down below, we can also see the *DNA binding* region, for example here, the amino acids are from 102 to 292 and that is also shown in the *Graphical view* as well.

GO-Molecular function or GO-Gene Ontologies, so gene ontologies are the different functional annotation term, there they define different functions, so amongst them we have molecular functions, biological processes, and we have cellular components. So here we just see a *Molecular function*, so it tells us that it performs the functions as shown in the figure , mainly it's a *ATP binding, it's p53 binding* with various other functions like *DNA binding*. So all those functions related to these proteins are present in the heading of GO-Molecular Function.

Keywords - Molec Activator	ular function <sup>1</sup>
Keywords - Biolog Apoptosis, Cell cycle	<b>jical process</b> <sup>1</sup> a, Host-virus interaction, Necrosis, Transcription, Transcription regulation
Keywords - Ligand DNA-binding, Metal-	d <sup>i</sup> -binding, Zinc
Enzyme and pathy	vay databases
Reactome <sup>i</sup>	REACT_118568. Pre-NOTCH Transcription and Translation. REACT_1194. Activation of NOXA and translocation to mitochondria. REACT_121. Activation of PUMA and translocation to mitochondria. REACT_169121. Formation of Senescence-Associated Heterochromatin Foci (SAHF). REACT_169185. DNA Damage/Telomere Stress Induced Senescence. REACT_169436. Oxidative Stress Induced Senescence. REACT_169436. Oxidative Stress Induced Senescence. REACT_20549. Autodegradation of the E3 ubiquitin ligase COP1. REACT_24970. Factors involved in megakaryocyte development and platelet production. REACT_309. Stabilization of p53.
SignaLink <sup>i</sup>	P04637.

Next, we move on to some other functions, in the Biological process category (shown in the figure) we see that it is related to Apoptosis (which is a cell death and it is related to cell-cycle and some other components).

In the below section, we see some *enzymes and pathway databases*,

and*Reactome* is a database in which we have a group of reactions which are categorized so these are the list of those reactions with which it is related.

Protein family/grou	up databases
TCDB	1.C.110.1.1. the pore-forming pnc-27 peptide of 32 aas from the p53 tumor suppressor protein (pnc-27) family.
Names & Taxo	onomy'
Protein names <sup>i</sup>	Recommended name: Cellular tumor antigen p53 Alternative name(s): • Antigen NY-CO-13 • Phosphoprotein p53 • Tumor suppressor p53
Gene names i	Name:TP53 Synonyma:P53
Organism <sup>1</sup>	Homo sapiens (Human)
Taxonomic identifier <sup>1</sup>	9666 [NCBI]
Taxonomic lineage <sup>i</sup>	Eukaryota > Metazoa > Chordata > Craniata > Vertebrata > Euteleostomi > Mammalia > Eutheria > Euarchontogilres > Primates - Haplornhini > Catarrhini + Hominidae > Homo D2
Proteomes <sup>1</sup>	UP000005640: Chromosome 17

#### When we move

further (as shown in the figure on the left) till we reach its Taxonomy.

On the top, we can see something written as *Protein family* or group databases which is TCDB. Basically, there is another classification in which the proteins are classified on the basis of being as transporter proteins so it is associated with the transportation across the membranes and there is 5-digit number, so there is a specific classification code which is given to each protein, and this protein has the specific code as shown in the figure.

So then we have the names and taxonomies, where there are protein names, and thetaxonomyof the individual can be seen in the Taxonomic lineage row. Let's see how we reach to its sequence and is shown in the figure below:

is also the s	equence that a	as the 'canonic appears in the	downloadable	versions of the entry.	y refers to it. IIguio
de					we ca
					see th
10	20	30	40	50	
AEEPQSDPSV	EPPLSQETFS	DLWKLLPENN	VLSPLPSQAM	DDLMLSPDDI	seque
60	70	80	90	100	of the
QWFTEDPGP	DEAPRMPEAA	PPVAPAPAAP	TPAAPAPAPS	WPLSSSVPSQ	of the
110	120	130	140	150	prote
KTYQGSYGFR	LGFLHSGTAK	SVTCTYSPAL	NKMFCQLAKT	CPVQLWVDST	Prote
160	170	180	190	200	which
PPPGTRVRAM	AIYKQSQHMT	EVVRRCPHHE	RCSDSDGLAP	PQHLIRVEGN	found
210	220	230	240	250	Tound
JRVEYLDDRN 260	TERHSVVVPY	EPPEVGSDCT	TIHYNYMCNS	SCMGGMNRRP	be at
	270	FUDUCACDCD	DEDTEFENTE	SUU RECEDULET D	oe ut
310	320	BVRVCACPGR 330	JAN 340	350	end o
PGSTKRALPN	NTSSSPOPKK	KPLDGEYFTL	OIRGREEFEM	FRELNEALEL	than
360	370	380	390		the pa
			WHI WOWDOOD	DCD	

Isoform 1, so different proteins have different isoforms, different alternative splice variants so this is Isoform 1 as exhibited by its name which is P04637-1, and is the kind of first isoform. We can see the sequence of the protein and starts with a methionine (always a first amino acid in those proteins) and ending at  $390^{\text{TH}}$  amino acid. So, it's a 393 aa long protein and the sequence is right here. You can click on the FASTA button on the top and then you can get this output in FASTA format (we'll discuss it later).

# **NCBI:**

it

We can also get the same protein from NCBI (as shown in the figure on the left)

ORIGIN 1 61 121 181 241 301	meepqsdpsv deaprmpeaa svtctyspal rcsdsdglap scmggmnrrp pgstkralpn	epplsqetfs ppvapapaap nkmfcqlakt pqhlirvegn iltiitleds ntssspqpkk	dlwkllpenn tpaapapaps cpvqlwvdst lrveylddrn sgnllgrnsf kpldgeyftl	vlsplpsqam wplsssvpsq pppgtrvram tfrhsvvvpy evrvcacpgr qirgrerfem	ddlmlspddi ktyqgsygfr aiykqsqhmt eppevgsdct drrteeenlr frelnealel	eqwftedpgp lgflhsgtak evvrrcphhe tihynymcns kkgephhelp kdaqagkepg	In NCBI, obviously the sequence is
301 361 //	pgstkralpn gsrahsshlk	ntssspqpkk skkgqstsrh	kpldgeyftl kklmfktegp	qirgrerfem dsd	frelnealel	kdaqagkepg	pretty similar and
							the

#### arrangement is slightly

#### different so it is

*ORIGIN*, where the sequence starts and sequence ends at those two slashes (//). So, we can get the protein sequence from NCBI as well and the link to this website is http://www.ncbi.nlm.nih.gov/.

### PDB:

ASTA   Sequence & DSSP   Image	Display Parameters
Polymer 2 Length: 93 residues Chain Type: polypeptide(L) Reference: UniProtKB P04637 (P	Currently displayed: SEQRES sequence. Display external (UniProtKB) sequence Mouse over an annotation to see more details. Click annotation to enable Jmol.
Annotations	
Secondary Structure: DSSP 4% helical (1 helices; 4 residues) [hide] [reference]	
DSSP PDB MEE P QS DP S V E P P LS QET F S DLWKLLPENN VLS P LP S QAMDDLM PDB DSSP PDB DEAP RMP EAAP P VAPAPAAP T PAAPAPAP SWP L PDB	ALSPDDIEQWFTEDPGP 50 60
DSSP Legend T: turn	
empty: no secondary structure assigned	
S: bend	
A A H aloba belix	

**PDB** gives us the structures, so we can go to PDB webpage (as shown in the figure on the left) and search for the same ID i.e. P04637 and it gives us the sections or the regions from where it can make up some specific structures.

You can see the *turns* in Annotations section, the black ones are the empty lines where no secondary structure can be formed, blue ones show those bends and the orange ones are designated as alpha helices regions. So in PDB, we can have structures in this format as well as the 3D-Structures as shown in the figure below:



# The link towards PDB is: http://www.rcsb.org/pdb/explore/remediatedSequence.do?structureId=2LY4 &bionumb er=1

#### **Conclusions:**

We conclude that:

- UniProt is the integrated resource between PIR, EBI and SIB and
- PDB is a good resource to get the protein structure.

# **TOPIC # 11 Sequence Formats**

Sequences are stored in different formats in databases and since software requires those sequences in specific format so it's good to have an idea about what major formats are, we'll look into few of them.

# **FASTA Sequence Format**

FASTA is the most recognized and well distributed format to present DNA and Protein sequences.

The sequence starts with a 'greater than' sign (>), whereas the actual sequence is always on the next line. It is recommended that all lines of text should be shorter than 80 characters in length (generally we have 60 characters).

# <u>Example:</u>

>gi|568815581:c7687550-7668402 Homo sapiens chromosome 17, GRCh38 Primary Assembly

GATGGGATTGGGGTTTTCCCCTCCCATGTGCTCATCTAGAGCCACCGTCCAGGGAG CAGGTAGCTGCTGGGCTCTCCACGACGGTGACACGC------

>gi|120407068|ref|NP\_000537.3| cellular tumor antigen p53 isoform a [Homo sapiens] MEEPQSDPSVEPPLSQETFSDLWKLLPENNVLSPLAPPVLGFLHSGTAKSVTCTYSPAL NKMFCQLAKT--\*

This sequence is of DNA in the *fasta* format (shown above), which starts with the '*greater than*' sign (>), same as in the case of the protein sequence in the *fasta* format (shown below) that also starts with the same symbol.

Then we have 'gi' written which stands for 'gene identification' and the numbers shown are the 'ID' in both of the sequences.

In the DNA sequence, we have 'c' followed by the 'ID', this 'c' basically means the sequence is of the complementary strand and the regions from where it is coming are designated here; the base positions in between them, this gene is located. Then we have a short description of this gene that it belongs to '*Homo sapiens*', 'chromosome 17' and the 'Primary Assembly' (assembly is where we get short sequence reads or small sequences and we put them together into a gene, known as assembly). Then finally, we have the actual sequence which is around 60 characters long in each line (as the sequence was quite long, we have used dashes to represent further characters).

In the protein sequence, we have 'ref' followed by the 'ID', which gives us an idea that it is a reference sequence (reference sequences are the curated sequence, there is a sub-section in NCBI called as *ref seq*, so they have reference sequences ; a kind of standard sequences to avoid any kind of redundancy. So, we can say these are the primary or the main sequences and we might have other alternative splice variants but references are the kind of true representative of the class). Followed by ref, we have another ID, which is the 'protein ID'. Then we have its brief description that it's a 'cellular tumor antigen' protein 'p53 isoform' and is also from '*Homo sapiens*'. Finally, we have the actual sequence of this protein and in the end we have dashes that represents it is an incomplete sequence and steric (\*) is shown (sometimes the steric (\*) is found to be seen in *fasta*files but sometime it don't, so the software must know what does this specific steric (\*) stands for).

# **GeneBank Sequence Format:**

GeneBank sequence format is found to be in the GeneBank Database which is a kind of standard format and other formats are pretty similar to it.

A sequence file in Gene Bank format can contain several sequences. One sequence starts with a line containing the word LOCUS and a number of annotation lines. The start of the sequence is marked by a line containing "ORIGIN" and the end of the sequence is marked by two slashes ("//").

# Example:

Here is the GeneBank format, which starts with the word '*LOCUS*', then we have it's 'ID', it is 237 base pairs long, we have some short description that it is a '*DNA*', '*PRI*' – primary sequence, submitted on '04-FEB-1995'.

Then we have a 'DEFINITION' line where we can have some description/explanation about this gene. Then again we have an 'ACESSION number'. It also provides us with the 'BASE COUNT' (i.e. how many A's (Adenines), G's (Guanines), C's (Cytocines), T's (Thymines) are there).

Then finally the word 'ORIGIN' tells us that the actual sequence is right here, we have these lines (60 bases on each line) that are separated into chunks of 10 bases and is a kind of standard practice. The sequences ends with the those slashes (//).

#### **EMBL Format:**

This format is similar to that of GeneBank Format. An example sequence in EMBL format is:

ID AA03518 standard; DNA; FUN; 237 BP. XX AC U03518; XX DE Aspergillusawamori internal transcribed spacer 1 (ITS1) and 18S DE rBNA and 5.8S rBNA genes, partial sequence

DE rRNA and 5.8S rRNA genes, partial sequence.

XX

SQ Sequence 237 BP; 41 A; 77 C; 67 G; 52 T; 0 other;

aacctgcggaaggatcattaccgagtgcgggtcctttgggcccaacctcccatccgtgtc60tattgtaccctgttgcttcggcgggcccgccgctgtgcggcgcgcctctg120ccccccgggcccgtgcccgcggagaccccaacacgaacactgtctgaaagcgtgcagtc180tgagttgattgaatgcaatcagttaaaactttcaacaatggatctcttggttccggc237

Here, we have ID, accession number (AC), descriptions (DE), and the sequence actually starts from where the word 'SQ' is there, and we can observe that we have pretty similar lines as seen in the previous example. Finally, the sequence ends with doubles slashes same as in GeneBank format.

# **SwissProt Format:**

SwissProt protein sequence format is similar to EMBL format but there is considerably more information about physical and biochemical properties of a protein (as you can see below there is more description).

ID - Identification.

- AC Accession number(s).
- DT Date.
- DE Description.
- GN Gene name(s).
- OS Organism species.
- OG Organelle.
- OC Organism classification.
- RN Reference number.
- RP Reference position.

- RC Reference comments.
- RX Reference cross-references.
- RA Reference authors.
- RL Reference location.
- CC Comments or notes.
- DR Database cross-references.
- KW Keywords.
- FT Feature table data.
- SQ Sequence header.
- // Termination line.

# XML Format:

It is a modern practice in which we try to put those sequences in kind of a machine language. So, XML stands for Extensible Markup Language. The format is similar to HTML (language for Web programming).

The good part is that this language is in between machine and man readable so it's kind of easy to code over this.

And it is becoming standard data format for transferring genome data.

# Example:

<xsd:annotation> <xsd:documentation> XML Schema for SBOL core data model compatible with RDF/XML serialization. <dc:date>2012-01-19</dc:date> <dc:creator>EvrenSirin</dc:creator> <dc:contributor>Michal Galdzicki</dc:contributor> </xsd:documentation> </xsd:annotation>

This format seems pretty weird but not for the people with computer science background.**NBRF Format:** 

>DL;seq1 seq1, 16 bases, 2688 checksum. agctagctagctagct\*

>DL;seq2 seq2, 16 bases, 25C8 checksum. aactaactaactaact\*

The format is pretty similar to fasta but in addition to that it gives us the checksum value (checksum- we take those nucleotides and since we know that in computers every digit is related to some 'ascii' value, we can take those values and add them up together and then we can come up with this number known as checksum. So, it's a good thing to have this number as when somebody is downloading the sequence, he can again check on his computer and find the checksum, if they are equivalent to one another, the sequences are correctly downloaded otherwise there must be some issues with the downloading)

# **GCG FORMAT:**

GCG stands for Genetics Computer Group (basically it was a group of scientists who were helping the biological community to develop different software and training programs to help with the biological sequence analysis problems, so they also came up with the sequence formats). This format is kind of similar to the NBRF format (we have checksum but we don't have greater than (>) sign as in fasta, we have length of the sequence). There can be multiple sequences in one file.

# Example:

seq1 seq1 Length: 16 Check: 9864 .. 1 agctagctagctagct seq2 seq2 Length: 16 Check: 9672 .. 1 aactaactaactaact

#### **Sequence converters:**

Sometimes, we need to convert between sequences so you can come up with your own script or you can come up with your own codes and there are also some programs meant for this purpose alone such as READSEQ is a useful sequence converter (developed by D.G.Gilbert at Indiana University, USA) basically it recognizes DNA or Protein sequence file and interconvert them between different formats.

### **Conclusions:**

What we conclude in the end of this lecture is the following:

- Databases store sequences in specified formats
- Genebank, DDBJ and EMBL has similar formats
- Different software need sequences in different formats

We might convert the sequences into other formats on our own or we can also simply use one of the programs available for converting like READSEQ

# **Topic -12 Data Retrieval**

#### **Data Retrieval:**

Nearly all biological databases are available for download as simple text (flat) files. Sometimes we are interested to download the database and do the analysis locally in our own machines which might save our time as the local version of the database allows one greater freedom in processing the data.

# **ENTREZ:**

It is an integrated search engine that works behind NCBI, so you can do lot of researches and can look for variety of data using it (It can be accessed from the site <u>www.ncbi.nlm.nih.gov/Entrez/</u>). It integrates PubMed and 39 other scientific literatures, nucleotide and protein databases. For example, it can be **protein domain data**, **population** 

studies, expression data, pathways, genome details and taxonomic information.



Here, we can see it integrates between GEO (gene expression sets), OMIM (Online Mendelian Inheritance in Man), Genome Databases, taxonomy Databases, etc. And we can see that in the middle we have Nucleotide, PubMed and Protein. So it is an integrated system which operates between different databases, so you can simply search for whatever you are looking after and ENTREZ will search it for you.

S NCBI Resources 🖂 How T			Sign in	to NCBI
Search NCBI databases			Search	Help
Literature		Genes		
Books MoSH NLM Catalog PubMed PubMed Central Health	books and reports ontology used for PubMed indexing books, journals and more in the NLM Collections scientific & medical abstracts/citations full-text journal articles	EST Gene GEO DataSets GEO Profiles HomoloGene PopSet	expressed sequence tag sequences collected information about gene loci functional genomics studies gene expression and molecular abundance profiles homologous gene sats for selected organisms sequence sets from phylogenetic and population studies.	
ClinVar dbGaP GTR MedGen OMIM PubMed Health Genomes	human variations of clinical significance genotype/phenotype interaction studies genetic testing registry medical genetics literature and links online mendelian inheritance in man clinical effectiveness, disease and drug reports	UniGene Proteins Conserved Domains Protein Protein Clusters Structure Chemicals	clusters of expressed transcripts conserved protein domains protein sequences sequence similarity-based protein clusters experimentally-determined biomolecular structures	
Assembly BioProject BioSample	genomic assembly information biological projects providing data to NCBI descriptions of biological source materials	BioSystems PubChem BioAssay	molecular pathways with links to genes, proteins an chemicals bioactivity screening studies	d

Here, is the page of ENTREZ that allows you to search anything by the help of a search bar at the top. It has different connections like we have Literature resources, we have Health Databases, Genomes, different Genes Databases, Proteins and Chemicals.

# **Bulk Data Retrieval:**

Sometimes, we need to obtain data in bulk amount and for this purpose normally we use Linux but for Windows users, there are some packages or programs available and are known as FTP clients so the best option is to use FTP (File transfer protocol). The File Transfer Protocol (**FTP**) is a standard network protocol used to transfer files Via command line or application programs like FTP clients (we'll be using it).

Once, we get the data which is mostly not in a proper format and every other software require different specific formats so we might want to use some programming languages to help convert the data into the required format. The programming languages like PERL and Python are good for processing Biological data in Bioinformatics.

# **Conclusions:**

We have learned that :

- Data is transferred over the internet.
- Data needs to be transformed or processed before handing it over to any software.

# Topic # 13 Genome Informatics

Now we talk about the main subject which we are seeking in this course that is the *Genome Informatics* and let's look into it.

# **Genome Informatics:**

It is about the Genome sequencing that provides the sequences of all the genes of an organism. The major application of Bioinformatics is the analysis of full genomes that have been sequenced. Whereas the challenge is to identify those particular genes that are predicted to have a specific biological function.

# **Genomics definition:**

NHGRI (National Human Genome Research Institute) defines Genomics as:

# "Study of all of a person's genes (the Genome), including interactions of those genes with each other and with the person's environment."

# So, *Genome Informatics* can be defined as:

It is the field in which computational and statistical techniques are applied to derive biological information from genome sequences.

# "Genome informatics includes methods to analyze DNA sequence information and to predict protein sequence and structure." (Iossifov, et al. 2014)

# **Genome Sequences:**

So, after we have the genome sequences, we do the analysis of those sequences and it includes the following:

- The discovery and utilization of sequence polymorphisms (different sequence vary from one another, so we can identify those polymorphisms).
- Opportunity to explore genetic variability both between and within the organisms (we can help identify different traits of those organisms by using those polymorphisms).

# **Genome Analysis:**

In Genome Analysis we mainly perform the following tasks;

# Sequencing

- Assembly (since the sequencing is done in a way that whole genome is broken down into short fragments and once those fragments are sequenced, we need to put them together, this step in genome analysis is known as Assembly).
- Repeat identification and masking out (once we assemble that genome, we try to find out the regions in which we have large number of repeats because assemblies jumble up where we have those repeats so we need to find those regions and it is one of the important task to go and look into those assemblies while keeping in mind those regions in which we have those repeats).
- Gene prediction (after we have assembled a finished genome, now we can go for the prediction of the genes where we can find the genes by using different patterns or features of those genes).
- Looking for EST (Expressed Sequenced Tags) and cDNA (complementary DNA) sequences.

(EST and cDNA are basically originated from the DNA where the genes that are expressed are transcribed into mRNA which is then reversed transcribed back into cDNA. So by the help of cDNA, we can look into where those expressing regions are present in the genes that will give us the idea of the gene expression or the regions from where the mRNAs are made).

Genome annotation (in which we can find out similar functions performed by different genes)

*Expression analysis* (once we have the idea about the regions of the gene in which we can have the gene expression then we can explore the quantification i.e. how much those genes are being expressed).

- Metabolic pathways and regulation studies (once those genes are expressed, their products interact with each-other and then they perform different metabolic roles in the shape of different metabolic pathways and networks).
- Functional genomics (where we are actually looking into the different functions performed by different regions of the genome that are under the control of different genes and what exactly would be the effect of changes in those genes specifically if we want to study about the genes related to diseases).
- Gene location/gene map identification (map the location of those genes on the chromosomes).
- Comparative genomics (in which we can take one genome and compare it with another genome, where we can find the comparative features; what is present in the first genome and not in the second one and the intersections between them, etc.).

- Identify clusters of functionally related genes (those genes they might be having similar structures, sequences and also performs similar functions, which can give us the idea about the evolution).
- Evolutionary modeling (so the identification of the clusters of functionally related gene can help us in making an evolutionary model).
- Self-comparison of proteome (sometimes we are interested in finding genes which are kind of duplicated within the same organism, so in order to do that this self-comparison of proteome is made, where proteome is the collection of the proteins which are derived from those genomes. Therefore, the whole collection of one organism's proteins can be termed as proteome and we can compare it with itself and can find about those sequences which are being duplicated in it).
- Model organisms:

Most of the times, while we are doing those genome sequencing projects, our objective is to find the cure of some disease, or improving some variety of the crop for enhancing its production, or looking into some drugs against different organisms so it's a good idea to have some model organisms that can be used for studying various processes in labs and there are is a range of model organisms which includes:

- ➢ E. coli − bacteria
- S. cerevisiae yeast
- C. elegans worm
- > D.melanogaster fly
- > Daniorerio zebrafish
- Musmusculus mouse
- Homo sapiens you and me
- Arabidopsis plant



Here, in this diagram we see auniversal tree of life that has been made with the structures of small ribosomal RNA unit. It divides whole living organisms into three groups, we have *Bacteria* at the top,

we have *Archaea*(they are special organisms that lives under hard conditions) and then we have Eucarya (which is obviously the biggest among all groups). We pick those model organisms from important branches of this tree of life so for example, E-coli is shown, yeast as an example of fungi, from animals we have worms, flies, fish, mice, and Arabidopsis, rice, soybean are the examples from plants. So, we try to get these organisms; best representatives from different classes from important branches on this tree of life.

# **Conclusions:**

We conclude the following:

- Sequencing and analysis of full genomes paves the way for future discoveries
- Different model organisms are best source to explore our Genome and to interpolate the results towards the higher organisms.

# **Topic -14 Prokaryotic Genome**

Now, we study the *prokaryotic genome*, prokaryotes are the organisms whose Genetic material (DNA) is not enclosed in a nuclear membrane, so there is no nucleus in them. As there is no nucleus in prokaryotes, there is no justification to have other membrane bound organelles. These are relatively simple cells.



Here, in this diagram we see a prokaryotic cell which is a bacterium (here). We have a genome (DNA) in the shape of a big chromosome in the middle, and ribosomes (small structures important for protein synthesis that occurs in every other organism so ribosomes can also be seen here). It's relatively simple cell, having cell wall with different layers.



Here, in this diagram we see a comparison between a eukaryotic cell and a prokaryotic cell. We can clearly see the membrane bounded organelles in the eukaryotic cell, like mitochondria (involved with the respiration process; food is broken down into the energy. There is a hypothesis that mitochondria actually evolved from bacteria and is known as

endosymbiont hypothesis). Here we can also see the difference in the size of both cells, so eukaryotic cells are complex and bigger than prokaryotes.

The first prokaryotic genome sequenced was that of *Hemophilus influenza* (we have seen in the previous section) and this organism was sequenced in a relatively moderate cost and with an efficient pace that paved the way for sequencing of many other organisms. So study of those prokaryotic organisms is important.

# **Selection Criteria:**

There are different criteria for the selections of the organisms which are then send to the genome sequencing projects. Following are the criteria for selection:

- They had been subjected to a detailed biological analysis/ extensive studies and thus were model organisms.
- They might be important human pathogens (so it's important for us to study its genome).
- > They were of phylogenetic interest.
- Sequences were annotated as they were sequenced.

Table 10.2.	Features of	representative	prokaryoti	c genome
Table 10.2.	Features of	representative	prokaryoti	c genon

Table 10.2.         Features of representative prokaryotic genomes				Here in this table.
Organism (reference)	Phylogenetic group	Genome size (Mbp) (no. protein- encoding genes)	Novel functions	we see different
Escherichia coli (Blattner et al. 1997)	Bacteria	4.6 (4288)	model organism	organisms:
Methanococcus jannaschu (Bult et al. 1996) Hemophilus influenzae (Eleischmann et al. 1995)	Archaea Bacteria	1.66 (1682) <sup>a</sup> 1.83 (1743)	grows at high temperature and pressure and produces methane human pathogen	representative
(Himmelreich et al. 1996) (Himmelreich et al. 1996)	Bacteria	0.82 (676)	human pathogen that grows inside cells; metabolically weak	organisms and
Bacillus subtilis (Kunst et al. 1997) Aquifex aeolicus (Deckert et al. 1998)	Bacteria Bacteria	4.2 (4098) 1.55 (1512) <sup>b</sup>	model organism ancient species, grows at high temperature and can grow in	their genomes.
			a hydrogen, oxygen, carbon dioxide atmosphere in the presence of only mineral salts	<i>E-coli</i> that was sequenced by
<i>Synechocystis</i> sp. (Kaneko et al. 1996a,b	) Bacteria	3.57 (3168)	ancient organism that produces oxygen by light-harvesting; may have oxygenated atmosphere	Blattner et al, the
			Mount	is bacteria, genome

size is 4.6 Mbp (4288 protein encoding genes), and the Novel functions or description of Ecoli is that it is a model organism.

In phylogenetic section, we just have one Archaea whereas rest of them are bacteria.

*Methanococcus* is an archaea with genome size of 1.66 Mbp (1682 protein encoding genes) and it grows at high temperature and pressure and produces methane (maybe a good source to have natural gas from it). Hemophilusis a bacterium with genome size of 1.83 Mbp (1743 protein encoding genes) and is a human pathogen. Mycoplasma is another bacterium with genome size of 0.82 Mbp (676 protein encoding genes) and is also a human pathogen that grown inside cells; metabolically weak. In the end, we have Synechocystis which is again a bacterium with 3.57 Mbp (3168 protein encoding genes) genome size and is an ancient organism that produces oxygen by light-harvesting; may have oxygenated atmosphere.

# **Conclusions:**

We conclude that:

- Prokaryotes are simple Genomes.
- > They are easy models to study Biochemistry, physiology and Molecular biology of life processes.
- > Sequencing is done on economically important organisms (i.e. first it's implemented on simpler genome which is then used to explore complex genomes).

# **Topic -15 Eukaryotic Genome**

# **Introduction:**

We have seen that prokaryotes are simple genomes in comparison to eukaryotes which are relatively complicated. So distinct properties of eukaryotes are mentioned below:

- Eukaryotes have larger genomes
- ➤ Have tandem repeats
- Have introns in their protein-coding genes (introns are between the exons which are the protein coding regions within the genes).
- > Heterochromatin and euchromatin region (eukaryotes have complicated genome, so the chromosome is grouped as heterochromatin; densely packed region and euchromatin; lightly packed region).


Here, in this diagram we can see a typical eukaryotic cell which is pretty stuffed as compare to prokaryotic one. We have nucleus in the middle, channels coming out of the nucleus known as *endoplasmic reticulum* (helps in transportation), ribosomes (for protein synthesis), mitochondria (energy synthesis), we can also see the cytoskeleton that makes the structure of this cell intact and Golgi apparatus (are concerned with the secretion). So complicated membrane bounded organelles are present in the eukaryotes



Here, in this diagram we see the connection between the DNA and the chromosomes.

On the left-hand side, we see a DNA strand that is a 2nm wide strand. So, the DNA wraps over the protein complex molecules (histones - labelled as 1, 2, 3...), and this structure is known as *nucleosome*. Then these histones, turn around and makes a wider structure and it makes a 3nm filament (in third section). Then these nucleosome structures supercoil on their selves to make those further bigger fibres and until they reach the chromosome the width is 1,400 nm. So, if we look into the chromosome, we can recognize that there are different arms in it, which are known as sister *chromatids* (remember this is just one chromosome but we have two chromatids), somewhere in the middle we see a constricted part known as *centromere*, whereas the terminals are known as *telomeres*, (remember these nomenclature while we are discussing the heterochromatin and euchromatin parts).

#### **Staining with dyes:**

So chromosomes if stained with the dyes, they give different coloring patterns, we can come up with the following:

- Dense heterochromatin (dense regions obviously take more color)
- Light Euchromatin (light regions take less color)

If we look into the gene expression, the *heterochromatic regions* are packed so the enzymatic machinery cannot reach there; hence these regions are poorly transcribed (expressed). Whereas, the *euchromatic regions* are highly expressed because they are loosely packed and the enzymatic machinery can easily reach to them.



Here, is the diagram in which we can see the relationship between heterochromatin and euchromatin. You can look into these nucleosomes (combination of DNA and histones), which are quite jammed packed with one another, so definetly the enzymes cannot access the DNA which is embedded inbetween. There are different modifications on the DNA or histones that bring about those structures (we can see on the top), so for example there are histone methylations (in which methyl groups are added to those histones) in that case the system moves towards downside (as shown in the diagram) so it becomes *euchromatin* and similarly, we see that there are some other methylations on some other aminoacids that can move back into the opposite direction also. So, histone methylation, histone deacetylation and there are some other complex proteins which gets attached and give us this *heterochromatin region* and in the reverse process, we get the *euchromatin region*. In the Euchromatin region, the histones are quite spaced and DNA can be accessible. So, this is the reason why the euchromatin region is expressed more as compared to the heterochromatin region.

#### **Conclusions:**

We conclude that:

- Eukaryotes are distinguished by the presence of prominent nuclei
- Eukaryotes have larger genomes, tandem repeats and introns in their protein-coding genes (i.e. they are complicated).

### Topic # 16 Epichromosomal Elements (EEs)

#### Introduction:

Genome is the total collection of genetic material and is made up of

- Chromosomes
- Epichromosomal Elements

#### **Prokaryotic EEs:**

- 1. Plasmids
- 2. Self-replicating
- 3. Additional rings
- 4. Bacteriophages
- 5. Host colonization

- 6. Transposons
- 7. Parasitic DNA elements

#### **Eukaryotic EEs:**

Eukaryotes have extra organelles that contain the genome (DNA) which we call it as *Organellar DNA*.

Examples are:

**Mitochondrial DNA** (both in animals and plants), **Chloroplasts DNA** (in plants), these are membrane-bound organelles and they may be present in hundreds to thousands of copies (so there is also multiple copies of these genomes). Mitochondrion is the site for respiration whereas chloroplast is the site for photosynthesis. Their DNA's can be labeled as mtDNA or cpDNA respectively.

Plasmids, yeast, Transposons, Viral genomes and retroviruses are other examples of organellar DNA.

#### **Endosymbiont hypothesis:**

How do these organelles evolved?

So there is a hypothesis known as *Endosymbiont hypothesis*. According to this hypothesis, these organelles originated as separate prokaryotic organisms that were taken inside a primordial eukaryotic cell. Such symbiotic relationships in which two species are dependent upon one another to varying extents served as crucial elements of the evolutionary progression of eukaryotic cells.

This hypothesis was originally proposed in 1883 by *Andreas Schimper*, but extended by *Lynn Margulis* in the 1980s.

So according to this theory, *Mitochondria* and *Chloroplast* are derived from endosymbiotic bacteria (that got incorporated into the cells).

#### **Organelle Genome:**

Organelle genome (of mtDNA/cell or cpDNA/cell) features are as following:

- Circular
- Double stranded
- Supercoiled
- No histones
- Multiple copies

Genome	Size & Organization
Plant plastid	150 kb circle
Plant mitochondria	150 – 2000 kb multipartite
Human mitochondria	17 kb circle
Saccharomyces mitochondria	75 kb circle

Here, in this table we can see the size of these genomes. For example, *plant genome* is 150kb circular genome, *plant mitochondria* is 150-2000kb multipartite, *human mitochondria* is 17kb circular and *saccharomyces mitochondria* is 75kb circular.

#### \*Mostly the genomes are circular.

As far as the expression of these organelle genomes is concerned, it has been observed that their functions are actually dependent on nuclear genomes (they cannot make functions for themselves).

They encode only a subset of genes required to elaborate a functional organelle like rRNAs, tRNAs, ribosomal proteins, membrane-associated respiratory or photosynthetic components.

Other components which are encoded by nuclear genome are translated in the cytosol of the cell and are imported into the organelle. It has been observed that 10% of nuclear genes are devoted to mitochondrial function whereas 15% to plastid function.

#### **Conclusions:**

We conclude the following:

- Organelle genome is similar to prokaryotes.
- It is in high copy number.
- The mtDNA and cpDNA depends on Nuclear DNA (genome) for their function.

## Topic # 17 Genome Repeats

We have seen that the major proportion of genomes in the eukaryotes is made up of repeats (they are also present in prokaryotes). So let's look into what these repeats actually are.

#### **Sequence Repeats:**

These repeats *skew the base composition* (normally the A's, T's, G's and C's relative proportion is similar to one another but if repeats are present and these repeats are of same types, for example if we have runs of GC's, then obviously they'll change the proportion of different bases) which can contribute to having differences in there buoyant densities (so those fragments can then be separated on the basis of those differential densities).

The repeat containing DNA can be separated as satellite DNA on the bases of these densities.



Here, is an example in which we can see the repeats, they can be *Tandem repeats* (array of repeats together), *interspersed repeats* (those repeats which are separated by normal genome followed by repeats i.e. normal genome is between those repeats).

The link to this figure is:

http://mcb1.ims.abdn.ac.uk/djs/web/lectures/repeats1.html#anchor10305

#### Satellite DNA:

Satellite DNA has following features:

- It may be one to several thousand bp long and it can also be present as *Tandem*; array of 100 million bases long.
- > They are present near centromere and telomere and
- > They can be classified as *Mini-satellite* and *Micro-satellite*.

#### Mini-satellite:

Mini-satellite features are as follows:

- They are 15 bases long in array of several hundred to thousands kb.
- They are typically present in euchromatin region.
- Example is VNTR and is used to identify human individuals in forensics.



Here, in this figure we can have different repeat patterns. In the individual #1, we have those GC repeats (runs of GC's together), and obviously the organisms are diploid i.e. they are having two chromosomes, from their parents. So, the same Allele A2 has two of those repeats, if we look into another locus (gene position), we can see there is Allele B2 that has

AGCT repeat (2 copies in both allele) and in the second individual we have 3 copies in 1<sup>st</sup> allele and 2 copies in second allele. So, when we digest them with restriction enzyme (restriction enzyme cut them on the repeat regions), we can have *differential banding patterns* when we run them on gel. In this way, we can recognize those individuals. In the figure (below), we can see the bands, on the left-hand side are the bands of the individual#1 and on the right-hand side are the bands of the individual#2. In this way, we are getting those repeat regions and by running them on the gel, we can detect which DNA belongs to which organism.

#### Microsatellite:

Micro-satellite features are as follows:

- 1. They are 2-6 bases long and can be in arrays of 10-100 bases.
- 2. They are inherited to offspring.
- 3. Mainly they are useful markers for genetic analysis and evolutionary studies (just like in previous example)
- 4. They are found in telomeres TTAGGG (one example of six nucleotides segment).
- 5. SSRs and STRs (Simple Segment Repeats and Short Tandem Repeats are typical examples).

#### **Transposable Elements (TEs):**

These are also important kind of repeats and their features are as follows:

- 1. They are making up the larger proportion of the eukaryotic genome.
- 2. They thought to play an important role in the evolution of these genomes.
- 3. They move (Jump) from one location to another even faster than chromosome replicate (and are known as jumping genes).
- 4. They have a potential to increase in number,
- 5. So they make up a large proportion of eukaryotic genome.
- 6. They are detectable but sometimes they blend into the genome due to mutations and cannot be detected.



Here, in this diagram, we can see the proportion of these transposable elements (red), whereas the green ones are the other sequences. We have different organisms like *Homo sapiens*, *Z. mays*, *Drosophila melanogaster*, *Arabidopsis*, *C. elegans and S. cerevisiae* (yeast).

We can see that in human genome, these Transposable elements make up 35% (huge number), in *Z. mays* they are even more than that i.e. 50%, in *Drosophila melanogaster*, they are comparatively less i.e. 15% genome

(y- axis represents the genome size), and in rest of the organisms, they have less proportions of those repeats.

So, mainly in humans and Z. mays, they are present as a major proportion.

#### **Conclusions:**

We conclude that:

- Large proportion of eukaryotic genome is composed of repeats
- Different repeats act as markers to detect genetic variation (of organisms) and are also used to study evolution of those organisms

### **Topic . 18 Transposable Elements (TEs)**

#### Introduction

Transposable Elements are the elements which can transpose i.e. they can move from one place to another in the genome and then they can cause the repetitive elements (repetitive DNA within that genome).

#### **Insertion Sequences (IS elements)**

These are the simplest transposable elements and their features are as follows:

- They code only for the ability to transpose and are found in prokaryotes.
- They are usually very small (< 1 Kb to 2 Kb)</p>
- They are flanked by inverted repeat sequences (IRs).
- > They encode at least one gene that provides their own transposition functions.
- They do not code for noticeable (phenotypic) traits.
- > They can cause mutations by transposition into genes.



Here, we see a structure of *Insertion Sequences* (IS elements). We have *Invert Repeats* (IR) which are antiparallel or facing each other (repetitive elements; these are the distinctions in them). We have a transposase gene that gives the transposition properties to transposable

elements. Another distinction in them is that when they get into the host genomes, they cause *target site duplication*- normally they create the sticky ends in the genomes of the host and later on when the complementary nucleotides are formed, they become those duplications (in pink).

#### **Transposons:**

Transposons are more complex transposable elements as compared to the simple IS (*Insertion Sequence*) elements and they code for additional characters in addition to the gene responsible for their transposition.

	3
-	
ORFZ	AAAAAAA
444	
	ORF2

Here, in this diagram, we can see that there are two major classes termed as Class I and Class II.

Within Class I, we have LTR (Long Terminal Repeats) retrotransposon (they have the ability of reverse transcription, so they have RNA's which gets converted into cDNA's; this is how they replicate). They have LTR at both ends (they are not invert but they are kind of forward repeats and are in the same direction).

In the same Class I, we have another category known as **Non-LTR** (Long

Terminal Repeats) retrotransposon, which doesn't have those LTR and are of two types namely LINE (Long Interspersed Nuclear Elements) and are larger in size and SINE (Small Interspersed Nuclear Elements) and are smaller in size. In Class II, we have DNA transposon.

#### **Eukaryotic TEs:**

As we have seen earlier, eukaryotic Transposable Elements has two classes in them; Class I and Class II.

In Class I, they have reverse transcriptase in them so they use RNA-mediated mechanisms of transcription. This Class I can be categorized into*LTR (Long Terminal Repeats) retrotransposons*, simple *Retrotransposons* and *Retrovirus like Elements*.

As we have seen earlier, that **Non-LTR** (Long Terminal Repeats) retrotransposon comprises of **SINES** and **LINES.SINES** are Short Interspersed Nuclear Elements and are **80-300 bp** long and the example is *Alurepeats* (found in humans). Whereas **LINES** (Long Interspersed Nuclear Elements) are **6-8 kb** long and are relatively longer than SINES.

The example of **Class II TEs** is *Ac-Ds* (which were studied in maize - the work of Barbara Mcclintock led to the discovery of these transposons where he gave different characteristics to the kernels of the maize). Another example of **Class II TEs** is *P elements in Drosophila*.

Additional elements are known as **MITES** (Miniature Inverted Repeats TEs) that has the features of both Class I and II and are 400 bp long.



Figure 10.3. Comparison of genome composition in four genomes. (A) Human  $\beta$  T-cell receptor locus on chromosome 7. V28 and V29.1 encode parts of the  $\beta$  T-cell receptor proteins that are joined during development of the immune system (Rowen et al. 1996). TRY4, the gene for trypsinogen, and TRY5, a pseudogene related to the trypsinogen family, are not related to the receptor sequence. Why they are located here is not known. (B) Segment of yeast chromosome III (Oliver et al. 1992). (C, D) 50-kb fragments of the maize and *E. coli* chromosomes, respectively (SanMiguel et al. 1996; Blattner et al. 1997). The maize repeats are LTR retrovirus-like elements (Fig. 10.2) that have inserted within the last 3 million years (SanMiguel et al. 1998). (Redrawn, with permission, from Brown 1999 [BIOS Scientific].)

Here, in this figure, we see the comparison of human, yeast, maize and E-coli.

In blue color, we see those repetitive regions (transposons), red are the exons, whereas orange are introns and brown colored are the pseudo-genes.

We are comparing the 50kb region of the genome where in human, we can see the blue color prevails i.e. many transposable elements can be seen. In yeast the transposons are less in number. Whereas in maize, they are found to be in huge proportion and in E-coli we can see the IS (Insertion Sequences) elements which are extremely low in number.

#### **Conclusions:**

We conclude the following about Transposable elements:

- They make up a significant part of organisms' genome especially in that of the eukaryotic genome.
- > They move within and across genomes and
- Causes genome expansion.

### **Topic # 19 Eukaryotic Gene Structure**

#### **Eukaryotic genes:**

Eukaryotic genes are relatively complicated and are not simple as prokaryotic genes. They possess exons and introns.

#### Exons:

Exons are protein coding regions and are interrupted with introns (i.e. in between exons are introns). During the gene expression, both exons and introns are first transcribed into mRNA and then the introns are removed out, so the remaining structure is known as ORF (Open Reading Frames) which only consists of the exons.

#### **Introns:**

They contribute a very small proportion in yeast i.e. only 239 introns in its genome whereas in human the introns makes up 95% of its genome.

The introns stay on the same location and they might also have embedded genes in them.

They can be distinguished by the presence of GT at the 5' ends and AG towards the 3' end (**GT**-----**AG**) and this trend is highly preserved all over the genome.



Here, in this diagram is the structure of a typical eukaryotic gene.

We see the chromosome and gene is the region which has specific patterns so we can observe the promoter region (in the beginning of a

gene), the blue ones are the exons and those orange ones are introns.

There is start of transcription (marked by black line) which ends at the exon3 (as shown here and marked by black line), this whole region is then transcribed into mRNA. We can see a 5'-UTR (Un-translated region) region, so this region is transcribed into mRNA but is not translated i.e. no protein is formed from this region, similarly we also have a 3'-UTR region.

When we see the ORF i.e. the region from start codon (initiator) to stop codon, and in between them we can see there are number of amino-acids, so this is the region from where translation takes place and we get a protein.

After transcription, the transcript is known as *Primary RNA transcript* and we can see that it also contains those introns. Which are later on removed through a process called as splicing and then we get a *Mature RNA transcript*, so that transcript is then translated into the proteins. This mature RNA transcript is also recognized by the presence of a poly-A tail (long runs of A's)

#### **Intron origin**

So, about the origin of introns, there are two theories which are as follows:

- **Intron-early** According to this, they used to assemble the genes from already existing exons (so they brought the exons together and then these structures became the genes).
- **Intron-late** According to this, the exons were already present with one another, then introns got into them (i.e. they Broke up previously continuous genes by inserting into them).

#### Number of Genes:

Now we talk about the degree of compactness, so the compact genomes whose size is small and the relative proportion of gene is higher which contributes to the variation in gene density. In short, we can say that compact genomes have higher genome density.

Organism	Genome size	Predicted genes
	(haploid MB)	
<i>A. thaliana</i> (plant)	130	~25,000
C. elagans (worm)	100	18,424
Drosphila melangaster	180	13,601
Escherichia coli	4.7	4,288
<i>Homo sapiens</i> (human)	3000	45,000 - 120,000
S. cerevisiae (yeast)	13.5	6,241
	Mount: Table	10.3 11.3 2 <sup>nd</sup> edr

Here, in this table, we have different genomes (mostly eukaryotes).

We can see that the size of genome in Arabidopsis is 130MB and number of genes is approximately around 25,000.

E-coli (prokaryote; bacteria) can be seen here, with genome size of 4.7MB and there are over 4,288 genes.

In humans, the genome size is 3000 MB (it's not megabytes, it's mega base pairs), and 45,000 to 120,000 genes (slightly around 30,000 have been identified).

So, if we look into those smaller organisms, like those having smaller genomes (E-coli), and if we take the number of genes and divide that to the genome size, we'll see that they have more densities as compared to the larger genomes.

#### **Pseudogenes:**

These are non-functional genes (sometimes there are mutations in the genes and if those mutations are present in some important regions then the genes' functions gets knocked out – known as psudogenes).

There is one category of them and is known as processed pseudogenes that lack introns and promoters.



Here, is the diagram in which we see the normal gene and a psuedogene.

In psudogene, we see that the bases are deleted from the

promoter region (promoter must have specific pattern of bases in them), so in this case, this deletion is lethal. Similarly, in start codon (initiation code) we can see GTG instead of ATG, and we can also see the mutations in the splice sites (its GC rather than GT and GA rather than GT respectively). And in exon2, there are 20 bases which are also been deleted. Hence, this is how a normal gene has now become a pseudogene.

#### Gene families:

Sometimes, the set of genes have similar sequences as well as similar functions. And if we try to find out similar genes within an organism or between different organisms, performing similar functions can be categorized as gene families. Gene families arise from gene duplication and subsequent divergence events.



FIGURE 27.29. Gene duplications during the evolution of the human globin gene families. The initial split gave rise to two lineages, one leading to the modern gene for myoglobin and the other to the globin genes. Subsequently, the proto- $\alpha$ -globin and proto- $\beta$ -globin lineages split following a duplication. Other duplications took place within the  $\alpha$  and  $\beta$  lineages. (Modified from Strachan T. and Read A.P. *Human Molecular Genetics 2*, Fig. 14.16,  $\bigcirc$  1999 Garland Science.)

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#### **Conclusions:**

In the end, we conclude the following:

- Eukaryotic genes have exons and introns.
- > Introns make up a significant portion of higher organisms' genome (Human genome).
- Pseudo genes are non-functional genes.
- > The genes which are similar in function, they make up the gene families

### **Topic # 20 Comparative Genomics**

#### **Introduction:**

Comparative genomics is where we compare the genomes of different organisms like we do comparison of gene number, gene content and gene location in both prokaryotic and eukaryotic groups of organisms.

The availability of genome makes it possible to have a comparison of all the proteins; proteome (so we can have the genome and translate them into the proteins or we can get the actual proteins and can do those protein comparisons – known as comparative proteomics).

#### **Orthologs:**

Orthologs are the genes that are present in two different organisms and are so similar that they must have the same function and the evolutionary history (Fitch, 1970).

#### **Paralogs:**

Paralogs are the gene families that originate from gene duplication events (they maybe within the same organism) over the evolutionary time.

Here, is the diagram showing the example of gene duplication, we have human globin gene.

So, first duplications give rise to two types of globins (one is myoglobin and other globins).

Within the second type (globins), we have alpha globin and and beta globin.

So this process is called as gene duplication.

(Unlike pseudogenes, 2<sup>nd</sup> copy of gene remains functional)



Here, in this diagram we can see that the similar genes are called as *homologs*, which is further divided into *orthologs* and *paralogs* categories.

The *Orthologs*, for example if we start from the bottom, we can see some globin gene (early) and then there is a gene duplication (as seen earlier), so it becomes a beta chain gene and an alpha chain gene.

Then those alpha chain genes

can be seen in three different organisms (frog, chick and mouse) and are similar so that's why they are known as orthologs. Similarly, those beta chain genes can also be seen in three different organisms.

Now, if we look into those genes that are similar but gets diverted (and are in the same organism) as in mouse, chick and frog (we have alpha globins and beta globins) so these organisms maybe classified as paralogs.

#### Drosophila and yeast:

When *Drosophila* is compared with yeast, we see that *Drosophila*has core proteome only twice the size of that of the yeast and *Drosophila* proteome is comparatively more similar to mammalian proteomes than worm or yeast.

#### Drosophila and C. elegans:

Now if we compare the fly *Drosophila* with the worm*C*. *elegans*, we can see that despite the large differences between them, the core proteome is of the similar size in both.

And nearly 30% of the fly genes have putative orthologs in the worm.

#### Drosophila and Human:

When we compare fly *Drosophila* with Human, interestingly some human disease genes are absent in *Drosophila* but we can see a number of previously unknown counterparts to human cancer and neurological disorders genes are present in *Drosophila*, so it can be used as a good model in cancer studies.

#### **Conclusions:**

We conclude the following:

- Comparative genomics reveals the relationship among different organisms.
- Fruit fly has more similarities with mammals, so we can utilize it especially as a model for cancer studies

### **Topic #21 Comparative Proteomics Introduction**

#### Introduction:

Since, genes encode proteins and these proteins perform actual life functions, we can have the genomes and get the translated protein from it and by comparing those translated proteins with one another, we can say it's a Comparative Proteomics study.

The collection of the protein sequences that are encoded by the genome makes up the proteome of that individual.



Here, is the figure of how comparative proteomics can be done.

So from the genome sequence, we can get the predicted genes out of it, then we do translations which gives us the proteome of that individual. We

can use those proteins and get some Database searches (we can find the homologous proteins, whereby we can predict their functions and roles).

We can also do the comparison of proteome by themselves so that will help us finding the paralogs (studied earlier) within that individual.

We can also do the comparison of proteins between different organisms and we can also find the clusters of co-related proteins in terms of their sequences and functions.

#### All against all, self-comparison:

Firstly we talk about the all against all; self-comparison, where we can do the following:

- Comparison of all proteins with each other within the individuals' proteome.
- Identify unique proteins from the ones having paralogs.
- Identify Gene families
- If we have a good match between query sequence and some other sequence, we can



suspect those two are the paralogs (because they are present within the same organism).

Here, in this figure we have the example of database searches.

We have some proteins which are aligned together.

Here, we can see some proteins which are similar to one another and as they are sharing the similar domains (domains are specific protein structure which is made up of specific amino-acids into particular structure). We do the sequence similarity searches by the software named as **BLAST** (will be discussed later).

We have some parameters (P/E), normally low P/E ratio is taken as a good score, for example if we have a good match we can have the range of P/E value less than  $10^{-20}$  and it keeps on decreasing if we go down to other similar sequences (as shown).

#### **Cluster Analysis:**

In cluster analysis, we make the groups of proteins which are quite similar to one another. And reasons of doing it are as follows:

• To sort out the relationships of all the related proteins.

- Clustering classify the proteins based on some objective criteria e.g.
  - E value cut off
  - *Distance in alignment* (so the proteins which are more similar will be grouped together and distant proteins will be grouped from them so in this way we can have sub-groups or clusters in our data).

There are different clustering methods and are explained briefly in the below section.

#### **Clustering by subgraph:**

The way of clustering or grouping by the method of sub-graph is as follows:

- Each sequence is a vertex (vertex or vertices are the point or dots by which the edges (links) in a graph are connected. There can also be a vertex that's without any edge connecting to it, known as isolated vertex).
- Significant alignment score is an edge (on the vertices, we put our sequences and on the edges we put our alignment scores).
- Trimming by removing weak edges (if we have High P/E ratio, we will remove them).



Here, in this diagram we are representing the clustering, we have those vertices (shown as balls) which are the proteomes here, we have the edges (lines) which are the connection between those vertices and are the alignment scores (the thickness of edges relates with the small P/E value).

And if they are weakly co-related like the values are greater than  $10^{-6}$ , so then they are not connected.

The dashed lines (or dotted lines) are where we have loose connections.

#### **Clustering by Linkage :**

There is another technique, where the clustering is done by linkage (almost similar to the previous ones except some changes). The method of doing it is as follows:

- Each sequence is a vertex.
- Significant alignment score is an edge.
- Trimming by removing weak edges (High P/E).
- Or remove >  $e^{-6}$  (we remove the ones which are not linked).
- Remaining sub-graph should share 2/3<sup>rd</sup> of edges.

#### Single Linkage:

Linkage is done by the following method:

- A group of sequences in all-against-all comparison is subjected to MSA (group those proteins which are co-related with multiple sequence alignment by first aligning them and then calculating their distances).
- Create distance matrix (by using those distance calculation just made).
- Neigbour joining is then used to do clustering (by distance matrices, we create those trees and the method used is *Neighbor joining* will be discussed later).



Here, in this figure we have the single linkage cluster analysis results, so we have those proteomes which are present in the end of those trees which we call them as leaves. The closely related ones are

connected with one another.

We can see two types of arrangements, one is circular arrangement (on left) and we have typical binary tree arrangement (on right).

In binary tree, we can see that we have come up with two big clusters, and within those clusters we find those further groups or sub-clusters.

#### **Core Proteome:**

Core Proteome is when we do *all-against-all comparison*; which tells us about the proteins which are duplicated and it also gives us the information about those proteins which are uniquely present in those organisms- the core proteome.

Organism	Total number of genes	Number of gene families <sup>a</sup>	Number of duplicated genes <sup>b</sup>		
Hemophilus influenzae (bacteria)	1709	1425 <sup>c</sup>	284		
Saccharomyces cerevisiae (yeast)	6241	4383	1858		
Caenorhabditis elegans (worm)	18,424	9453	8971		
Drosophila melanogaster (fly)	13, 600	8065	5536		
<sup>a</sup> The number of clustered groups core proteome of the organism. <sup>b</sup> Count of number of duplicated p	in the all-against-all analysis u genes within the protein famil	sing the algorithm described in the y clusters.	e text. This number represents the		

Here, in this table, we compare the core proteome with the total number of genes.

Here, for example in

bacteria we have 17 hundred genes, 14 hundred are the gene families (represents the core proteome) and we have number of duplicated genes. In worm, we can see that we have 18 thousand genes where almost half of them are in the shape of gene families and rests of them are duplicated genes (so almost half of them are duplicated amongst them). In *Drosophila*, we have 13 thousand genes, 8 thousand gene families and 5536 are the duplicated genes.

#### **Conclusions:**

We conclude the following:

- Genome is translated into proteome.
  - Self-comparison of proteome yields gene families and duplications

### **Topic # 22 Between-Proteome comparisons**

#### Introduction:

In between-proteome comparisons, we compare the proteomes from different organisms with one another, so in this way we can find the genes which are similar between those organisms, and we call them as orthologous genes.

Here we take the proteome as a query and we do a database similarity search against another proteome or there can be a whole database where we have the set of proteomes.

If the proteome is not available, we can search the EST (Expressed Sequence Tag) Database as well.

#### Significance:

As mentioned earlier, this helps in finding the <u>orthologs</u>, <u>gene families</u> and the <u>domains</u> (between different organisms). There can be other significance of *between-proteome comparisons search* and are as follows:

- Proteins that have a highly significant alignment score can be suspected as the orthologs.
- Mostly the proteins that are related to core biological functions (basic functions of life) are likely to be orthologs.

#### Finding true orthologs:

How can we find true orthologs? There is a technique which we call it as:

#### Method 1:

Where we have the "**Reciprocal Hits**", in this method, you take one organism at a time as a query and search against the other as a database and then you flip around and take first as database and second one as a query. So, if you are getting similar end results, it means that the same genes are co-related with one another (or highly similar to one another), and we can keep them as the best hits.

We can also apply some criteria on hit, for example here we can do a E-value cut off when we do BLAST (BLAST gives us the parameter called as E-value and a lower value is considered good and we'll talk about BLAST algorithm in coming up lectures but here the point is to let you know that we do some E-value cut off) and we can retain those gene pairs in those Reciprocal hits.

So, for example here we say that E < 0.01, we can retain them.

Similarly, while we are doing BLAST, since it is a local search tool, we can compare different regions amongst different genes or proteins, we look for the matches between different regions and sometimes both proteins are not greatly covered in the alignment, so we want to have at least some coverage criteria, so here we have like for example 60% coverage.

So we keep those matched pairs normally with a very conservative or low P value like  $10^{-10}$  to  $10^{-100}$ .

#### **Clusters of Orthologous Group (COG):**

In this way, while we are finding the true orthologs, we can group those organisms which are similar to one another and this method is known as Clusters of Orthologous Group (COG).

Orthologs are assumed to be derived from common ancestor, so they might also have paralogs (within the organisms, the orthologs might also go through the duplications).

Orthologs are clustered to form COG (can be studied as COGs).

Table 10.5.         Numbers of closely related yeast and worm sequences										
Cut-off P value	$< 10^{-10}$	$< 10^{-20}$	$< 10^{-50}$	$< 10^{-100}$						
Total number of sequence groups	1171	984	552	236						
Number of groups with more than two members	560	442	230	79						
Number and percent of all yeast proteins (6217) represented in	2697 (40)	1848 (30)	888 (14)	330 (5)						
groups										
Number and percent of all worm proteins represented in groups	3653 (19)	2497 (13)	1094 (6)	370 (2)						

Adapted, with permission, from Chervitz et al. 1998 (copyright AAAS).

Here, in this table we have the example of the closely related organisms' sequences i.e. Orthologs (between yeast and worms).

We have different P cut-offs (like  $<10^{-10}$ ,  $<10^{-20}$ ,  $<10^{-50}$ ,  $<10^{-10}$ ).

We have the '*total number of sequence groups*' and at different cut offs if we go on a stringent criteria, we have less common orthologous groups and if keep the criteria less strict we can have more of these orthologous groups.

Then we have the 'number of groups with more than two members' (as shown). Lastly, we have the 'percentage of yeast' and 'percentage of worm' (i.e. how many amongst the total, they are present) and are presented in these two groups (the yeast and the worm), say we have 40 percent and 19 percent on  $<10^{-10}$  cut-off P-value, and we have 5 percent and 2 percent on  $<10^{-100}$  cut-off P-value (if our criteria is strict).

So, in this way we can group the similar proteins at different cut offs of P-values, and we can have the various results.

#### Proteomes to EST databases:

Sometimes, we take those proteomes and we match them or align them with Expressed Sequence Tags (EST) (which is cDNA copies of cell's mRNA sequences). We do this procedure for those organisms' genomes whose sequences are not available.

ESTs are single DNA reads and are mostly 3' biased (since we get them from mRNA and mRNA extraction protocols relies on getting those mRNA by using their 3-prime poly A-tail which is present on their 3-prime end, so that is why they are kind of more tiled or oriented towards 3-prime ends as they are mainly extracted from this site).

EST may be incomplete because it is wholly dependent upon the gene expression, so if we do not have genomes rather than we only have the ESTs, we might be biased towards only those genes which are expressed.

The softwareor the package in BLAST which is being frequently used for this purpose is **TBLASTN**.

#### Family and Domain Analysis:

Proteins are organized into domains that represent modules of structure or function (as domains are specific arrangements of amino-acids). And domain comparison sometimes is corelated with their biological functions.



Here, in this figure for an example, we take the domains from different proteins and we put them into *Domain Databases*, and then in the end we can come up with the shared domains; the domains which are present in these different

groups and lastly we can co-relate this information to have an idea about their functions.

#### **Conclusions:**

- Proteome comparison helps finding orthologs, gene families and protein domains
- Domain comparison reveals their biological roles.

## **Topic #23 Horizontal Gene Transfer**

#### **Introduction:**

Horizontal Gene Transfer is where the genes are transferred at the same levels or horizontally (so two organisms they transfer their genes and it's not like transferring the genes from top to bottom like vertical transfers for example, from parent to the offspring) and is relatively slow process in evolution.



Here, in this chart we have the length of the coding sequences and the proportions which are coming from the foreign elements (in black). (Grey are the native ones).

So this is an indication of the horizontal gene transfer. We can see here that mostly in E-coli K12, there is the biggest portion of foreign gene and in

Synechocystis, we can observe the huge part of foreign genes as well.



0157-H7 has about 1,400 genes not present in K12 K12 has about 500 genes not present in 0157-H7

Green: Genes conserved in both bacteria Red: Genes present in 0157:H7 but not in K12 Here, is another case in which there is the comparative map of two genomes from E-coli 0157-h7 and K12.

We can see that here are about 1400 genes present in H7 which are not present in K12.

Similarly there are 500 genes found in K12 which are not seen to be in

#### H7.

In this circular arrangement, we see the red which indicates the genes present in H7 but not in K12. Whereas green are the ones which are conserved in both the organisms.

Other Examples of Genes of Foreign Origin in Complex Eukaryotes								
Eukaryote Fo	reign Genes Sour	ce						
Various Plants	Hormone synthases	Bacterial						
Aphids	Carotenoids	Fungal						
Sturgeons	Various (15 genes)	Trematodes						
Sea Slug	psoB, encodes a nuclear f	actor Alga						

Here, is the table of another example in which we observe the horizontal gene transfer in eukaryotes.

We have different plants, which has *hormone synthases gene* (a foreign gene) and is suspected to be coming from the bacteria.

The Aphids (insects) which has from fungal

carotenoids as a foreign gene and it might be coming from fungal.

Sturgeons have some 15 genes which are foreign and are thought to be coming from Trematodes.

Sea Slug that has a *psoB*gene which encodes a nuclear factor and the source is some Alga.

#### **Conclusion:**

Horizontal Transfer of genes between different organisms is a relatively slow process that leads to acquisition of new traits.



#### **Gene Mapping**

• Gene mapping is determining the location of and relative distances between genes on a chromosome

#### Genetic vs physical Mapping

• Genetic map distances are based on the genetic linkage information measured in Centi-Morgans (CM)

#### **Genetic vs physical Mapping**

• Physical maps use actual physical distances usually measured in number of base pairs

#### Synteny

- Arrangement of genes on the chromosomes
- Comparing gene orders provides a good measure of similarities and differences among different organisms

#### **Conserved Synteny**

- Species diverged from common ancestor have similar chromosomes and gene order
- Gene duplication and rearrangements changes synteny
- As a results species diverge



#### Gene Order (Synteny)

tr	PE	trp	D	trp C	trp B	trp A	
1 Haemop	264 hilus ir	126 nfluenza	e i	1262	1261	1260	
trp E 🔰 trj	G	trp D	trp (	- /	trp B	trp A	
1387 13 Helicoba	cter py	1389 Jori	1389.	•	1431	1432	
trp E	trp	G trp	D	trp C	trp B	trp A	
1282	128	1 128	BO 1	1279	1278	1277	
Bacillus	ubtilis						
t	rp E	trp D	trp C	trp F	trp B	trp A	trp G
	2264	2263	2262	2261	2260	2259	75





Species diverged from a common, expected to have similar synteny

Functionally related genes stay close as clusters

## **Topic # 25 Genome Annotations**

#### Genome annotation:

After the genomes are assembled together, the genome annotation is a very important task whereby we could gain information of the genome. It has several procedures involved and which are as follows:

- We try to locate some important genes which are protein coding genes and also their products.
- > We also try locating the RNA-encoding genes.
- > We recognize the non-coding regions in a genome.
- > We can also predict the function of the genes.
- > The Biochemistry and structure of gene products can also be obtained.
- > We can explore the Literature links.

- We can also explore the links to genetic maps where they are located on the chromosomes.
- We can look into the location of the repeats.
- ➢ We can also look into the location of STS (sequence tag sites).
- > We can also look into the location of sequence polymorphisms.
- And we can find the significant alignment to some protein sequences of known function in databases (by comparison).

#### **Annotations steps:**

Annotations are divided into two types and are as follows:

- Structural annotation
- Functional annotation

#### **Structural Annotation:**

Structural Annotation is where we try to identify certain gene features like;

- Promoters
- Terminators
- Shine-Dalgarno sites; the ribosomal binding sites during the protein synthesis)
- DNA motifs (patterns of nucleotides within the genes)
- Co-transcription units
- Operons in microbes (in micro-organisms, lots of genes are transcribed together known as operons)

#### **Annotations Tools:**

There are two tools which are important worth mentioning here, one is MAGPIE and the other is GENEQUIZ and these are designed to assist with gene the genome annotations.

**MAGPIE** (Multipurpose Automated Genome Project Investigation Environment) - It's an automated genome analysis tool that is used for structural annotation.

**GENEQUIZ-** Focuses on deriving a predicted protein function based upon the available evidence; including evaluation of similarity to the closest homologue in the database (i.e. it is good tool for functional annotation).



Here, in this figure, we have an outline and is a kind of a work flow of how exactly MAGPIE works.

We take some *source sequence*, and we give it to some software program known as *Magpie Daemon;* it takes the sequences (which are added to the database, so it automatically gets them) and sends that data to the *local tools*,

*remote tools* (over the internet) and then it explores some specific features or annotation patterns and it puts them into the *Feature Database*.

Then later on those results are interpreted (*Interpretation and Reconstruction*) and then we get the *reports*.

In this way, we have a kind of automated annotation gathering tool

#### **Functional Annotations:**

The attributing biological information to the genes is called as functional Annotations and we can have it via;

- Biological function
- Biochemical function
- Gene expression (transcription of the gene is considered as gene expression)
- Regulation and interactions among different genes

#### 8 Group Classifications:

There are different classification schemes, which are meant for functional classification, to classify the genes and their products into one of these groups;

- Enzymes
- Transporters
- Regulators
- > Membranes
- Structural elements
- Protein factors
- > Leader peptides (control transcription and translation)
- Carriers (transporters)

In this way, scientists have seen that 90% of the *E-coli* genes fit into these categories (so their annotations can be explained).

#### **Enzyme Commission (EC) numbers:**

It is another scheme which was put forward by the Enzyme Commission (EC) that was working under the IUBMB (International Union for Biochemistry and Molecular Biology).

They say that the enzymes are classified on the basis of the reactions they catalyze and have a 4-digit scheme which is actually the enzyme commission number: **EC a.b.c.d** 

**'a'** (first digit) informs that it is from one of the 6 classes of biochemical reactions (enzyme might be coming from one of these classes).

**'b'** (second digit) informs that is from the group of substrate (the thing on which the enzyme attacks).

**'c'** (third digit) informs us that it is anaccepter molecule.

'd' (fourth digit) gives the details of biochemical reaction

#### For example,

tripeptideaminopeptidases

#### EC 3.4.11.4

Where 3 - tells us that it is a Hydrolase (use water to break substrate).

This **3.4-** tell us that it is a Hydrolase that acts on the peptide bonds.

The **3.4.11-** tells us that it is a Hydrolase that cleaves the amino terminal amino acids of polypeptide.

While putting everything together, **EC 3.4.11.4**- it tells us that it is a Hydrolase that cleaves the amino terminal amino acids of a tri-peptide.

# With Enzyme Commission Scheme, they classified that 70% of E-coli genes shared a and b (first two classes), which means that they catalyzes the same biochemical reaction.

#### **Three Groups Scheme:**

This is another classification scheme known as a 'Three Groups Scheme', where we divide all those functions which are related to the following:

• Energy

- Information
- Communication

It was found that plants devotes half of their genome to the **energy metabolism** (they make food), whereas animals devotes half of their genome to **communication** (they talk a lot :D )

#### **Conclusions:**

We conclude the following:

- Finding genes and their coding regions is an important task in Genome annotations.
- Functional annotations correlate the genes to different classes of functions

# **Topic # 26 Genome Sequencing**

#### Introduction

Genome sequencing involves recognition of nucleotides in a Genome and determining their precise order of arrangement. And we have seen that the advances in sequencing technologies have revolutionized the pace of scientific discovery.

DNA sequencing began in 1970s with the development of Maxam-Gilbert's method and later got the pace with Sanger's method so most of the modern day sequencing employs the variants of Sanger's methods.

### First published Nucleotides



Here, we see the first published nucleotide, which was published by Gilbert and Maxam.

They presented the first 24 base pairs of the DNA (working on the *lac* Operator).

#### First genome sequenced:

So first genome was RNA of a virus (phage) **MS2** and was sequenced by **Walter Fiers** and colleagues at University of Ghent, Belgium (1972-76).We can say that RNA was first sequenced among nucleic acids (first complete genome sequence made was that of RNA).

Among the free living organisms, *Haemophilusinfluenzae* was the first ever published genome by Fleischmann et al. in 1995. (Since we are in a discussion that viruses are living or non-living creatures, so if we talk about the living ones, it is the bacterium which was the first living organism whose published genome was made).

#### MS2:

MS2 is a virus that infects *E-Coli* and *Enterobacteriaceae*. It is a single stranded sense RNA (in the host it gets duplicated and where anti-sense is formed and from that the RNA copies are again formed) and it encodes MS2 coat protein.

#### MS2 sequencing:

The entire nucleotide sequence was established by nuclease digestion and characterization of a fragment (it had the enzymatic digestion of the nucleotides and that's how it was sequenced).

#### MS2 genome

MS2 has 49 different codons in the genetic code that specify the sequence of the 129 aminoacids long coat polypeptide (virus has a coat which is made up of proteins on its outer side and it has a RNA; it's genome).



Here, we see the virus genome, which has the genes like *mat* (helps in assembly; putting those proteins together), *cp* (codes for coat protein), *rep* (codes for replicase protein) and *lys*(codes for lysis

protein; that breaks the host cells). When we observe cp, rep, andlys, we can see the lysgene is

DNA Sequencing History	
1870	Miescher/Discovered DNA
1940	Avery/Proposed DNA as "Genetic Material"
1953	Watson & Crick/Discovered Double Helix Structure
	<ul> <li>Specific RNA digestion and chromatography methods were used to sequence RNA; it required large quantities of sample.</li> </ul>
1965	Holley/Sequenced Yeast tRNA <sup>ALA</sup>
	<ul> <li>Primed synthesis concept and 2-D electrophoresis were use samples were labeled and less material was required.</li> </ul>
1970	Wu/Sequenced $\lambda$ cohesive end DNA

embedded between these two genes, so we can have the genes within the genes (here).

Here, is the history of different milestones, which were done while we are moving towards this DNA sequencing.

So we start way back in 1870 with the discovery of DNA, then in 1953, we see the major breakthrough is Watson and Crick Model

#### Significant Milestones in the history of DNA Sequencing



Here, in 1977, we can see that Sanger's method arose. And worth mentioning here is of Hood's name which gave us the automated sequencer that also changed the pace of technologies in this

sequencing business.

#### Automated sequencing:

Leroy E. Hood's laboratory at the California Institute of Technology invented the first **semi-automated DNA sequencer**in 1986, which is the key technology used in **Human Genome Project.** 



Leroy E. Hood Institute of system biology Seatle Washington Conclusions:

We conclude the following:

- Genome sequencing involves recognition and determining the precise order of nucleotides in a Genome.
- Advances in sequencing technologies have revolutionized the pace of scientific discover

## **Topic # 27 Advanced Computing Approaches**



#### MUHAMMAD IMRAN 62

52





		Second base										
		U	С	Α	G							
	U	UUU UUC UUA UUG Leu	$\left. \begin{matrix} UCU \\ UCC \\ UCA \\ UCG \end{matrix} \right\} Ser$	UAU UAC Tyr UAA Stop UAG Stop	UGU UGC UGA Stop UGG Trp	U C A G						
(5' terminus)	С	CUU CUC CUA CUG	CCU CCC CCA CCG	$ \begin{array}{c} CAU \\ CAC \\ CAC \\ CAA \\ CAG \\ \end{array} \right\} Gln $	CGU CGC CGA CGG	D V D C (3' terminus)						
First base	А	AUU AUC AUA AUG Met	$\left. \begin{array}{c} ACU \\ ACC \\ ACA \\ ACG \end{array} \right\} Thr$	$ \begin{array}{c} AAU \\ AAC \\ AAC \\ AAA \\ AAG \\ Lys \end{array} $	AGU AGC AGA AGA AGG	D C A C Third base						
	G	GUU GUC GUA GUG	GCU GCC GCA GCG	GAU GAC GAA GAA GAG Glu	GGU GGC GGA GGG	U C A G						

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# **Topic # 30 31 Dynamic Programming**



3+4	Ļ,	
R <sub>3,4</sub>	=	w
R <sub>3,4</sub>	=	L
R <sub>n.n</sub>	n	

0       1       2       3       4       5       6       7       8       9       1         0       W       W       I												
0       W		0	1	2	3	4	5	6	7	8	9	10
1       W	0		W									
2     3       3     4       4     4       5     4       6     4       7     4       8     4	1	W	W									
3     4       4     4       5     4       6     4       7     4       8     4	2											
4     4 <th>3</th> <th></th>	3											
5     6       7       8	4											
6     7       8     1	5											
7       8	6											
8	7											
	8											
9	9											

Table "**R**"

10

Topic #31 Dynamic Programming Block Game (*i*-1,*j*), (*i*-1,*j*-1), (*i*,*j*-1)  $R_{0,1} R_{1,0} R_{1,1}$ 

(1,0), (1,0), 1,1)  $(2,0) \rightarrow (1,0) = W$   $(0,2) \rightarrow (0,1) = W$   $(2,1) \rightarrow |$  (1,1), (2,0), (1,0) (2,0) = L (1,2) = L  $R_{2,2} (2,2) \rightarrow$ (2,1), (1,2), (1,1)=W

	0	1	2	3	4	5	6	7	8	9	10	
0	L	W	L	W	L	W	L	W	L	W	L	
1	W	W	W	W	W	W	W	W	W	W	W	
2	L	W	L	W	L	W	L	W	L	W	L	
3	W	W	W	W	W	W	W	W	W	W	W	
4	L	W	L	W	L	W	L	W	L	W	L	
5	W	W	W	W	W	W	W	W	W	W	W	
6	L	W	L	W	L	W	L	W	L	W	L	
7	W	W	W	W	W	W	W	W	W	W	W	
8	L	W	L	W	L	W	L	W	L	W	L	
9	W	W	W	W	W	W	W	W	W	W	W	
10	L	W	L	W	L	W	L	W	L	W	L	

BLOCKS (n, m)
1. $R_{0,0} = L$
2. for i ← 1 to n
3. <b>if</b> $R_{i-1,0} = W$
4. R <sub>i,0</sub> <b>←</b> ⊥
5. else
5. R <sub>i,0</sub> ← W
7. <b>for</b> j1tom
B. <b>if</b> $R_{0,j-1} = W$
9. R <sub>o,i</sub> 🕳 L
10. else
11. R <sub>0,i</sub> ← W
12. for i – 1 to n
13. for j 🛶 1 to m
14. <b>if</b> $R_{i-1/j-1} = W$ and $R_{i,j-1} = W$ and $R_{i-1,j} = W$
15. <u>Rij</u> 🛶 L
16. else
17. <u>Rij</u> ← W
18. return R <sub>n.m.</sub>

#### FASTBLOCK(n, m)

- 1. if *n* and *m* are both even
- 2. return L
- 3. else
- 4. return W

### **Topic # 32 Algorithm for Dynamic Programming**

**Block Game Algorithm** 

BLOCKS 
$$(n, m)$$
  
1.  $R_{0,0} = L$   
2. for  $i \leftarrow 1$  to n  
3. if  $R_{i-1,0} = W$   
4.  $R_{i,0} \leftarrow L$   
5. else  
6.  $R_{i,0} \leftarrow W$   
7. for j 1 to m  
8. if  $R_{0,j-1} = W$   
9.  $R_{0,j} \leftarrow L$   
10. else  
11.  $R_{0,j} \leftarrow W$ 

i = 2 $R_{2-1,0} = R_{1,0} = W$ j = 2 $R_{0,2\text{-}1} = R_{0,1} = W$ 

BLOCKS (n, m) 12. for *i* ← 1 to n for j ← 1 to m 13. **14.** if  $R_{i-1,j-1} = W$  and  $R_{i,j-1} = W$  and  $R_{i-1,j} = W$ 15. <u>R</u>i,i ← L 16. else <u>R<sub>i,i</sub></u> ← W 17. return R<sub>n,m</sub> 18.

 $R_{1,1} = W$  $R_{2,1} = W$  $R_{1,2} = W$ 

#### FASTBLOCK(*n*, *m*)

1. if n and m are both even

- 2. return L

else
 return W

Rn,m

 $R_{2,2} = L$ 

 $R_{4,4} = L$ 

 $R_{4,5} = W$ 

# **Topic #33 Restriction Mapping**

EcoRI (Escherichia coli)

GAATTC

ATGTTTGCATTACGATAGAATTCCGTCAAAGTGCTAG TACAAACGTAATGCTATCTTAAGGCAGTTTCACGATC GCCGTTATACGCTGGATTTAAATTGCTGTGAAATGGT CGGCAATATGCGACCTAAATTTAACGACACTTTACCA TACTGCCAAGACCGAATTCCTGCGAGTGCTGAAACG ATGACGGTTCTGGCTTAAGGACGCTCACGACTTTGC GCGATATTACGAATGTGCTTACAGCACCGAATTCATC CGCTATAAAGCTTACACGAATGTCGTGGCTTAAGTAG

ATGTTTGCATTACGATAGAATTCCGTCAAAGTGCTAG TACAAACGTAATGCTATCTTAAGGCAGTTTCACGATC GCCGTTATACGCTGGATTTAAATTGCTGTGAAATGGT CGGCAATATGCGACCTAAATTTAACGACACTTTACCA TACTGCCAAGACCGAATTCCTGCGAGTGCTGAAACG ATGACGGTTCTGGCTTAAGGACGCTCACGACTTTGC GCGATATTACGAATGTGCTTACAGCACCGAATTCATC CGCTATAAAGCTTACACGAATGTCGTGGCTTAAGTAG





{2, 2, 2, 3, 3, 4, 5} If  $X = \{x_1 = 0, x_2, \dots, x_n\}$   $\Delta X = \{x_j - x_i : 1 \le i < j \le n\}$   $X = \{0, 2, 4, 7, 10\}$ , then  $\Delta X = \{2, 2, 3, 3, 4, 5, 6, 7, 8, 10\}$ , Representation of  $\Delta X$ 

	0	2	4	7	10	
0		2	4	7	10	
2			2	5	8	
4				3	6	
7					3	
10						

The element at (i, j) in the table is the value  $x_j - x_i$  for  $1 \le i < j \le n$ .

# **Topic # 34 Restriction Mapping**

{2, 2, 2, 3, 3, 4, 5} If  $X = \{x_1 = 0, x_2, \dots, x_n\}$   $\Delta X = \{xj - xi : 1 \le i < j \le n\}$  $X = \{0, 2, 4, 7, 10\}$ , then  $\Delta X = \{2, 2, 3, 3, 4, 5, 6, 7, 8, 10\}$ ,

Representation of  $\Delta X$ 



The element at (i, j) in the table is the value  $x_j - x_i$  for  $1 \le i \le j \le n$ .

# **Topic # 35 Partial Digest Problem**

#### **Partial Digest Problem:**

Given all pairwise distances between points on a line, reconstruct the positions of those

points. Input: The multiset of pairwise distances L, containing (n) integers.

Output: A set X, of n integers, such that  $\Delta X =$ 

 $\Delta A$  is equal to  $\Delta(A \oplus \{v\})$ , where  $A \oplus \{v\}$  is defined to be  $\{a + v : a \in A\}$ , Also  $\Delta A = \Delta(-A)$ , where  $-A = \{-a : a \in A\}$  $A = \{0, 2, 4, 7, 10\}, \Delta(A \oplus \{100\}) =$  $\{100, 102, 104,$  $107, 110\}, and <math>-A$  $= \{-10, -7, -4, -2,$  $0\}$ 

 $\{0, 1, 3, 8, 9, 11, 12, 13, 15\}$  and  $\{0, 1, 3, 4, 5, 7, 12, 13, 15\}$ 

	0	1	3	4	5	7	12	13	15
0		1	3	4	5	7	12	13	15
1			2	3	4	6	11	12	14

	0	1	3	8	9	11	12	13	15
0		1	3	8	9	11	12	13	15
1			2	7	8	10	11	12	14
3				5	6	8	9	10	12
8					1	3	4	5	7
9						2	3	4	6
11							1	2	4
12								1	3
13									2
15									

3		1	2	4	9	10	12
4			1	3	8	9	11
5				2	7	8	10
7					5	6	8
12						1	3
13							2
15							

 $\{14, 24, 34, 43, 52, 62, 72, 83, 92, 102, 112, 123, 13, 14, 15\}$ 

 $U \bigoplus V = \{u + v : u \in U, v \in V\}$ 

 $U \ominus V = \{\mathbf{u} - \mathbf{v} : \mathbf{u} \in \mathbf{U}, \mathbf{v} \in \mathbf{V}\}$ 

1	$U = \{6, 7, 9\}$ and $V = \{-6, 2, 6\}$							
	U⊕V	-6	2	6				
					$\mathcal{N}$	6		
	6	0	8	12				
						7		
	7	1	9	13				
	0	2	11	1.7		9		
	9	3	11	15				

UΘV	-6	2	6
6	12	4	0
7	13	5	1
9	15	7	3

#### BRUTEFORCEPDP(L, n)

- 1. M maximum element in L
- 2. for every set of n 2 integers  $0 < x_2 < \cdots < x_{n-1} < M$
- 3.  $X = \{0, x_2, \ldots, x_{n-1}, M\}$
- 4. Form  $\Delta X$  from *X*
- 5. **if** X = L
- 6. return X
- 7. **output** "No Solution"

#### ANOTHERBRUTEFORCEPDP(L, n)

- 1. M maximum element in L
- 2. for every set of n-2 integers  $0 < x_2 < \cdots < x_{n-1} < M$  from L
- 3.  $X \{0, x_2, \ldots, x_{n-1}, M\}$
- 4. Form  $\Delta X$  from X
- 5. if X = L
- 6. **return** *X*
- 7. output "No Solution"

### **Topic # 36 Practical Restriction Mapping Algorithm**

#### **Brute Force Algorithm**

Largest distance in "L"

Outermost points of "X"

Remaining distance "δ"

 $L = \{2, 2, 3, 3, 4, 5, 6, 7, 8, 10\}$ 

Size of *L* is  $\binom{n_2}{n_2} = n_{\frac{n_2-1}{2}} = 10$  where is "*n*" is number of points in the solutions.

Here "*n*" is 5 and positions of '*X*' as  $x_1 = 0$ ,  $x_2$ ,  $x_3$ ,  $x_4$  and  $x_5$ .

<i>x</i> <sub>1</sub> =0	X2	X3	X4	Xs
-				

#### **Practical Restriction Mapping Algorithm**

$\binom{n}{n}$ $\binom{n}{n}$ $\frac{n!}{n!}$ =	n!
$\binom{k}{k} = \binom{n}{2} = \binom{n-k}{k!}$	(n-2)!2!
$=\frac{n(n-1)}{(n-1)}$	(n-2)!
(1-)	2):2:
$=\frac{n(n)}{n}$	
	2
$\frac{n(n-1)}{2}$	$\frac{1}{2} = 10$
n <sup>2</sup> -n = 2	x 10
n²-n =	20
n²-5n + 4n	-20 = 0
n (n-5) + 4	(n-5) = 0
(n-5) (n+	4) = 0
n= 5; n	i= -4
$\binom{5}{2}$	$\binom{-4}{2}$

X1=0	X2	Xa	X <sub>4</sub>	X5=10
-				

 $L = \{2, 2, 3, 3, 4, 5, 6, 7, 8, 10\}$ 

X	$f = \{0, 10\}$	$L = \{2, 2, 3, 3, 4, 5, 6, 7, 8\}$				
<b>X</b> 1=0	X <sub>2</sub> =2	X <sub>3</sub>	X <sub>4</sub> =8	X <sub>5</sub> =10		

 $x_5 - x_2 = 8$  and  $x_2 - x_1 = 2$ 

 $X = \{0, 2, 10\} \qquad L = \{2, 3, 3, 4, 5, 6, 7\}$
$X = \{0, 2, 10\}$   $L = \{2, 3, 3, 4, ...\}$ 5, 6, 7  $x_4 = 7$  or  $x_3 = 3$ If  $x_3 = 3$  then  $x_3 - x_2 = 1$  must be in *L*, but not, so  $x_4$  must be 7,  $x_5 - x_4 = 3$ ,  $x_4 - x_2 = 3$ 5, and  $x_4 - x_1 = 7$  from *L*  $X = \{0, 2, 7, 10\}$  $L = \{2, 3, 4, 6, \}$ X<sub>3</sub>= 4 or 6 X5=10  $X_1 = 0$ X2=2 X4=7 Two choices,  $x_3 = 4$  or  $x_3 = 6$ , If  $x_3 = 6$ , then  $x_4 - x_3 = 1$  must be in *L* but not. Then  $x_3 = 4$  $X = \{0, 2, 4, 7, 10\}$ X2=2 X3=4 *x*<sub>1</sub>=0 X4=7 X5=10

### **Topic #37** Partial Digest Algorithm

Brute Force Algorithm

List of pairwise distances, L, and uses the function DELETE(y, L)

Value *y* from *L*, notation (y, X) to denote the multiset of distances

For example,

 $\Delta(2,\{1,3,4,5\}) = \{1,1,2,3\}$ 

#### PARTIALDIGEST (L)

- 1. *width* Maximum element in L
- 2. DELETE (width, L)
- 3.  $X = \{0, width\}$
- 4. PLACE (L,X)

#### PLACE (L,X)

- 1. if L is empty
- 2. output X
- return
   y
  - y Maximum element in L
- 5. if  $\Delta(y,X) \subseteq L$
- 6. 6 Add y to X and remove lengths  $\Delta$  (y,X) from L
- 7. PLACE (*L*,*X*)
- 8. Remove *y* from *X* and add lengths (y, X) to *L*
- 9. if  $(width y, X) \subseteq L$
- 10. Add width -y to X and remove lengths (width -y,X) from L
- 11. PLACE (*L*,*X*)
- 12. Remove width -y from X and add lengths (width -y,X) to L
- 13. return

## **Topic # 38 Partial Digest Algorithm**

PARTIALDIGEST (L)

10
10, L
0, 10

 $X = \{0, 10\}$ 

PLACE (L,X) 1. if L is empty  $L = \{2, 2, 3, 3, 4, 5, 6, 7, 8, 10\}$ 2. output X  $L = \{2, 2, 3, 3, 4, 5, 6, 7, 8\}$ 3. return  $X = \{0, 10\}$ 4. y - Maximum element in L 8  $\Delta(y,X) = 8(0, 10) = (8, 10)$ 5. if  $\Delta(y, X) \subseteq L$ Add y to X and remove 6 lengths  $\Delta(y, X)$  from L PLACE (L,X) 7. 8. Remove y from X and add lengths (y,X) to L  $L = \{2, 2, 3, 3, 4, 5, 6, 7, 8\}$  $X = \{0, 10\}$ if (width -y,X)  $\subseteq L$ width - y,X = 10-8 (0,10) 9. = 2 (0, 10) 10. Add width - y to X and = (2, 8) remove lengths (width - y, X) from L PLACE (L,X) 11.  $L = \{2, 3, 3, 4, 5, 6, 7\}$ 12. Remove width - y from X and  $X = \{0, 2, 10\}$ add lengths (width - y,X) to L 13. return

1. if <i>L</i> is empty 2. <b>output</b> <i>X</i> 3. <b>return</b> 4. $y \leftarrow$ Maximum element in <i>L</i> 5. if $\Delta(y,X) \subseteq L$ 6 Add <i>y</i> to <i>X</i> and remove lengths $\Delta(y,X)$ from L 7. PLACE ( <i>L,X</i> ) 8. Remove <i>y</i> from <i>X</i> and add lengths ( <i>y,X</i> ) to <i>L</i>	$L = \{2, 3, 3, 4, 5, 6, 7\}$ $X = \{0, 2, 10\}$ $7$ $\Delta (y, X) = 7 (0, 2, 10) = (7, 5, 3)$ $L = \{2, 3, 3, 4, 5, 6,\}$ $X = \{0, 2, 7, 10\}$
PLACE ( <i>L</i> , <i>X</i> ) 1. if <i>L</i> is empty 2. <b>output</b> <i>X</i> 3. <b>return</b> 4. $y \leftarrow$ Maximum element in <i>L</i> 5. <b>if</b> $\Delta(y,X) \subseteq L$ 6 Add <i>y</i> to <i>X</i> and remove lengths $\Delta(y,X)$ from L 7. PLACE ( <i>L</i> , <i>X</i> ) 8. Remove <i>y</i> from <i>X</i> and add lengths ( <i>y</i> , <i>X</i> ) to <i>L</i>	$L = \{2, 3, 3, 4, 5, 6\}$ $X = \{0, 2, 7, 10\}$ 6 $\Delta (y, X) = 6 (0, 2, 7, 10) = (6, 4, 1, 4)$

#### PLACE (L,X)

- 1. if *L* is empty
- 2. output X
- 3. return
- y ← Maximum element in L
- 5. if  $\Delta(\underline{y},\underline{X}) \subseteq L$
- 6 Add y to X and remove lengths Δ (y,X) from L
- 7. PLACE (*L*,*X*)
- Remove y from X and add lengths (y, X) to L

 $L = \{2, 3, 3, 4, 5, 6\}$   $X = \{0, 2, 7, 10\}$  6 $\Delta (y, X) = 6 (0, 2, 7, 10) = (6, 4, 1, 4)$ 

### **Topic # 39 Regulatory Motifs in DNA Sequences**

Sequence Motifs

**Biological function** 

Nucleases and transcription factors

Processes at RNA level

Specific sequence located upstream of genes TCGGGGATTTCC

NF-kB binding sites (nuclear factor kappa-light-chain-enhancer of activated B cells)

Regulatory motifs, transcription factors

Set of upstream regions in genes in the genome, each region containing one NF-kB

sites Suppose we do not know either location or sequence of NF-kB sites

"The Gold Bug" by Edgar Allan provided some clue of finding DNA motifs, one of the character find parchment written below

```
53++!305))6^{*};4826)4+.)4+);806^{*};48!8^{\circ}60))85;]8^{*}:+*8!83(88)5^{*}!;46(;88^{*}96^{*}?;8)
*+(;485);5^{*}!2:^{*}+(;4956^{*}2(5^{*}-)8^{\circ}8^{*};4069285) ;)6!8)
4++;1(+9;48081;8:8+1;48!85;4)485!
528806^{*}81(+9;48;(88;4(+?34;48)4+;161;:188;+?;
```

```
";48" codes for "THE"
53++!305))6*THE26)H+.)H+)TE06*THE!E'60))E5T]E*:+*E!E3(EE)5*!TH6(T
EE*96*?TE
)* +(THE5) T5*!2:*+(TH956*2(5*-H)E'E*T
H0692E5)T)6!E)H++T1(+9THE0E1TE:E+1THE!E5TH)HE5!52EE06*E1(+9TH
ET(EETH(+?3HTHE)H+T161T:1EET+?T
```

";" for "T"

"4" for "H"

**"8" for "E"** 

## Topic # 40 Profiles 1

#### **Conserved Pattern**

32 nucleotide

7 sequences

- 1. CGGGGCTGGGTCGTCACATTCCCCTTTCGATA
- 2. TTTGAGGGTGCCCAATAACCAAAGCGGACAAA
- 3. GGGATGCCGTTTGACGACCTAAATCAACGGCC
- 4. AAGGCCAGGAGCGCCTTTGCTGGTTCTACCTG
- 5. AATTTTCTAAAAAGATTATAATGTCGGTCCTC
- 6. CTGCTGTACAACTGAGATCATGCTGCTTCAAC
- 7. TACATGATCTTTTGTGGATGAGGGAATGATGC

#### Figure 1

P = ATGCAACT

*l* = 8

- 1. CGGGGCT<u>ATGCAACT</u>GGGTCGTCACATTCCCCTTTCGATA
- 2. TTTGAGGGTGCCCAATAAATGCAACTCCAAAGCGGACAAA
- 3. GGATGCAACTGATGCCGTTTGACGACCTAAATCAACGGCC
- 4. AAGG<u>ATGCAACT</u>CCAGGAGCGCCTTTGCTGGTTCTACCTG
- 5. AATTTTCTAAAAAGATTATAATGTCGGTCC<u>ATGCAACT</u>TC
- 6. CTGCTGTACAACTGAGATCATGCTGCATGCAACTTTCAAC
- 7. TACATGATCTTTTG<u>ATGCAACT</u>TGGATGAGGGAATGATGC

#### Figure 2

#### P = ATGCAACT

*l* = 8

- 1. CGGGGCTATGCAACTGGGTCGTCACATTCCCCTTTCGATA
- 2. TTTGAGGGTGCCCAATAAATGCAACTCCAAAGCGGACAAA
- 3. GGATGCAACTGATGCCGTTTGACGACCTAAATCAACGGCC
- 4. AAGGATGCAACTCCAGGAGCGCCTTTGCTGGTTCTACCTG
- 5. AATTTTCTAAAAAGATTATAATGTCGGTCCATGCAACTTC
- 6. CTGCTGTACAACTGAGATCATGCTGCATGCAACTTTCAAC
- 7. TACATGATCTTTTGATGCAACTTGGATGAGGGAATGATGC

#### Figure 3

P = ATGCAACT

*l* = 8

7 x (32 + 8) = 280 nucleotides

Probability =  $280/4^8 = 0.004$ 

- 1. CGGGGCT<u>ATcCAgCT</u>GGGTCGTCACATTCCCCTTTCGATA
- 2. TTTGAGGGTGCCCAATAAggGCAACTCCAAAGCGGACAAA
- 3. GG<u>ATGgAtCT</u>GATGCCGTTTGACGACCTAAATCAACGGCC
- 4. AAGG<u>AaGCAACc</u>CCAGGAGCGCCTTTGCTGGTTCTACCTG
- 5. AATTTTCTAAAAAGATTATAATGTCGGTCCtTGgAACTTC
- 6. CTGCTGTACAACTGAGATCATGCTGC<u>ATGCcAtT</u>TTCAAC
- 7. TACATGATCTTTTG<u>ATGgcACT</u>TGGATGAGGGAATGATGC

#### Figure 4

## **Topic # 41 Profiles 2**

#### **Conserved Pattern**

	1- TCGGGGATTTCA
18 sequences	2- ACGGGGATTTTT
NF-kB	3- TCGGTACTITAC
TCGGGGGATTTCC	4- TIGGGGACTIT
/= 12	6- GCGGGGGAATIIC
	7- TCGGGGATICCT
	8- TCGGGGATTCCT
	9- TAGGGGAACTAC
	10- TCGGGTATAAAC
	11- TCGGGGGTTTTT
Alignment	12- CCGGTGACTTAC
	13- CCAGGGACTCCC
	14- AAGGGGACTICC
	15- TIGGGGACTIII
	16- TTTGGGAGTCCC
	17- TCGGTGAT <u>TI</u> CC
	18- TAGGGGAAGACC
	A: 2 3 1 0 0 1 16 3 1 2 4 1
Profile	T: 12 3 1 0 4 1 0 9 15 11 5 6
	G: 10 16181416 1 1 1 0 00 C: 3 12 0 00 0 15 1 5 911
Consensus	TCGGGGATTTCC
	score = 12, 12, 16, 18, 14, 16, 16, 9, 15, 11, 9, 11= 159

1- position 8 - Sequence 1



- 3- position 3- Sequence 3
- 4- position 5- Sequence 4
- 5- position 31- Sequence 5
- 6- position 27- Sequence 6
- 7- position 15- Sequence 7

CGGGGCT<u>ATcCAgCT</u>GGGTCGTCACATTCCCCTT

2-

1-

TTTGAGGGTGCCCAATAAggGCAACTCCAAAGCGGACAAA 3-

GG<u>ATGgAtCT</u>GATGCCGTTTGACGACCTA

4-

AAGG<u>AaGCAACc</u>CCAGGAGCGCCTTTGCTGG

- 5- AATTTTCTAAAAAGATTATAATGTCGGTCCtTGgAAC<u>T</u>TC
- 6- CTGCTGTACAACTGAGATCATGCTGC<u>ATGCcAtTT</u>TC AAC

TACATGATCTTTTG<u>ATGgcACT</u>TGGATGAGGGAATGATGC

#### Figure 6

7-

## **Topic # 42 Profiles 3**

#### **Conserved Pattern**

		1 2 3 4 5 6 7 8	
Alignment		ATCCAGCT	
		GGGCAACT	
		ATGGATCT	
Angiment		AAGCAACC	
		TIGGAACT	
		ATGCCATI	
		ATGGCACT	
	A	5 100 5 500	
Duefile	т	1 5000 116	Figure 7
Profile	G	1 1 6 3 0 1 0 0	
	С	00142061	
Consensus		ATGCAACT	
		$\sim$	

If s = (8, 19, 3, 5, 31, 27, 15) then

Score(s) = 5 + 5 + 6 + 4 + 5 + 5 + 6 + 6

= 42 Set of t DNA sequences

"n" nucleotides one position in each of

these "t" sequences  $s = (s_1, s_2, \ldots, s_t)$ ,

with  $1 \le s_i \le n - l + 1$  *l*-mers can be

compiled into  $t \times l$  alignment matrix

whose (i, j)th element is the nucleotide in the  $s_i + j - 1$ th element in the *i*th sequence of figure 7

## **Topic # 43 Motif Finding Problem**

#### **Profile Matrix**

If P(s) denotes profile matrix, starting from s

 $M_{P(s)}(j)$ -largest count in column *j* of P(s)

		ATCCAGCT	
		GGGCAACT	AA (1) E
Allanant		ATGGATCT	$IVI_{P(s)}(1) = 5$
Alignment		AAGCAACC	$M_{P(s)}(2) = 5$
		TIGGAACT	$M_{P(s)}(3) = 6$
		ATGCCATT	$M_{P(s)}(4) = 4$
		ATGGCACT	$M_{P(s)}(5) = 5$
Profile	Α	5 1005 500	$M_{P(s)}(6) = 5$
	т	1 5000 116	$M_{\rm p(a)}(7) = 6$
	G	1 1 6 3 0 1 0 0	$M_{-1}(8) = 6$
	С	00142061	101P(s)(0) = 0
Consensus		ATGCAACT	

For the starting positions in above figure, Score(*s*,*DNA*) =

5 + 5 + 6 + 4 + 5 + 5 + 6 + 6 = 42. Score(*s*,*DNA*) can be used to measure

The strength of a profile corresponding to the starting positions s. consensus score of  $l \cdot t$  corresponds to the best possible alignment *Score*(*s*,*DNA*) consensus score is defined to be *Score*(*s*,*DNA*) =  $\sum_{j=1}^{l} MP_{(s)}(j)$  Score(*s*,DNA) = 5 + 5 + 6 + 4 + 5 + 5 + 6 + 6 = 42 consensus score of  $l \cdot t$  corresponds to the best possible alignment

A consensus score of l.t/4, is worst possible alignment

Motif Finding Problem:

Given a set of DNA sequences, find a set of l-mers, one from each sequence, that maximizes the consensus score.

Input: A  $t \times n$  matrix of DNA, and, the length of the pattern to

find. Output: An array of t starting positions  $s = l(s_1, s_2, ..., s_n)$ 

s<sub>t</sub>) maximizing Score(*s*,*DNA*).

## **Topic # 44 Motif Finding Problem 1**

#### **Median String**

Median string

Given two *l*-mers *v* and *w*, can compute the *Hamming distance* between them,  $d_H(v,w)$ , as the number of positions that differ in the two strings

ATTGTC

### : x : x : :

ACTCTC

 $\mathbf{s} = (\mathbf{s}_1, \mathbf{s}_1, \ldots, \mathbf{s}_t)$ 

*v* is some *l*-mer  $d_{\rm H}(v, s)$  to denote the total Hamming distance between *v* and the *l*-mers starting at positions s:  $d_{\rm H}$  $(v, s) = \sum_{i=1}^{t} d_{\rm H}(v, si)$  where  $d_{\rm H}(v, s_i)$  is the Hamming distance between *v* and the *l*-mer that starts at  $s_i$  in the *i*th DNA sequence TotalDistance(*v*,*DNA*) = min<sub>s</sub>( $d_{\rm H}(v, s)$ )

Finding Total Distance(*v*,*DNA*) is a simple problem:

find the best match for *v* in the first DNA sequence (i.e., a position minimizing  $d_{\rm H}(v, s_1)$  for  $1 \le s_1 \le n-l+1$ ), then the best match in the second sequence and so on

Median string for DNA as the string v that minimizes TotalDistance(v,DNA); this minimization is performed over all 4<sup>1</sup> strings v of length *l*.

#### **Median String Problem:**

Given a set of DNA sequences, find a median

*string.* **Input**: A  $t \times n$  matrix DNA, and l,

the length of the pattern to find

Output: A string v of *l* nucleotides that

minimizes TotalDistance(v,DNA)

over all strings of that length

**Double minimization**: finding a string v that minimizes TotalDistance(v, DNA), which is in turn the smallest distance among all choices of starting points s in the DNA sequences.

Min min

d<sub>H</sub>(v, s) all choices of

all choices *l*-mers v

starting positions s

The Median String problem- Minimization problem

The Motif Finding problem- Maximization problem

Computationally equal

Let *s* be a set of starting positions with consensus score Score(s, DNA), and let *w* be the consensus string of the corresponding profile. Then  $d_H(w, s) = lt - Score(s, DNA)$ 

AT CC AGCT
 G GGC AACT
 AT GGATCT
 AAGC AACC
 TI GG AACT
 ATG C CATI
 ATGC CACT
 ATGC AACT

Hamming distance between the consensus

string w and each of the seven implanted patterns is 2, and  $d_{\rm H}(w, s) = 2 \ge 7 = 14$ 

 $= 7 \times 8 - 42 = 14$ 

## **Topic # 45 Motif Finding Problem 2**

#### Median String

Median string

Given two *l*-mers *v* and *w*, can compute the *Hamming distance* between them,  $d_H(v,w)$ , as the number of positions that differ in the two strings

#### A T T G T C

: x : x : :

ACTCTC

 $\mathbf{s} = (\mathbf{s}_1, \mathbf{s}_1, \dots, \mathbf{s}_t) \ v$  is some *l*-mer  $d_H(v, s)$  to denote the total Hamming distance between *v* and the *l*-mers starting at positions s:  $d_H(v, s) = \sum_{i=1}^t d_H(v, si)$  where  $d_H(v, s_i)$  is the Hamming distance between *v* and the *l*-mer that starts at  $s_i$  in the *i*th DNA sequence

TotalDistance(v,DNA) = min<sub>s</sub>( $d_{\rm H}(v, s)$ 

Finding Total Distance(*v*,*DNA*) is a simple problem:

find the best match for v in the first DNA sequence (i.e., a position minimizing  $d_{\rm H}(v, s_1)$  for 1

 $\leq$  s<sub>1</sub>  $\leq$  n-*l*+1), then the best match in the second sequence and so on

Median string for DNA as the string v that minimizes TotalDistance(v,DNA); this minimization is performed over all 4<sup>1</sup> strings v of length *l*.

#### **Median String Problem:**

Given a set of DNA sequences, find a median

*string.* **Input**: A  $t \times n$  matrix DNA, and l,

the length of the pattern to find

**Output**: A string v of *l* nucleotides that

minimizes TotalDistance(v,DNA)

over all strings of that length

**Double minimization**: finding a string v that minimizes TotalDistance(v,DNA), which is in turn the smallest distance among all choices of starting points s in the DNA sequences.

 $d_{\rm H}(v, s)$ 

Min min

all choices of

es of all

choices *l*-mers

starting positions s

The Median String problem- Minimization problem

The Motif Finding problem- Maximization problem

Computationally equal

Let *s* be a set of starting positions with consensus score Score(*s*,*DNA*), and let *w* be the consensus string of the corresponding profile. Then  $d_H(w, s) = lt - Score(s, DNA)$ 

1. AT CC AGCT 2. GGGC AACT 3. AT GGATCT 4. AAGC AACC 5. TI GG AACT 6. ATG C CATI 7. ATGG CACT ATGC AACT

Hamming distance between the consensus

 $= 7 \times 8 - 42 = 14$ 

## **Topic #45 Search Trees-Introduction**

#### Introduction



BBB, BBG, BGB, BGG, GBB, GBG, GGB, GGG  $2^3 = 8$ 

### **Topic # 46 Search Trees Best Alternative**

Median String and

Motif Finding Problems

Number of alternatives

To find best one

 $(n - l + 1)^t$ 

#### **Search Trees**

(	1,	1, ,	1,	1)
(	1,	1,,	1,	2)
(	1,	1,,	1,	3)
(	1,	1,,	1,	n – l + 1)
(	1,	1,,	2,	1)
(	1,	1,,	2,	2)
(	1,	1,,	2,	3)
(	1,	1,,	2,	n – l + 1)
(n -	-l+1,	n – l + 1, ,		n – l + 1, 1)
(n -	-l+1,	n – l + 1,,	n – l + 1,	2)
(n -	- +1,	n – l + 1, ,	, n – l + 1,	3)
(n -	-l+1,	n – l + 1,,	n – l + 1,	n – l + 1)

Figure 1

#### Figure 1

8

6

3



Figure 3



1	for A	1
2	for <sup>¬</sup>	Γ
3	for (	3
4	for (	2

(1,	1,	•	•	•	,	1,	1)
(1,	1,	•	•	•	,	1,	2)
(1,	1,	•	•	•	,	1,	3)
(1,	1,	•	•	•	,	1,	4)
(1,	1,	•	•	•	,	2,	1)
(1,	1,	•	•	•	,	2,	2)
(1,	1,				,	2,	3)

(1, 1, ..., 2, ..., 2, ..., 2, ..., 2, ..., 3, 3)(4, 4, ..., 3, 3)(4, 4, ..., 3, 4)(4, 4, ..., 4, 1)(4, 4, ..., 4, 2)(4, 4, ..., 4, 3)(4, 4, ..., 4, 4)1 for A

2 for T

3 for G

4 for C

Consider all  $k^L L$ -mers in a *k*-letter alphabet

For Motif Finding problem, k =

n-l+1, For Median String

problem, k = 4.

All  $2^4$  4-mers in the two-letter alphabet of 1 and 2.

### **Topic # 47 Algorithm for Search Trees 1**

6.

#### Search tree Algorithms

Next leave All leaves Preorder

*L*-mer  $a = (a_1 a_2...a_L)$ 

```
NEXTLEAF(\underline{a}, \underline{L}, k)

1. for \underline{i} \leftarrow L to 1

2. if \underline{a}_{\underline{i}} < k

3. \underline{a}_{\underline{i}} \leftarrow \underline{a}_{\underline{i}} + 1

4. return a

5. \underline{a}_{\underline{i}} \leftarrow 1

6. return a
```

ALLLEAVES(L, k)

a = elements of L-mer L = l-mer k = number of elements

```
    a ← (1, ..., 1)
    while forever
    output a
    a ← NEXTLEAF(a,L, k)
    if a = (1, 1, ..., 1)
```

return



## **Topic # 48 Algorithm for Search Trees 2**

#### PREORDER(v)

- 1. output *v*
- 2. if *v* has children
- 3. **PREORDER**(left child of v)
- 4. **PREORDER**( right child of v )



1. (-,-,-,-)	
2. (1,-,-,-)	
3. (1,1,-,-)	
4. (1,1,1,-)	
5. (1,1,1,1)	
6. (1,1,1,2)	
7. (1,1,2,-)	
8. (1,1,2,1)	
9. (1,1,2,2)	
10. (1,2,-,-)	
11. (1,2,1,-)	
12. (1,2,1,1)	
13. (1,2,1,2)	
14. (1,2,2,-)	
15. (1,2,2,1)	
16. (1,2,2,2)	
17. (2,-,-,-)	
18. (2,1,-,-)	
19. (2,1,1,-)	
20. (2,1,1,1)	
21. (2,1,1,2)	
22. (2,1,2,-)	
23. (2,1,2,1)	
24. (2,1,2,2)	
25. (2,2,-,-)	
26. (2,2,1,-)	
27. (2,2,1,1)	
28. (2,2,1,2)	
29. (2,2,2,-)	
30. (2,2,2,1)	
31. (2,2,2,2)	

# **Topic # 49 Next Vertex Algorithm**

**Traversing vertices** 

Input:  $a = (a_1, ..., a_L)$  at

level *i* Output: Next vertex in

the tree values  $a_1, \ldots, a_i$  and

ignores  $a_{i+1}, \ldots, a_L$ 

NEXTVERTEX takes inputs that are similar to NEXTLEAF, with the exception that the

"current leaf" is now the "current vertex," and uses

parameter *i* for vertices



When i < L, NEXTVERTEX (**a**, *i*, *L*, *k*) moves down to the next lower level and explores that subtree of **a**. If i = L, NEXTVERTEX either moves along the lowest level as long as  $a_L < k$  or jumps back up in the tree.

### **Topic # 50 Advanced Computing Approaches**

#### Algorithm

- Some entity needs to carry out the steps specified by the algorithm
- Humans are generally slow

- A computer is less intelligent but can perform simple steps quickly and  $\geq$ reliably
- > Algorithm must be rephrased in programming language
- Pseudocode: language often used to describe algorithm
- > Complex operations are grouped together into mini-algorithms called subgroups
- **Variable** is written as *x* or *total*  $\geq$
- > An **array** of *n* elements is an ordered collection of *n* variables  $a_1$ , a<sub>2</sub>,....a<sub>n</sub>
- An algorithm is a pseudocode is denoted by a name, followed by the list of arguments

## **Topic #51 ByPass Algorithm**

#### **Branch and Bound Algorithm**

NEXTVERTEX Algorithm **BYPASS** Algorithm

Skip the subtree rooted at

vertex (a, i) Increment  $a_i$ 

(unless  $a_i = k$ 



A tree that has uninteresting subtrees. The numbers next to a leaf represent the "score" for that L-mer. Scores at internal vertices represent the maximum score in the subtree rooted at that vertex. To improve the brute force algorithm, we can "prune" subtrees that do not contain highscoring leaves. For example, since the score of the very first leaf is 24, it does not make sense to analyze the 4th, 5th, or 6th leaves whose scores are 20, 4, and 5, respectively. Therefore, the subtree containing these vertices can be ignored.

BYPASS(a, *i*, *L*, *k*)  
1. for 
$$j \leftarrow i$$
 to 1  
2. if  $a_i < k$   
3.  $a_j \leftarrow a_i + 1$   
4. return (a, *j*)  
5. return (a, 0)

# **Topic # 52 Finding Motifs**

#### Brute force approach

#### BRUTEFORCEMOTIFSEARCH(DNA, t, n, l)

0

1. bestScore

5

- 2. for each  $(s_1, \ldots, s_t)$  from  $(1, \ldots, 1)$  to  $(n l + 1, \ldots, n l + 1)$
- 3. if *Score*(*s*,*DNA*) > *bestScore*
- 4. *bestScore Score*(*s*,*DNA*)
- bestMotif  $(s_1, s_2, \ldots, s_t)$
- 6. return bestMotif n-l+1 choices

for the first index  $(s_1)$ , then for  $s_2, s_3$ )

number of positions is  $(n - l + 1)^t$ 

Score(*s*, *DNA*), which requires O(l) operations- $O(ln^t)$ .

#### BRUTEFORCEMOTIFSEARCHAGAIN(DNA, t, n, l)

- 1. s  $(1, 1, \ldots, 1)$
- 2. *bestScore Score*(s,*DNA*)
- 3. while forever
- 4. s NEXTLEAF(s, t, n l + 1)
- 5. if *Score*(*s*,*DNA*) > *bestScore*
- 6. *bestScore* Score(s,DNA)
- 7. bestMotif  $(s_1, s_2, \ldots, s_t)$
- 8. if s = (1, 1, ..., 1)
- 9. return bestMotif

### **Topic # 53 Simple Motif Search Algorithm**

Simple Motif Search Algorithm

SIMPLEMOTIFSEARCH(DNA, t, n, l)

1.	s (1,,1)
2.	bestScore 0
3.	<i>i</i> 1
4.	while $i > 0$
5.	if <i>i</i> < t
б.	(s, i) NEXTVERTEX(s, i, t, $n - l + 1$ )
7.	else
8.	if Score(s,DNA) > bestScore
9.	bestScore Score(s,DNA)
10.	bestMotif $(s_1, s_2, \ldots, s_t)$
11.	(s, <i>i</i> ) NEXTVERTEX(s, <i>i</i> , <i>t</i> , $n - l + 1$ )
12.	return bestMotif

#### Simple Motif Search Algorithm

Some sets of starting positions can be ruled out

If the first *i* of *t* starting positions [i.e.,  $(s_1, s_2, ..., s_i)$ ]

Sequences  $i+1, i+2, \ldots, t$ ,

s = ( $s_1$ ,  $s_2$ ,...,  $s_t$ ), define the partial consensus score, Score( $s_i$ ,  $i_i$ , DNA)- $i \times l$  alignment matrix

Partial consensus score for  $s_1, \ldots, s_i$ , remaining t-i rows can only improve the consensus score by  $(t - i) \cdot 1$ 

First *i* starting positions  $(s_1, \ldots, s_1)$  could be at most Score $(s, i, DNA) + (t-i) \cdot l$ 

Score(s, i,DNA)+  $(t-i) \cdot 1$ 

is less than the currently best score,

bestScore t - i sequences in the

sample

 $Score(s, i, DNA) + (t - i) \cdot l$ 

$$(n-l+1)^{t-i}$$

### **Topic # 54 Branch and Bound Algorithm**

#### **Branch and Bound**

**Motif Search** 

#### BRANCHANDBOUNDMOTIFSEARCH(DNA, t, n, l)

1. s (1	$1, \ldots, 1$	)
---------	----------------	---

0

- 3. *i* 1
- 4. while i > 0
- 5. if i < t
- 6. *optimisticScore* Score (s, i, DNA)+

 $(t-i) \cdot l$ 

if optimisticScore < bestScore

8. (s, i) BYPASS(s, i, t, n - l + 1)

9. else

7.

10. (s, i) NEXTVERTEX(s, i, t, n - l + 1)

11. else

12. if *Score*(*s*,*DNA*) > *bestScore* 

13.bestScoreScore(s)

14. bestMotif  $(s_1, s_2, \ldots, s_t)$ 

15. 
$$(s, i)$$
 NEXTVERTEX $(s, i, t, n - l + 1)$ 

16. return bestMotif

### **Topic # 55 Brute Force Algorithm**

#### **Brute Force**

#### Median Search

#### BRUTEFORCEMEDIANSEARCH(DNA, t, n, l)

- 1. bestWord AAA···AA
- 2. *bestDistance*
- 3. for each *l*-mer *word* from AAA...A to TTT...T
- 4. if TOTALDISTANCE(word,DNA) < bestDistance
- 5. *bestDistance* TOTALDISTANCE(word,DNA)
- 6. *bestWord* word
- 7. return *bestWord*



A search tree for the Median String problem. Each branching point can give rise to only four children, as opposed to the n-l+1 children in the Motif Finding problem.

#### SIMPLEMEDIANSEARCH(DNA, t, n, l)

1. s  $(1, 1, \dots, 1)$ 

2. *bestDistance* 

3.	i 1
4.	while i > 0
5.	if i < 1
6.	(s, i) NEXTVERTEX $(s, i, l, 4)$
7.	else
8	<i>word</i> nucleotide string corresponding to $(s_1, s_2, \ldots s_l)$
9.	if TOTALDISTANCE(word,DNA) < bestDistance
10.	bestDistance TOTALDISTANCE(word,DNA)
11.	bestWord word
12.	(s, i) NEXTVERTEX $(s, i, l, 4)$
13.	return bestWord
BRANC	HANDBOUNDMEDIANSEARCH(DNA, t, n, l)
1.	$s (1, 1, \ldots, 1)$
2.	bestDistance
3.	i 1
4.	while i > 0
5.	if i <1
6.	<i>prefix</i> nucleotide string corresponding to $(s_1, $
	$s_2,\ldots,s_i)$
7.	optimisticDistance TOTALDISTANCE(prefix,DNA)
8.	if optimisticDistance > bestDistance
9.	(s, i) BYPASS $(s, i, l, 4)$
10.	else
11.	(s, i) NEXTVERTEX $(s, i, l, 4)$
12.	else
13	word nucleotide string corresponding to
	$(s_1, s_2, \ldots s_l)$
14.	if TOTALDISTANCE(word,DNA) < bestDistance
15.	bestDistance TOTALDISTANCE(word,DNA)
16.	bestWord word
17.	(s, i) NEXTVERTEX $(s, i, l, 4)$
18.	return bestWord
Top	ic # 56 Genomic Rearrangements

#### Waardenburg's syndrome

- ➤ Hearing loss
- Two Different colored eyes
- Gene present on chromosome 2

- Splotch gene in mice
- Human genome-mouse genome
- Cut into 300 genomic fragments-Synteny blocks
- Chromosome 2 in humans-mouse chromosomes 1, 2, 3, 5, 6, 7, 10, 11, 12, 14, and 17
- Genome rearrangement results in a change of gene ordering
- > Analysis of human and mouse genomes-250 genomic rearrangements

#### Mouse X chromosome



#### Human X chromosome

Transformation of the mouse gene order into the human gene order on the X chromosome

### **Topic # 57 Greedy Approach for Motif Search**

```
Approximation algorithm
```

Brute force algorithm to solve the Motif Finding

problem running time of  $O(l \cdot n^{t})$ 

Cannot run it on

biological samples

Faster greedy

technique-not correct,

good performance

Approximation

algorithm

CONSENSUS- as good

#### or better

GREEDYMOTIFSEARCH scans each DNA sequence only once. Once we have scanned a particular sequence i, we decide which of its l-mer has the

best contribution to the partial alignment score Score(s, i, DNA) for the first i sequences and immediately claim that this l-mer is part of the alignment.

#### GREEDYMOTIFSEARCH (DNA, t, n, l)

1. bestMotif (1	, 1,	• •		,	1)
-----------------	------	-----	--	---	----

- 2. s  $(1, 1, \ldots, 1)$
- 3 for  $s_1$  1 to n l + 1
- 4 for  $s_2$  1 to n l + 1

5. if Score(s, 2,DNA) > Score(bestMotif,

2,DNA)

 $6. \qquad BestMotif1 \quad s_1$ 

7. BestMotif2 s<sub>2</sub>

8.  $s_1$  *BestMotif*<sub>1</sub>

9.  $s_2$  BestMotif<sub>2</sub>

10. for i 3 to t

11 for  $s_i$  1 to n - l + 1

12. if Score(s, *i*, *DNA*) > Score (*bestMotif*,

i, DNA)

13.  $BestMotif_i$  s<sub>i</sub>

14.  $s_i$  bestMotifi

15. return bestMotif

#### Approximation algorithm

Two

closest l-

mers  $2 \times l$ 

seed

matrix l(n

 $(-l+1)^2$ 

operations

t – 2 iterations-by scanning the *i*th sequence (for  $3 \le i \le t$ )

t-2 sequences and selecting the one *l*-mer that has

themaximumScore(s, i)  $l \cdot (n - l + 1)$  operations

#### Approximation algorithm

Running time of this algorithm is  $O(ln^2 + lnt)$ , which is vastly better than the O(lnt) of

IMPLEMOTIFSEARCH or even the O(4lnt) of BRUTEFORCEMEDIAN-

STRING

### **Topic # 58 The Power of DNA Sequence Comparison**

#### Introduction

Discovery of new gene-no idea of functions

Find similarities with genes of known function

Newly discovered cancer-causing -sis oncogene matched a normal gene involved in growth and development called platelet-derived growth factor

Oncogene v-sis is the simian sarcoma virus

Scientists became suspicious that cancer might be caused by a normal growth gene

Discovery of cystic fibrosis gene

Abnormal secretions, and is diagnosed in children produce abnormally thick mucus that clogs the lungs

10 million Americans are carriers of the cystic fibrosis gene

There is a 25% chance that the child will have cystic fibrosis

In 1989, 1 million nucleotides on the chromosome 7

The area around the cystic fibrosis gene was sequenced

Database of all known genes

Similarities between a gene that had already been discovered, code for *adenosine triphosphate* (*ATP*) *binding proteins* 

These proteins constitute the ion transport channel

The disease involves sweat secretions with abnormally high sodium content

Link between cancer-causing genes and normal growth genes and elucidating the nature of cystic fibrosis

### **Topic # 59 Brute Force vs Greedy Algorithm**

Dynamic Programming

Manhattan Tourist Problem:

Find a longest path in a

weighted grid Input: A

weighted grid G with

two distinguished

vertices: a source and

a *sink* 

Output: A longest

path in *G* from *source* 

to sink

Tourists only move south and east, any grid positions west or north of the source are unusable

Any grid positions south or east of the sink are unusable, the source vertex is at (0, 0) Sink vertex at (n, m) defines the southeastern most corner of the grid



The brute force approach search for the longest path

Not an option for medium to large grid

Use a greedy strategy, choose between two possible directions (south or east) by comparing them and selecting one with large increase for one step (local maximum)

Good at beginning

May lead to area with few attractions

No known greedy strategy for the Manhattan Tourist problem provides an optimal solution to the problem



Instead of solving the Manhattan Tourist problem directly, that is, finding the longest path from source (0, 0) to sink (n,m), we solve a more general problem: find the longest path from source to an arbitrary vertex (i, j) with

 $0 \le i \le n, 0 \le j \le m$ . We will denote the length of such a best path as  $s_{i,j}$ , noticing that  $s_{n,m}$  is the weight of the path that represents the solution to the Manhattan Tourist problem

## **Topic #60 Sequence Similarity**

Meaning of "sequence similarity or "distance" between DNA sequences. Hamming distance is not typically used to compare DNA or protein sequences. Calculation rigidly assumes that the *i*th symbol of one sequence is already aligned against the *i*th symbol of the other

It is common case that the *i*th symbol in one sequence corresponds to a symbol at different position in other. Mutation in DNA-evolutionary process: DNA replication- substitutions, insertions, and deletions of nucleotides, leads to "edited" DNA texts. Whether the *i*th symbol in one DNA sequence corresponds to the *i*th symbol in the other

ATATATAT and TATATATA

**А Т А Т А Т А Т** тататата align the (i+1)-st letter in ATATATAT against the ith letter in TATATATA for  $1 \le i \le 7$ 

> ATATATAT and TATAAT- subtle similarities 4 5

> > Α

Δ

Т

Т

3

6 7 8 TA

A

5

т

Т

6

# **Topic # 61 Edit Distance**

1

2 3

ТА

ΤА

2

1

Edit distance between two strings as the minimum number of editing operations needed to transform one string into another, where the edit operations are insertion, deletion, and substitution of one symbol for another

It is often the case that the *i*th symbol in one sequence corresponds to a symbol at different position in other. Mutation in DNA-evolutionary process: DNA replication- substitutions, insertions, and deletions of nucleotides, leads to "edited" DNA texts. Whether the *i*th symbol in one DNA sequence corresponds to the *i*th symbol in the other



# **Topic # 62 Alignment**

The *alignment* of the strings v (of *n* characters) and w (of *m* characters, with *m* not necessarily the same as *n*) is a two-row matrix such that the first row contains the characters of v in order while the second row contains the characters of w in order, where spaces may be interspersed throughout the strings in different places

As a result, the characters in each string appear in order, though not necessarily adjacently.

No column of the alignment matrix contains spaces in both rows, so that the alignment may have at most n + m columns.

Α	Т		G	Т	Т	Α	Т	
Α	Т	С	G	Т		Α		С

Columns that contain the same letter in both rows are called *matches*, while columns containing different letters are called *mismatches*. The columns of the alignment containing one space are called *indels*, with the columns containing a space in the top row called *insertions* and the columns with a space in the bottom row *deletions*. Five matches, zero mismatches, and four indels. The number of matches plus the number of mismatches plus the number of indels is equal to the length of the alignment matrix and must be smaller than n + m

Α	Т		G	Т	Т	Α	Т	
Α	Т	С	G	Т		Α		С

Each of the two rows in the alignment matrix is represented as a string interspersed by space symbols "–"; for example AT--GTTAT-- is a representation of the row corresponding to v = ATGTTAT, while ATCGT--A--C is a representation of the row corresponding to w = ATCGTAC



from (0,0) to (<u>n,m</u>) in that grid

## Topic # 63 Edit Graph 1

The grid that is achieved after alignment is similar to the Manhattan grid where each entry in the grid looks like a city block

The graph called Edit Graph

The main difference between the Manhattan and Edit Graph is that we can move along the diagonals in the Edit Graph



# **Topic # 64 Edit Graph 2**

Analyzing the merit of an alignment-corresponding path in the edit graph. For any two strings- different alignment matrices and corresponding paths.

Surplus of mismatches and indels and a small number of matches, while others have many matches and few indels and mismatches.

Relative merits of one alignment over another-scoring function- input an alignment matrix and produces a score that determines the "goodness" of the alignment.

Variety of scoring functions-higher scores to alignments with more matches.

Column as a positive number if both letters are the same, and as a negative number if the two letters are different. The score for the whole alignment is the sum of the individual column scores.

### **Topic # 65 Longest Common Subsequences**

The simplest form of a sequence similarity analysis is the Longest Common Subsequence (LCS) problem, where we eliminate the operation of substitution and allow only insertions and deletions. A subsequence of a string v is simply an (ordered) sequence of characters (not necessarily consecutive) from v.

v = ATTGCTA AGCA and ATTA are subsequences

TGTT and TCG are not subsequences

A common subsequence of two strings is a subsequence of both of them. Common subsequence of strings

 $\mathbf{v} = v_1 \dots v_n$  and  $\mathbf{w} =$ 

 $w_1$  . . . $w_m$  as a

sequence of positions

in v,

 $1 \leq i_1 < i_2 < \cdots < i_k$ 

 $\leq n$  and a sequence of

positions in w,

 $1 \leq j_1 < j_2 < \cdots < j_k \leq m$  such that the symbols at the

corresponding positions in v and w coincide:

 $v_{\rm it} = w_{\rm jt}$  for  $1 \le t \le k$ 

TCTA is a common to both ATCTGAT and TGCATA

Typically many common subsequences between

two strings v and w-how to find the longest one

s(v,w)-be the length of the longest common subsequence of v and w

edit distance between v and w—under the assumption that only insertions and deletions are allowed—is d(v,w) = n + m - 2s(v,w) and corresponds to the minimum number of insertions and deletions needed to transform v into w

### Alignment v — A T - C - T G A T w — - T G C A T - A -

Above figure presents an LCS of length 4 for the strings v = ATCTGAT and w = TGCATA and a shortest sequence of two insertions and three deletions transforming v into w



Computing similarity s(V,W)=4 V and W have a subsequence TCTA in common



Computing distance d(V,W)=5 V can be transformed into W by deleting A,G,T and inserting G,A

Longest Common Subsequence Problem:

Find the longest subsequence common to two strings

Input: Two strings, v and w

Output: The longest common subsequence of v and w

### **Topic # 66 Recurrence for LCS problem**

Every common subsequence corresponds to an alignment with no mismatches.

This can be obtained simply by removing all diagonal edges from the edit graph whose characters do not match, thus transforming it into a graph like that shown in the figure



An LCS Edit Graph

The relationship between

the Manhattan Tourist problem and the LCS Problem is further illustrate by showing that these two problems lead to very similar recurrences

Define  $s_{i,j}$  to be the length of an LCS between  $v_1 \dots v_i$ , the *i*-prefix of v and  $w_1 \dots w_j$ , the *j*prefix of w. Clearly,  $s_{i,0} = s_{0,j} = 0$  for all  $1 \le i \le n$  and  $1 \le j \le m s_{i,j}$  satisfies the following recurrence

$$Si-1,j$$
  
 $Si,j = \max$   $Si,j-1$   
 $S_{i-1,j-1} + 1$ , if  $v_i = w_j$ 

The first term- when  $v_i$  is not present in the LCS

of the *i*-prefix of v and *j*-prefix of w (deletion of  $v_i$ ); the second term- when  $w_j$  is not present in this LCS (an insertion of  $w_i$ ); and the third term-when both  $v_i$  and  $w_j$  are present in the LCS ( $v_i$  matches  $w_j$ ).

These recurrences can be rewritten by adding some zeros here and there as



's' is used to represent dynamic programming table, the data structure

The length of an LCS between v and w can be read from the element (n,m) of the dynamic programming table, but to reconstruct the LCS from the dynamic programming table, one must keep some additional information about which of the three quantities,  $s_{i-1,j}$ ,  $s_{i,j-1}$ , or  $s_{i-1,j-1} + 1$ , corresponds to the maximum in the recurrence for  $s_{i,j}$ .

## **Topic #67 Algorithms for LCS**

The length of an LCS

Some additional information about which of the three quantities ,  $s_{i-1,j}$  ,  $s_{i,j-1}$ , or  $s_{i-1,j-1} + 1$ 

The following algorithm achieves this goal by introducing backtracking pointers that take one of the three values , or .



#### Algorithms for LCS

```
PRINTLCS(v,w, i,j)
1. for i = 0 or j =
2.
           return
         b<sub>i,j</sub> = K
з.

    PRINTLCS (b, v, i-1, j-1)

5.
    print v_i
6.
    Else
    if b<sub>i,j =</sub> ↑
PRINTLCS (b, v, i-1, j)
7.
8.
9. Else
        PRINTLCS (b,v i, j-1)
10.
```



Dynamic programming table-computation of the similarity score s(v,w) between v and w, while the table on the right-computation of the edit distance between v and w-insertions and deletions are the only allowed operations. The edit distance d(v,w) is computed according to the initial conditions  $d_{i,0} = i$ ,  $d_{0,j} = j$  for all  $1 \le i \le n$  and 1 j m and the following recurrence:

The edit distance d(v,w) is computed according to the initial conditions  $d_{i,0} = i$ ,  $d_{0,j} = j$  for all  $1 \le i \le n$  and 1 j m and the following recurrence:

 $d_{i-1,j+1}$ 

 $\mathbf{d}_{i,j} = \min \qquad \mathbf{d}_{i,j-1} + 1$ 

 $d_{i-1,j-1}$ , if  $v_i = w_j$ 

## **Topic #68 Scoring Alignments 1**

Scoring matrices for DNA sequence-  $\mu$  and  $\sigma$ 

Scoring matrices for protein sequences are complicated – *pointed accepted mutation* (PAM) and *block substitution* (BLOSUM), frequency amino acid 'x' replaces amino acid 'y' in evolutionary related sequences.

Random mutagenesis-change amino acid sequence

Some mutations- do not alter but other do

Some amino acid substitutions are commonly found through the process of molecular evolution-

### **Scoring Alignments**

Asparagine (Asp), Serine (Ser) Glutamate (Glu), Aspartate (Asp) More mutable

#### Cysteine (Cys), Tryptophan (Trp)

ProbabilitySer  $\rightarrow$  Phe3 times moreTry  $\rightarrow$  Phe

Types of changes- most and least common- amino acid scoring matrix, sequence alignment

Amino acids sequences- very few matches-scoring matrix  $\delta(i, j)$  – how often a. a *'i'* substitutes a. a *'j'* 

## **Copic # 69 Scoring Alignments 2**

Large set of alignments of related sequences

Computing  $\delta(i,j)$ 

Amounts to counting how many times the amino acid 'i' is aligned with amino acid ʻj'

Needs to know scoring matrix

Met Ala Phe Ser Gly Asp Glu Ser. . . . . .

Met Ala Phe Ser -- Asp Glu Ser.....

If proteins are 90% identical, premium +1 for matches and -1 for mismatches and indels will do the job. Then "obvious" alignments are constructed that are used to compute scoring matrix  $\delta$ .

The simplified description hides subtle details are important in the construction of scoring matrix

Ser Phe Try Phe

LESS (related proteins in mouse and (related proteins in mouse and rat) human)

15 million years

80 million years

The best scoring matrix to compare two proteins depends on similarity of these organisms

## **Topic #70 PAM matrix**

Problem is rectified- analyzing similar proteins e.g, one mutation/100 a.a. Proteinshuman and chimpanzee, Such sequences, one PAM unit diverged. PAM unit as the amount of time in which an "average" protein mutates 1% of it's a.a. PAM1 scoring matrix- many alignments of extremely similar proteins For a given set of base alignments

define f(i,j) as the total number of times amino acids i and j are aligned against each other, divided by the total number of aligned positions.

f(i,j)

Also define g(i,j) as all proteins from data set.

f(i,j)

where f(i) is the frequency of the amino acid *i* in

g(i,j) defines the probability that an amino acid *i* mutates into amino acids *j* within 1 PAM unit. The (i,j) entry of the PAM 1 matrix is defined as  $\delta(i,j) =$ 

log

g(i, j)

 $=\log f(j)$ (f(i), f(j)), the frequency of aligning amino acid *i* against amino acid *j* that one expects simply by chance). The *PAM* n matrix can be defined as the result of applying the PAM 1 matrix n times
If g is the 20 x 20 matrix of frequencies g(i,j), then  $g^n$  (multiplying this matrix by itself *n* times) gives the probability that amino acid *i* mutates into amino acid *j* during *n* PAM units. The (i,j)

## $\frac{gni, j}{f(j)}$

entry of the PAM *n* matrix is defined as  $\log \frac{f(j)}{f(j)}$ 

For large *n*, the resulting PAM matrices often allow one to find related proteins even when the practically no matches in the alignment. In this case the underlying nucleotide sequences are so diverged that their comparison usually fails to find any statistically significant similarities.

### Similarity between the

Cancer – causing v-sis oncogene

Growth factor PDGF (Platelet derived growth factor)

### **Remained undetected**

Russell Doolittle and colleagues has not transformed the nucleotide sequences into amino acids sequences prior to performing the comparison.

### **Topic #71 Local Sequence Alignment**

Global Alignment- similarity 2 strings- extends over entire length, eg, same protein family- very conserved, same length – fruit flies to humans Score of alignment- 2 substrings 'v', 'w'- larger than score of alignment between entire length of 'v' and 'w'.

*Homeobox* genes, regulate embryonic development- variety of species. Different in different species- one region- *homeodomain*- highly conserved. How to find conserved area- ignore areas-little similarity. Temple Smith and Micheal Waterman proposed modification of the global sequence alignment dynamic programming algorithm-LAP



#### CAGTTATGTCCAG



The conserved domain are identical and cover one third of the entire length, n, of ese genes. Path th from source to sink will include approx  $\frac{2}{3}n$ horizontal edges,  $\frac{1}{3}n$  diagonal match edges (corresponding to homeodomains) and  $\frac{2}{3}n$  vertical edges

The score of this path is



The score of this path is



This path contains so many indels that it is unlikely to be the highest scoring alignment.

Biologically irrelevant diagonal paths - likely have score- mismatches are

penalized less than indels. Score of diagonal path is  $n(\frac{1}{4} - \frac{3}{4}\mu)$ , since every diagonal edge  $\frac{1}{4}$  and mismatch with probability . Since  $(\frac{1}{3} - \frac{4}{3}\sigma) < (\frac{1}{4} - \frac{3}{4}\mu)$  for indels and mismatch penalties, global alignment- miss correct solution of real biological problem, biologically irrelevant near-diagonal path.

**Topic # 72** 

same TOPIC 71

### **Topic #73 Local Alignment Problem**

Biological significant in certain parts of DNA fragments, maximize the alignment score s ( $v_i$ 

... $v_{i'}$ ,  $w_j$ ... $w_{j'}$ ) over all substrings  $v_i$ .... $v_{i'}$  of v and  $w_j$ .... $w_{j'}$  of w. Local Alignment Problem- not extend over entire length as in Global Alignment Problem

Local Alignment Problem:

Find the best local alignment between two strings

Input: Strings v and w and a scoring matrix  $\boldsymbol{\delta}$ 

Output: Substring of v and w whose global alignment, as defined by  $\delta$ , is maximal amoung all global alignment of all substrings of v and w.



Global Sequence Alignment- finding the longest path between vertices (0,0) and (*n*,*m*) in the edit graph

Local Alignment-finding the longest path among the paths between arbitary *vertices* (i, j) and (i', j') in the edit graph.

Find the longest path between every pair of vertices (i, j) and (i', j')- then select longest of these computed paths. Instead of finding the longest path from (i, j) to (i', j'), LAP- finding the longest path from the *source* (0,0) to every other vertex by adding edges to weight 0

Local Alignment Problem

 $s_{i,j} = \max \begin{bmatrix} 0\\ s_{i-1,j} + \delta(v_i, --)\\ s_{i,j-1} + \delta(--, w_j)\\ s_{i-1,j-1} + \delta(v_i, w_j) \end{bmatrix}$ 

the edit graph

These edges make the source vertex (0,0) a predecessor provide a "free ride" from the source to any other vertex (i,

reflects the transformation of

j). Following recurrence

## **Topic #74 Progressive Multiple Alignment**

Another approach-strong pairwise alignment Greedy progressive multiple alignment heuristic-pair of strings with greatest similarity-new string-"once a gap, always a gap." Multiple alignment of k sequences is reduced to the multiple alignment of k-1 sequences.

The motivation for the choice of the closest strings at the early steps of the algorithm is that close strings often provide the most reliable information about a real alignment

Many popular iterative multiple alignment algorithms including the tool CLUSTAL, use similar strategies

### ATGTCATATTCGGAC

### ATGTCATATTCGGAC

ATGTCATATTCGGAC

### ATG- CATA

### ATGTCATA

Progressive multiple alignment algorithms- problem with CLUSTAL-may be misled by some spuriously strong pairwise alignment effect, a bad seed. The error in initial pairwise alignment will propagate all the way through to the whole multiple alignment. Many algorithms have been proposed,-even with systematic deficiencies are quite useful in computational biology

Multiple alignment for k sequences

Generalization of the Pairwise Alignment problem Existence of a k-dimensional

scoring matrix k-dimensional scoring matrices are not very practical Describe two

other scoring approaches that are

more biologically relevant. The choice of the scoring function can drastically affect the quality of the resulting alignment, and no single scoring approach is perfect in all circumstances.

Multiple alignment of k sequences-a path of edges in a k-dimensional-Manhattan gridlike edit graph.

The weights of the edges-scoring function

Intuitively, assign higher scores to the columns with a low variation in letters-high scoreshighly conserved sequences

*Multiple Longest Common Subsequence* problem, the score of a column is set to 1 if all the characters in the column are the same, and 0 if even one character disagrees

## **Topic #75 Gene Prediction 1**

Sydney Brenner and Francis Crick

Every triplet codes for one amino acid

Introduce deletions in DNA-dramatically alters its protein product. Deleting three consecutive nucleotides results in minor changes in the protein

The phrase

THE SLY FOX AND THE SHY DOG

(written in triplets)

Turns into nonsense after deleting one letter Y from SLY

THE SYF OXA NDT HES HYD OG

or two letters LY from SLY

#### THE SFO XAN DTH ESH YDO G

but makes some sense after deleting three nucleotides SLY

THE SOX AND THE SHY DOG

Charles Yanofsky proved that a gene and its protein product are collinear. Yanofsky's experiment was so influential that nobody even questioned about codons and for almost 2 decades biologists believed that a protein was encoded by a long string of adjacent triplets.

Discovery of split human genes-collection of substrings

Raised the computational problem-prediction of genes

Human genome is larger and complex than bacterial genome

Salamander genome is ten times larger than the human genome amounts of so-called junk DNA Large

Human genes - exons that are separated by this junk DNA. The difference in the sizes of the salamander and human genomes thus presumably reflects larger amounts of junk DNA and repeats in the salamander genome.

Split genes are analogous to a magazine article- page 1, 13, 43, 51, 74, 80, and 91, with pages of advertising appearing in between. Junk DNA represents "advertising" that separates exons.

## **Topic #76 Gene Prediction 2**

The jump is inconsistent from species to species.

A gene in insect genome- different in worm

Number of exons

The information in one part in human-broken up into two in the mouse.

While the genes themselves are related, they may be quite different in terms of the parts' structure

Split genes-1977

Phillip Sharp and Richard Roberts- Adenovirus

"An Amazing Sequence Arrangement at the 5' End of Adenovirus 2Messenger

RNA." Sharp's group- hexon.

Hexon mRNA, mRNA was hybridized to adenovirus DNA- the hybrid moleculeselectron microscopy.

mRNA-DNA hybrids-three loop structures-continuous duplex segment- classic continuous gene model



newly sequenced gene has a good chance of being related to one that is already known. For example, 99% of mouse genes have human analogs.

Simply look for a similar sequence-based on the genes known in another-exon sequence and the exon structure are different

The commonality-produce similar proteins

Similarity-based methods attempt to solve a combinatorial puzzle-putative exons in a genomic sequence – mouse-human

Know a human protein, and we want to discover the exon structure of the related gene in the mouse genome. The more sequence data we collect, the more accurate and reliable similarity based

methods become. Consequently, the trend in gene prediction has recently shifted from statistically motivated approaches to similarity-based algorithms

### **Topic #79 Statistical Approach to Gene Prediction 1**

Statistical approaches to finding genes-statistical variations between coding (exons) and noncoding regions

*Open reading frames* (ORFs) Fenome of length n as a sequence of n/3 codons

The three "stop" codons, (TAA, TAG, and TGA)

The subsegments of these that start from a start codon, ATG, are ORFs.

ORFs within a single genomic sequence may overlap since there are six possible "reading frames": three on one strand starting at positions 1, 2, and 3, and three on the reverse strand



### The six reading frames for the DNA sequence

One would expect to find frequent stop codons in noncoding DNA, since the average number of codons between two consecutive stop codons in "random"

DNA should be  $64/3 \sim 21$ . This is much smaller than the number of codons in an average protein, which is roughly 300. Therefore, ORFs longer than some threshold length indicate potential genes. However, gene prediction algorithms based on selecting significantly long ORFs may fail to detect short genes or genes with short exons

Many statistical gene prediction algorithms rely on statistical features in proteincoding regions, such as biases in *codon usage*. We can enter the frequency of occurrence of each codon within a given sequence into a 64-element *codon usage array* 

Π	U		C		A		G	
	UUU Phe	57	UCU Ser	16	UAU Tyr	58	UGU Cys	45
	UUC Phe	43	UCC Ser	15	UAC Tyr	42	UGC Cys	55
U	<b>UUA</b> Leu	13	UCA Ser	13	UAA stp	62	UGA stp	30
	UUG Leu	13	UCG Ser	15	UAG stp	8	UGG Trp	100
-2	CUU Leu	11	CCU Pro	17	CAU His	57	CGU Arg	37
0	CUC Leu	10	CCC Pro	17	CAC His	43	CGC Arg	38
C	CUA Leu	4	CCA Pro	20	CAA Gln	45	CGA Arg	7
	CUG Leu	49	CCG Pro	51	CAG Gln	66	CGG Arg	10
	AUU Ile	50	ACU Thr	18	AAU Asn	46	AGU Ser	15
٨	AUC Ile	41	ACC Thr	42	AAC Asn	54	AGC Ser	26
A	AUA Ile	9	ACA Thr	15	AAA Lys	75	AGA Arg	5
	AUG Met	100	ACG Thr	26	AAG Lys	25	AGG Arg	3
	GUU Val	27	GCU Ala	17	GAU Asp	63	GGUGly	34
0	GUC Val	21	GCC Ala	27	GAC Asp	37	GGC Gly	39
G	GUA Val	16	GCA Ala	22	GAA Glu	68	<b>GGA</b> Gly	12
	GUG Val	36	GCG Ala	34	GAG Glu	32	GGGGly	15

### **Topic # 80 Statistical Approach to Gene Prediction 2**

Π	U		С		A	01000	G	
Π	UUU Phe	57	UCU Ser	16	UAU Tyr	58	UGU Cys	45
TT	UUC Phe	43	UCC Ser	15	UAC Tyr	42	UGC Cys	55
U	UUA Leu	13	UCA Ser	13	UAA stp	62	UGA stp	30
	UUG Leu	13	UCG Ser	15	<b>UAG</b> stp	8	UGG Trp	100
	CUU Leu	11	CCU Pro	17	CAU His	57	CGU Arg	37
0	CUC Leu	10	CCC Pro	17	CAC His	43	CGC Arg	38
-	CUA Leu	4	CCA Pro	20	CAA Gln	45	CGA Arg	7
	CUG Leu	49	CCG Pro	51	CAG Gln	66	CGG Arg	10
Π	AUU Ile	50	ACU Thr	18	AAU Asn	46	AGU Ser	15
٨	AUC Ile	41	ACC Thr	42	AAC Asn	54	AGC Ser	26
A	AUA Ile	9	ACA Thr	15	AAA Lys	75	AGA Arg	5
	AUG Met	100	ACG Thr	26	AAG Lys	25	AGG Arg	3
Π	GUU Val	27	GCU Ala	17	GAU Asp	63	GGU Gly	34
0	GUC Val	21	GCC Ala	27	GAC Asp	37	GGC Gly	39
9	GUA Val	16	GCA Ala	22	GAA Glu	68	<b>GGA</b> Gly	12
	GUG Val	36	GCG Ala	34	GAG Glu	32	GGGGly	15

Some more facts about genetic code and codon usage in humans

#### in-frame hexamer count Mark Borodovsky

Gene prediction in bacterial genomes-several conserved sequence motifs often found in the regions around the start of transcription

Such sequence motifs are more elusive in eukaryotes

The approaches-prokaryotes-eukaryoyes

Exons-130 nucleotides-reliable peaks in the likelihood ratio plot while analyzing ORFs-do not differ enough from random fluctuations to be detectable. Moreover,

codon usage and other statistical parameters probably nothing in common in splicing machinery recognizes exons

### **Topic # 81 Statistical Approach to Gene Prediction 3**

Biologically oriented

Approach

Recognize the locations of splicing signals at exon-intron junctions

There exists a weakly conserved sequence of eight

nucleotides at the boundary of an exon and an intron (donor splice site) and a sequence of four nucleotides at the boundary of an intron and exon (acceptor splice site)



Numbers = Frequency (%)

Profiles for splice sites are weak-limited success Hidden Markov Model (HMM) approaches that capture statistical dependencies between sites GENSCAN developed by Chris Burge and Samuel Karlin. GENSCAN combines coding region and splicing signal predictions into a single framework.

Splice site prediction-coding region appear on one side of the site Such statistics are used in the HMM framework of GENSCAN that merges splicing site statistics, coding region statistics, and motifs near the start of the gene The accuracy of GENSCAN decreases for genes with many short exons or with unusual codon usage

### **Topic # 82 Similarity Based Approached to Gene Prediction 1**

A similarity-based approach-Previously sequenced genes Unknown genes in newly sequenced DNA fragments Combinatorial puzzle-find a set of substrings (candidate exons) whose splicing best fits the target Brute force approach-find all local similarities

Each substring from the genomic sequence that exhibits sufficient similarity to the target protein could be considered a putative exon Exon-flanking dinucleotides AG and GT

Overlapping

Model a putative exon with a *weighted interval* in the genomic sequence, parameters (l, r, w)

l is the left-hand position, r is the right-hand position, and w is the weight of the putative exon "w"- local alignment score

Likelihood that this interval is an exon

A chain is any set of non overlapping weighted intervals. Total weight of a chain

A maximum chain



### **Topic #83 Similarity Based Approached to Gene Prediction 2**

Model a putative exon with a weighted interval in the genomic sequence



Five weighted intervals, (2, 3, 3), (4, 8, 6), (9, 10, 1), (11, 15, 7), and (16, 18, 4), shown by bold edges, form an optimal solution to the Exon Chaining problem. The array at the bottom shows the values  $s_1, s_2, \ldots, s_{2n}$  generated by the EXONCHANING algorithm

### **Topic # 84 ORF Prediction**

### **ORF (Open Reading Frame)**

Gene finding, especially in prokaryotes starts form searching for open reading frames (ORF)

An ORF is a sequence of DNA that starts with start codon "ATG" (not always) and ends with any of the three termination codons (TAA, TAG, TGA)



Here, in this figure we see a comparison between prokaryotic and eukaryoticcells. Prokaryotes maybe bacteria or some related organisms whereas rest of the organisms is classified in eukaryotes.

If we look into the prokaryotic situation, we

have different genes which are separated by Intergenic regions and these ORF can be spanning these two genes whereas in case of the eukaryotes, we have only one gene, we have exon1 and exon2 whereas the ORF spans cross these different exons.

So on the top (in the figure), we have the longer ORF whereas in the bottom we have a shorter one as compared to the genome size those shorter ORF which are actually spanning different exons.

### **ORF** and gene finding:

- ORF provide important evidence in gene finding.
- Generally longer ORFs are preferred.
- However presence of ORF not necessarily means the region is translated to a functional product

### **Reading Frames:**

Depending on the start point, we can define different ORFs, so if we want to go for those triplet codons, we can start with any nucleotide, in this way we have three different possibilities for one of the strands and since we have two strands in the DNA, so in total we can have six ORFs (so 3 of them are from 3' to 5' direction whereas the other 3 are from 5' to 3' direction).

Six Frame translation +3ATGGTTTGGGA +2ATGGTTTGGGA +1ATGGTTTGGGA 5'-ATGGTTTGGGAACCGAAGTCAATT-3' 3'-TACCAAACCCTTGGCTTCAGTTAA-5' GCTTCAGTTAA-1 GCTTCAGTTAA-2 GCTTCAGTTAA-3 \*Three on forward strand and three on complementary strand

This is how those 6 ORFs looks like. So, there are the six frame translations (in this picture). We observe that we have a top strand which is a forward strand and starts at 5' end and ends at 3' end.

A complementary runs in an anti-parallel fashion which starts with 3' end and ends at 5' end.

So, how can we get the reading frames out of it?

We can start with the position number 1 on the first strand and we label it as +1 (as shown), in this way we can start with A, and have triplet codons like ATG, GTT, TGG and so on. In the second strand or second possibility (denoted by +2), we can start from the position number 2, so this frame starts from the second nucleotide like T and makes a triplet codon as TGG, TTT, GGG and so on.

So this how the third strand  $(3^{rd}$  possible ORF; denoted by +3), where it starts from the position number 3 can be made. We don't start with the position number 4 because it will be same as the position number 1.

Similarly, we can do like this for the opposite direction strands with the possibility of position number 1, 2 and 3.

So, this how the 6 ORF are made.

### **Conclusions:**

An ORF is a sequence of DNA that starts with start codon "ATG" (not always) and ends with any of the three termination codons (TAA, TAG, TGA) and there are 6 reading registers as far as the ORFs are considered from the sequence.

### **Reference:**

**Biological Sequence Analysis** 

R Durbin, S Eddy, A Krogh and G Mitchison

Cambridge University Press, 1998.

### Bioinformatics The machine learning approach

P Baldi and S Brunak

The MIT Press, 1998

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M Kanehisa

## **Topic # 85 ORF Finders**

### **ORF Finding:**

- Long ORF may be a gene.
- Expected  $64/3 \sim 21$  codons before we see a stop codon.
- Genes are longer than this.
- We might scan for ORF longer than a threshold

### Codon usage and likelihood ratio:

- An ORF is more 'reliable' if it has 'likely' codons
- We can do sliding window calculations (focus on some nucleotides like for example if the segment is 1000 long, we can make a window of say size 50 and we can look into the frequencies of different codons in that window) to ORF having 'likely' codon usage.
- An ORF is more 'reliable' if it has 'likely' codons
- However average vertebrate exon length (130 nucleotides) is too small for reliable peaks

An improvement may be;

### In-Frame hexamer count

i.e. frequencies of pairs of consecutive codons.

### **ORF Finders:**

S ncbi

Tools are mainly based on pattern finding algorithms

- 1. NCBI's ORF Finder
- 2. ORF Investigator
- 3. OrfPredictor

#### ORF Finder (Open Reading Frame Finder)

The ORF Finder (Open Reading Frame Finder) is a graphical analysis tool which finds all open reading frames of a selectable minimum size in a user's sequence or in a sequence already in the database. This tool identifies all open reading frames using the standard or alternative genetic codes. The deduced amino acid sequence can be saved in various formats and searched against the sequence database using the WWW BLAST server. The ORF Finder should be helpful in preparing complete and accurate sequence submissions. It is also packaged with the Sequin sequence submission software. or sequence in FASTA format Enter GI or ACCESSION

atgcccaagctgaatagcgtagaggggttttcatcattt gaggacgatgtataa

If you go to NCBI webpage, the ORF Finder is there.

You can place you sequence into the text box and upload it while simply clicking onto the 'OrfFind' button. You can even grasp the sequence automatically by writing the accession number in the space give after the word 'ACESSION'.



When you click 'OrfFind', it will take you to this page where there are different registers or ORFs (explained in the previous lecture).

The steric (\*) here means that the protein synthesis stops here.

1 atgcccaagctgaatagcgtagaggggttttcatcatttgaggacgatgtataa 54 M P K L N S V E G F S S F E D D V \*

### **Conclusions:**

- ORF provides important evidence in gene finding.
- Generally longer ORFs are preferred.
- However presence of ORF not necessarily means the region is translated to a functional product

### **TOPIC 86 Translation Start Site (TSS)**

### **Translation Start Site (TSS)**

Translation starts with ATG that codes for methionine in a polypeptide

GCCATGGCGA ... ACGATGCTGT ... GACATGGTAC ... AGGATGGGCT ...

Here, in this example shown in the figure, we have those TSS in the middle (it's actually a file where sequences are aligned with one another), we look deeply, we can see that their neighbors are C and G which are most frequently seen (if not always). We might use these neighbors to identify the presence of these TSS.

### Assumption:

- Certain nucleotides prefer to be around TSS than others.
- The "biased" nucleotide distribution is information is a basis for translation start prediction

### **Coding Potential:**

**Hexamer** frequencies in coding versus non-coding regions may provide important insights Frequency of X(A,G,C,T) at position i is

 $F_i(X) = \Sigma \log(C_i(X)/N_i(X))$  (frequency of any nucleotide can be found by taking the sum of the log of ratio of the counts of the particular nucleotide in that particular position divided by the total)

\*where c is the counts.



Based upon the frequency equation, we can come up with a frequency table as shown in the figure.

The frequencies shown here indicates

that you have the presence of true TSS here or you can expect that there are some Transcription Start Sites (TSS) here.

You can observe that on different positions like -4, -3, -2 and -1, similarly at +3, +4,+5, and +6 you have which nucleotides and what are their percentages (in the table).

### Example

Which one is more probable to be a Translation Start?

### CACC ATA GC TCGA ATG TT

### Solution

We can use frequency table and the scoring function as under;

### $S_i = \Sigma \log (F_i (X)/0.25)$

-frequency from the frequency table divided by the expected frequency and then we convert it into log scale because we want to play with big numbers.

We can call this equation as the Information Content (IC)

### Frequency table (Biased Nucleotide distribution) ATG

A17.36,19.01,17.36 48.76 28.93,15.70, 21.49,23.14,19.83,21.49,25.62,15.70, C16.53,28.93 57.85 5.79,39.67,50.41, 22.31,38.84,23.97,27.27,31.40,38.02, G46.28,29.75,19.01,42.98,14.88,26.45, 42.98,23.97,33.88,32.23,25.62,29.75, T 9.83,22.31, 5.79, 2.48,16.53, 7.44, 13.22,14.05,22.31,19.01,17.36,16.53]

-4 -3 -2 -1 +3 +4 +5 +6

CA

Here, is our frequency table, we pick the frequencies from here.

### CACC ATA GC

log (58/0.25) + log  $(49/0.25) + \log (40/0.25) + + \log (15/0.25) + \log$ log (50/0.25) + log (43/0.25) + log (49/0.25)

**TCGA ATG TT** log (6/0.25) + log (6/0.25)  $(7/0.25) + \log(13/0.25) +$ log (14/0.25)

= 9.44

And we add those frequencies here in the equation. So, in this way we get a positive number in the end,

so we have 13.69 on the left side whereas we have 9.44 on the right side.

So, which sequence has the strong evidence to have a translation site. In our case, we will prefer the one with higher value so its probably the green sequence.

### Algorithm:

=13.69

- > Build a mathematical model, based on collected translation start sequence
- > For each candidate translation start sequence, apply the model and get a score
- > If the score is larger than zero, predict it is a "translation start"; the higher score, the higher the probability the prediction is true

### **Conclusions:**

- TSS prediction can be an important step in gene prediction •
- TSS can be predicted while using the frequency of neighboring nucleotides

### **References:**

### **Biological Sequence Analysis**

R Durbin, S Eddy, A Krogh and G Mitchison

Cambridge University Press, 1998.

### Bioinformatics The machine learning approach

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The MIT Press, 1998

### **Post-Genome Informatics**

M Kanehisa

Oxford University Press, 2000

### **TOPIC 87 Prediction of splice junctions**

### **Splice Junctions:**

### **Donor site**

Coding region | GT (introns starts)

### Acceptor

- (introns ends)YAG | coding region -Y can be any pyrimidine.
- **Canonical form** 
  - GT-AG: 99.24%

There are also some non-canonical forms.

Like TSS, the flanks of splice junctions show "biased" distributions of nucleotides in certain positions

• These biased distributions of nucleotides are the basis for prediction of splice junctions



Here, is the example where we can see we have the exon then we have the donor site where we see a big GT (this representation is known as sequence logos), then we have the intron region which is followed by the acceptor region which ends up with AG and then again a next exon starts.

### **Sequence LOGOS:**

- A visual representation of a position-specific distribution
- Easy for nucleotides, but we need colour to depict up to 20 amino acid proportions.

Overall height at position is proportional to the information content

• Proportions of each nucleotide/amino acid are in relation to their observed frequency, with most frequent on top, next most frequent below

### Non Canonical Splice Junctions:

In addition to canonical GT-AG (99.24%);

GC-AG: 0.69%

AT-AC: 0.05%

Others: 0.02%

### **Information Content (IC):**

### $S_i = \Sigma \log (F_i (X)/0.25)$

- If every nucleotide has 0.25 frequency in a position, then the position's information content is ZERO.
- Use "information content as a criterion for determining the length of flanks

### Accepter site prediction

	-6	-5	-4	-3	-2	-1	1
Α	12.7	9.5	26.2	6.3	100	000	21.4
С	40.5	36.5	33.3	68.2	000	000	2.0
G	2.4	6.3	13.5	000	000	100	62.7
T/U	44.5	47.6	27.0	25.2	000	000	7.90

### Multiple positions have high information content

### Donor site prediction

	-3	-2	-1	1	2	3	4
Α	34.0	60.4	9.2	000	000	52.6	71.3
С	36.3	12.9	3.3	000	000	2.8	7.6
G	18.3	12.5	80.3	100	000	41.9	11.8
T/U	11.4	14.2	7.3	000	100	2.5	9.3

Here, is the acceptor site distributions and we can observe that at position number -2 and -1, there are the presence of the acceptor sites.

100% frequencies of A and G, so these are predicted but if we look into the neighbors like at position -3, there are mostly Cs.

We can look into the donor sites; we have 100% frequency for Gs and Ts, we label them as 1 and 2, and different positions which are neighboring to them, we can include 10 or 15 neighbors.

## Multiple positions have high information content

### Algorithm:

Mathematical model: F<sub>i</sub> (X): frequency of X (A, C, G, T) in position I

Score a segment as a candidate donor/acceptor site by

 $\Sigma \log (F_i (X)/0.25)$ 

For each candidate sequence, apply the model and get a score

If the score if larger than zero, predict it is "donor/acceptor"; the higher score, the higher the probability the prediction is true

### **Conclusions:**

Like TSS, the flanks of splice junctions show "biased" distributions of nucleotides in certain positions

These biased distributions of nucleotides can be used for prediction of splice junctions

### **References:**

**Biological Sequence Analysis** 

R Durbin, S Eddy, A Krogh and G Mitchison

Cambridge University Press, 1998.

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M Kanehisa

Oxford University Press, 2000

## **TOPIC 88 Prediction of Exons**

### Introduction:

Exons can be predicted within ORF from the information gathered about

splice junctions

### Approach:

For each segment [acceptor, donor], we get three scores (coding potential, donor score, acceptor score)

### Various possibilities

- all three scores are high probably true exon.
- all three scores are low probably not a real exon.
- all in the middle -- ?.
- some scores are high and some are low -- ??

So here, we can get the evidence by the help of that information which we gathered from those splice sites.

### **Prediction:**

- Collect a set of exons and non-exons
- Score them using our scoring schemes
- Plot them as follows
- "draw" a separating line between exons and non-exons

### Linear Discriminate Analysis:

- linear discriminate analysis (LDA) finds an optimal plane surface that best separates points that belong to two classes.
- For example, if there are ten true exons and ten introns, and two feature.
- These samples could be represented by 20 points in a two-dimensional space.
- LDA would compute a straight line through the space that can best separate the two classes with the minimal classification error



For example, if we look into this picture, there is the coding region and the noncoding region and there is a line in the middle that tries to separate these two which helps in discrimination.

We can draw a central line or linear regression line, we can see the equation

over there and by drawing this line we can have the prediction, this line fits in such a way that it fits into most of the data points.

If we don't know about data point, we can predict it while using this prediction line (angular line in the figure).

### **Conclusion:**

- Collect a set of exons and non-exons
- Score them using our scoring schemes
- Plot them as follows
- "draw" a separating line between exons and non-exons

### **References:**

### **Biological Sequence Analysis**

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### **TOPIC 89** Annotation of Assembled Genome

### A genome sequence is useless without annotation

Three steps in genome annotation:

• Find features not associated with protein-coding genes (e.g. tRNA, rRNA, snRNA, SINE/LINE, miRNA precursors)

A genome sequence is useless without annotation

Three steps in genome annotation:

• Build models for protein-coding genes, including exons, coding regions, regulatory regions

### A genome sequence is useless without annotation

Three steps in genome annotation:

• Associate biologically relevant information with the genome features and genes

### Ab initio methods:

Based on sequence alone

- Gene prediction algorithms (e.g. AUGUSTUS, Glimmer, GeneMark)
- RepeatMasker(repeat families)

### **Evidence-based Methods:**

- Require transcriptome data for the target organism (the more the better)
- Align cDNA sequences to assembled genome and generate gene models: TopHat/Cufflinks, Scripture

### **Biological Annotations:**

BLAST of gene models against protein databases

- Sequence similarity to known proteins
- InterProScan of predicted proteins against databases of protein domains
  - Pfam, Prosite, HAMAP, PANTHER, ...
- Mapping against Gene Ontology (function of those genes) terms
  - GO terms
  - BLAST2GO

### **Pattern Finding:**

- Much of the data processing in bioinformatics involves searching and recognizing certain patterns within DNA, RNA or protein sequences.
- In Biology it means finding motifs in DNA or proteins while in computational means it is finding a pattern in a string

### **Conclusions:**

After a genome is assembled, genome annotations are performed to identify gene and other features in a genome

### **TOPIC 90** Pattern Finding in a Genome

### Vocabulary:

- A pattern (keyword) is an ordered sequence of symbols .
- Symbols of the pattern and the searched text are chosen from a predetermined finite set, called an **alphabet** ( $\Sigma$ )

### Four Cases of Pattern Finding:

Look for a perfect match

### CGTA CGTA

- Allow errors due to substitutions

CGTA CG<mark>G</mark>A

- Allow errors due to insertions-deletions

(InDels).



Rank possible matches according to a weight function and keep matches above a certain threshold

### Four Cases of Pattern Finding:

*p1* and *p2* are two patterns of length 5 CTGTA CCGGA

 W is the weight of complete patterns defined via a nucleotide-nucleotide weight function w()

$$W(p1, p2) = \sum_{i=1}^{5} w(p1[i], p2[i])$$

### **Generalized Algorithm:**

Goal: Finding all occurrences of a pattern in a text

#### Input:

Pattern  $\mathbf{p} = [\mathbf{p1...pn}]$  of length n

Text  $\mathbf{t} = [\mathbf{t1...tm}]$  of length m

### **Output:**

An indication that pattern P exist in T

or it does not exist in text T

Index	0	1	2	3	4	5	6	7	8	9	10	11	12	13
Text:	а	t	a	С	а	а	t	а	t	а	С	a	t	a
						Т	Т	Т	Т	Τ	Т			
Pattern:								a	t	a	c	a	t	

Here P is a substring of T i.e., P=T[7,...,12]

from 0 and ending at 13), and the pattern length is 6.

For every pair, aligned together, we come up with the weight function and in the end we combine all those weight functions

Here we are using arrays of data structure, so we have text in those arrays and the indexes are the positions of those letters.

The total length is 14 (starting

So, the pattern matches on the seventh index, so what we will get in the end is that P is a substring of T, starting from 7th index to 12th.

### **Conclusion:**

Pattern searching algorithms search specific sequences in strands of DNA, RNA and proteins having important biological meaning

## **TOPIC 91** Pattern Finding Algorithms

#### **Methods Devised for Pattern Finding:**

- Exact searching methods
- Approximate searching methods
- Position weight matrices
- Suffix trees

### **Exact Pattern Matching:**

- Given a pattern p of length **m**
- and a string or text **T** of length  $n \ (m \le n)$

- Find all the occurrences of p in T

### The matching needs to be exact, which means that the exact word or pattern is found

### **Exact Pattern Matching Algorithms:**

- Naïve Brute Force algorithm
- Boyer-Moore algorithm
- Knuth Morris Pratt algorithm

### **Approximate Pattern Matching**:

Also referred as approximate string matching or matches with k mismatches or differences

Also referred as approximate string matching or matches with k mismatches or differences

Given: a pattern *p* of length *m* 

and

#### a string or text **T** of *length* $n (m \le n)$

Find: all the occurrences of substring X in T that are similar to p, allowing a limited number, say k different characters in similar matches

### **Approximate Pattern Matching Algorithms**

- Dynamic programming approach
- Automata approach
- Filtering and automation algorithms

### **Position Weight Matrices:**

Also known as position specific scoring matrices (PSSM)

• A matrix representing the frequencies of residues observed for a position in multiple alignment

### **Suffix Trees:**

It is a compressed tree containing all the suffixes and allows many problems on strings to be solved quickly

### Conclusions:

- Exact searching or pattern matching methods
- Approximate searching or pattern matching methods
- > Position weight matrices.
- Suffix trees

### **TOPIC 92** Methods Devised for Pattern Finding

#### **Introduction:**

Also known as exhaustive search algorithm

- > All the possibilities are explored and the best one is chosen
- > For the task with many possibilities brute force will take too much time

### Working:

- Searches patterns by going through the whole sequence nucleotide per nucleotide.
- Always shifts the window by exactly one position to right
- Requires 2n expected text character comparisons

When a mismatch the comparison stops and starts again by moving the pattern one position forward

### Algorithm:

### Brute\_Force(T,P)

n length[T]

m length[P]

For s 0 to n-m

Do if P[1..m]= T[s+1...s+m]

### print "pattern occurs at position" s+1

Here, Brute\_Force is the function which has two arguments (T= text and P= pattern) where n records the length of T and m records the length of P and we start with a for loop which goes from 0 to n-m, say for example we have 'T' which is of length 10 and 'P' which is of length 5 so it starts from 0 and will go till 5.

### Do if P[1..m]= T[s+1...s+m]

This line is where we are saying that nucleotide number 1 of the pattern and we go up to its whole length. In case of the text, we started with 0, so we add one over here, so 0+1 i.e. the 1<sup>st</sup> nucleotide is compared till the last nucleotide.

So, if we find the occurrence of the patterns we will put that in the print statement and s+1 will give its position.

working



### Drawback:

The repetitive use of residues in comparison leads to runtime of O(mn), which makes it very slow

### **Conclusion:**

Brute force is an exhaustive search method that takes long time as it does nucleotide by nucleotide comparison

### **TOPIC 93 Knuth-Morris-Pratt Algorithm**

### **Introduction:**

A linear time algorithm for string matching

• Does not involve backtracking on string *s* i.e., repetitive comparison of nucleotide residues

### **Components:**

- The Prefix Function
- The KMP Matcher

### **The Prefix Function Π:**

Encapsulates knowledge about how the pattern matches against shifts of itself

- This information can be used to avoid useless shifts of the pattern 'p'
- This enables avoiding backtracking on the string 'S'

#### The KMP Matcher:

Given: string 'S', pattern 'p' and prefix function 'II'

**Find:** the occurrence of '*p*' in '*S*'

Return: the number of shifts of 'p' after which occurrence is found

```
Compute-Prefix-Function (p)
1
  m ← length[p]
                     //'p' pattern to be matched
2 Π[1] ← 0
3 k ← 0
4
    for q \leftarrow 2 to m
5
      do while k > 0 and p[k+1] != p[q]
6
       do k ← Π[k]
      If p[k+1] = p[q]
7
8
      then k ← k +1
9
      Π[q] ← k
10
     return Π
```

The KMP Matcher (S, p)

```
1 n \leftarrow \text{length[S]}
```

```
2 m \leftarrow length[p]
```

3 Π ← Compute-Prefix-Function(p)
 4 q ← 0 //number of characters matched

4 q ← 0 //number of characters matched 5 for i ← 1 to n //scan S from left to right

- 6 do while q > 0 and p[q+1] != S[i]
- 7 do q ← Π[q] //next character does not match
- 8 if p[q+1] = S[i]
  9 then q ← q + 1 //next character matches



### **Conclusions:**

A linear time algorithm for string matching

- avoid useless shifts of the pattern 'p'
- Does not involve backtracking on string *S*

### **Topic 94 Knuth-Morris-Pratt Algorith**

### Components

- The Prefix Function
- The KMP Matcher

Cor	npute-Prefix-Functi	on (	P)						
1 m	n ← length[p]	a	1	2	3	4	5	6	7
2	<b>[1] ← 0</b>	-	-	-	-	-	-	-	t-
3 k	←0	р	a	ι	a	τ	a	C	a
4	for q ← 2 to m		0						
5	do while k > 0 and	p[k+	-1]	!=	p[q	1]			
6	do k ← ∏[k]	Imi	4:-			1214			
7	lf p[k+1] = p[q]	Ini	ua						-
8	then k ← k +1		m	= 1	en	gtn	[p]	=	1
9	∏[a] ← k	-	1.11	11	= 0	<			
10	return П		к :	= 0					
								-	

### Compute-Prefix-Function (p)



## Compute-Prefix-Function (p)

1 m ← length[p]	q	1	2	3	4	5	6	7	'
2 ∏[1] ← 0	n	a	t	a	t	a			
3 k ← 0		a 0		4					4
4 for q ← 2 to m	Ш	0	0	1	2				
5 do while k > 0 and p	)[k+	-1]	!=	p[o	q]				
6 do k ← Π[k]	Ste	n 3	۰. u		4 4	-	1		
7 If p[k+1] = p[q]	510		n∐ n	[1+	41	= r		1	
8 then k ← k +1	E	2	K	5	1+1		74 2	4.	
9 ∏[q] <del>(</del> k			I.		714	1 =	2		
10 return П					.1.4	1	-		
	is.							-	
Compute-Prefix-Euncti	on	(n	1						
1 m $\leftarrow$ longth[n]			1						
$2 \Pi (1) \leftarrow 0$	(	9	1	2	3	4	5	6	7
		o	a	t	а	t	а	С	a
$\int \mathbf{k} \mathbf{c} 0$	Ī	7	0	0	1	2	3		
$4  101 \mathbf{q} \leftarrow 2 \\ 101 \mathbf{n}$	ין שר		411			1	•		
	ρĮ	Κ+	11:	- I	νĮq	1			
	S	ite	p 4	<u>4: c</u>	1 =	5,	k =	2	
7 If $p[k+1] = p[q]$				p[2	2+1	1] =	: p	<b>q</b> <sub>5</sub>	
				K	= ;	2+1	=	3	
a li[d] ← k			3		П	[5]	= ;	3	
10 return II									
	-	5							



### **Topic # 95 Knuth-Morris-Pratt Algorithm**

### The KMP Matcher (S, p) $1 \quad n \leftarrow \text{length}[S]$

- 2  $m \leftarrow \text{length}[p]$
- 2 m ( iongui[p]
- 3  $\Pi \leftarrow \text{Compute-Prefix-Function}(p)$
- 4  $q \leftarrow 0$  //number of characters matched
- 5 for i  $\leftarrow$  1 to n //scan S from left to right
- 6 do while q > 0 and p[q+1] != S[i] //if next character does not match
- 7 do  $q \leftarrow \Pi[q]$
- 8 if p[q+1] = S[i] //next character matches
- 9 then  $q \leftarrow q + 1$
- 10 if q = m //is all of p matched?
- 11 then print "Pattern occurs at position"
- 11 q  $\leftarrow \Pi[q]$  // look for the next match

Let us execute the KMP algorithm to find whether 'p' occurs in 'S'.

For 'p' the prefix function,  $\Pi$  was computed previously and is as follows:

q	1	2	3	4	5	6	7
р	а	t	a	t	а	с	а
П	0	0	1	2	3	1	0

i – m+1

Initially: n = size of S = 15

m = size of p = 7

Step 1: i = 1, q = 0 comparing p[1] with S[1]

t	а	С	t	a	t	a	t	а	t	a		2	a	а	t
1				1	1			р	а	t	а	t	a	c	a
а	t	a	t	a	С	а		<b>H</b>	0	0	1	2	3	1	0

P[1] does not match with S[1] 'p' will be shifted to the right as i increments

### Components

- The Prefix Function
- The KMP Matcher

MUHAMMAD IMRAN 137







processing very large files

### Drawback

- > Doesn't work so well as the size of the alphabets increases Conclusion
- > A fast linear time algorithm for string matching

## **96.Scoring Scheme**

### **Scoring System: Introduction:**

• Total score assigned to an alignment is sum of terms for each aligned pair of residues, plus terms for each gap

 $\sum S(x_i,y_j) + d$ 

### d = linear gap penalty

### Simple Alignment Scores:

• A simple way (but not the best) to score an alignment is to count 1 for each match and 0 for each mismatch



### **Scoring or Substitution Matrices:**

- Used for scoring amino acid substitutions in pairwise alignments
- They reflect substitution rates that are originated by evolutionary events

Some of the substitution matrices to compute sequence alignments are:

- PAM: Point Accepted mutations
- BLOSUM: BLOCK Substitution Matrix

### **Substitution Matrices:**

For a set of well known proteins:

- Align the sequences
- Count the mutations at each position
- For each substitution set the score to the log-odds ratio

For each substitution set the score to the log-odds ratio

 $log\left(\frac{observed}{expected by chance}\right)$ 

#### **Positive Score:**

The amino Acids are similar, mutations from one into the other occur more often than expected by chance during evolution

### **Negative Score:**

The amino Acids are dissimilar, mutations from one into the other occur less often then expected by chance during evolution





### Conclusion

• Substitution matrices are the log-odds matrices used for scoring amino acid substitutions in pairwise alignments

### **97.Substitution Matrices**

### Introduction:

• Substitution scores can be derived from probabilistic model

#### **Notations:**

- $\succ$  Let a pair of sequences x and y of length *n* and *m*
- $\succ$  X<sub>i</sub> be the ith symbol in x y<sub>j</sub> be the jth symbol in y
- Symbols are from alphabet A
  - $\circ$  A={A,T,G,C}
- A={twenty amino acids}
- Symbols from alphabet be a,betc

### **Objective:**

Given a pair of aligned sequences, we want to assign a score to the alignment that gives a measure of the relative likelihood that the sequences are related as opposed to being unrelated

### **Unrelated or random Model R:**

Letter a occurs independently with frequency  $q_a$  and the probability of two sequences is the product of probability of each amino acid

 $P(x,y|R) = \mathbf{q}_{xi} \mathbf{q}_{yj}$ 

### Match Model M:

- Aligned pairs occur with a joint probability **p**<sub>ab</sub>.
- $\mathbf{p}_{ab}$  can be thought of as the probability that the residues a and b have been independently derived from some unknown original residue c in their common ancestor.
- The probability of the alignment is

$$P(x,y|M) = \prod_{i} p_{xiyj}$$

- The ratio of two likelihoods can be calculated as;

$$\frac{P(\mathbf{x},\mathbf{y}|\mathbf{M})}{P(\mathbf{x},\mathbf{y}|\mathbf{R})}$$

### **Odds ratio:**

- The ratio of two likelihoods can be calculated as;

$$\frac{P(x,y|M)}{P(x,y|R)} = \frac{\prod_{i} P_{xiyj}}{\prod_{i} q_{xi} \prod_{j} q_{yj}} = \prod_{i} \frac{P_{xiyj}}{q_{xi} q_{yj}}$$

- Logarithm can be used to have an additive score

$$S = \sum S(x_i, y_j)$$

where  $S(a,b) = log(p_{ab}/q_aq_b)$ 

### log likelihood of (a,b) as aligned vs unaligned pair

### **Dayhoff PAM matrices:**

Dayhoff, Schwartz and Orcutt (1978) presented their famous PAM (Point accepted mutations) using substitution data from similar proteins then extrapolating this information to longer evolutionary distances

 $S(a,b) = log(p_{ab}/q_aq_b)$ 

incorporating the evolutionary time

### $S(a,b|t) = \log P(b|a,t)/q_b$

Since  $p_{ab}/q_a = P(b|a)$ 

Values are rounded to near integer for computational convenience

• PAM250 is scaled by 3/log2 to give scores in third-bits

### **BLOSUM matrices:**

- Dayhoff matrices do not capture true difference between short time substitutions and long term ones.
- PAM matrices do not perform well in case of distantly related proteins.
- BLOSUM matrices are derived from set of aligned, ungapped regions from protein families called BLOCKS database (Henikoff&Henikoff 1992).
- Sequences from each block was clustered together with score >L%.
- Matrices with *L*= 62 and *L*= 50 known as **BLOSUM62** and **BLOSUM50** respectively.

BLOSUM62 is good for ungapped alignments and BLOSUM50 is good for gapped alignments

	A	С	D	E	F	G	н -	~
А	4	0	-2	- 1	-2	0	-2	
С	0	9	-3	$^{-4}$	-2	-3	-3(	
D	-2	-3	6	2	-3	-1	-1(	
Е	-1	$^{-4}$	2	5	-3	-2	- 9'	
F	-2	-2	-3	-3	6	-3	5	
G	0	-3	-1	-2	-3	1		
н	-2	-3	-1					
↓					BLC	วรบ	M 6	2

### Conclusions

- Substitution scores can be derived from probabilistic models
- PAM and BLOSUM are famous substitution matrices

# 98.Optimal Algos

### Introduction:

Finding the path whose total score is maximal will give the best sequence alignment

- Two methods
  - Local alignment
  - Global alignment

### **Global Alignment:**

It is an alignment that essentially spans the full extents of input sequences

- Hence it covers the entire length of sequences involved
- The Needleman-Wunsch algorithm finds best global alignment between two sequences

### Local Alignment:

- It only covers parts of the sequences to be aligned
- Smith-Waterman algorithm finds the best local alignment between two sequences

### **Dynamic Programming:**

- Dynamic programming is used to find an optimal alignment of two sequences and its scores
- It is a method by which a larger problem may be solved by first solving smaller, partial versions of the problem
- Three steps in dynamic programming:
  - Initialization
  - Matrix filling (scoring)

### Traceback (alignment

- <u>Initialization</u>: Create matrix with M+1 columns and N+1 rows where M and N correspond to the size of sequences to be aligned
- <u>Matrix filling</u>: Fill the matrix with highest possible score
- <u>Trace back:</u>Move from the last corner and follow the arrow

#### **Conclusion:**

• Dynamic programming is used to find an optimal alignment of two sequences and its scores

## 99.Needleman\_wunch Algos

#### Introduction:

It performs global alignment on two sequences

• The algorithm was developed by Saul B. Needleman and Christian D. Wunsch and published in 1970

Basic idea is

- to build up the best alignment by using optimal alignments of smaller subsequences
- It was the first application of dynamic programming to compare biological sequences

#### Steps:

Three steps

- 1. Initialization
- 2. Matrix filling
- 3. Traceback

#### **Creating the Matrix:**

Initial matrix is created with M+1 columns and N+1 rows

• Where M and N correspond to the length of sequences

#### Initialization:

- The cell of first row and first column of the matrix is initially filled with zero
- Add gap penalty for each shift to the right

### Initialization:

a.	F(0, 0)	= 0
b.	F(0, j)	= - j × d
c.	F(i, 0)	= - i × d

#### Matrix Fill:

- Move through the cells row by row, calculating the score for each cell
- Compute three scores:
  - A match score
  - Vertical gap score
  - Horizontal gap score
- The *match score* is the sum of the diagonal cell score and the score for a match

- The horizontal gap score is the sum of the cell to the left and the gap score
- The vertical gap score is computed analogously

```
Matrix filling: Filling-in partial alignments
For each i = 1.....M
   For each j = 1.....N
                 F(i-1,j-1) + s(x_i, y_i),
                                          [case 1]
   F(i, j) = max
                   F(i-1, j) – d,
                                          [case 2]
                   F(i, j-1) – d
                                          [case 3]
                      DIAG,
                                if [case 1]
                      LEFT,
                                if [case 2]
         Ptr(i,j)
                   =
                      UP,
                                 if [case 3]
```

#### **Traceback:**

- > The final step in the algorithm is the trace back for the best alignment
- Start at the bottom-right corner
- ➢ Follow where maximum value comes from
- F(M, N) is the optimal score, and from Ptr(M, N) can trace back optimal alignment

#### **Scoring Scheme:**

- Scoring scheme introduced can be user defined
- It contains specific scores for match and mismatch residues as well as gap

#### **Conclusions:**

• Needleman and Wunsh Algorithm performs global alignment on two sequences using a dynamic programming approach

### 100.Smith\_waterman Algo

#### Introduction:

Finds the best local alignment between two subsequences

#### Steps:

- Initialization
- ➢ Matrix filling
- ➢ Traceback or alignment

#### Algorithm

- 1. Initialization:
  - a. F(0, 0) = 0
  - b. F(0, j) = 0
  - c. F(i, 0) = 0

Trace-back:

#### Matrix filling: Filling-in partial alignments For each i = 1.....M



#### $F(M,\,N)$ is the optimal score, and from $Ptr(M,\,N)$ can trace back optimal alignment

#### Creating the Matrix:

- Initial matrix is created with M+1 columns and N+1 rows
  - Where M and N correspond to the length of sequences

#### Initialization:

• First row and first column of the matrix is filled with zero

		Т	G	G	Т	G
	0	0	0	0	0	0
Α	0					
Т	0					
С	0				Y	
G	0					
Т	0					

#### Matrix Filling:

		т
	0	0
Α	0	0

- Add the match or mismatch scores diagonally
- Add gap penalties vertically and horizontally
- Replace the negative values by zero

		T	G
	0	0	0
Α	0	0	0
Т	0	5	3

			T	G	G	
		0	0	0	0	
	Α	0	0	0	0	
	Т	0 -	5	3	1	
c	С	0	3	2	0	

		Т	G	G	Т	G
	0	0	0	0	0	0
Α	0	0	0	0	0	0
т	0	5	3	1	0	0
С	0	3	2	0	0	0
G	0	1	0	7	5	5
Т	0	5	3	5	12	10

Match=+5 Mismatch=-3 Gap penalty=-2

#### Traceback:

• Traceback starts with maximum value in the matrix and then go backwards

		Т	G	G	Т	G
	0	0	0	0	0	0
Α	0	0	0	0	0	0
Т	0	5	3	1	0	0
С	0	3	2	0	0	0
G	0	1	0	7	5	5
Т	0	5	3	5	12	10

Alignment:

**Conclusion:** 

Finds the best local alignment between two subsequences

# **Topic 101-102 DNA Sequencing**

Magazine cut into millions of pieces. Problem of fragment assembly in DNA sequencing Short 500 to 700 nucleotide sequences per experiment, Assembling entire genome reassembling the magazine Both problems are complicated by unavoidable experimental error. The data are frequently incomplete. DNA sequencing methods-Fred Sanger and Walter Gilbert. Cells make copies of DNA DNA fragments of different lengths- if one base is missing.



For A and C missing, will also result in DNA fragments by length. Each of four starvation experiments produces a *ladder* of fragments of varying lengths called the *Sanger ladder* 

Sequencing of a 5386- nucleotide virus, DNA sequence data, Human Genome Project, 3 billionnucleotide sequence, DNA sequencing technology, Sequencing reads, Continuous genome, DNA reading process, DNA sequencing machines, Single DNA fragment.

#### Shotgun sequencing

Sonicated, Inserts, Vector, Bacterial host, Cloning proce, DNA sequencing- inserts and computational

# Topic-103 DNA Array

Human Genome Project, Sequencing *by Hybridization*, DNA array (DNA chip), Probes, Hybridization, Weak chemical bond. DNA probes, Biochemical problem and the combinatorial problem *Science*, SteveFodor and colleagues, Light-directed polyme synthesis –similarities to



computer chip manufacturing.

In Fig. An array with all  $4^l$  probes-  $4 \cdot l$  separate reactions Affymetrix-64kb DNA array in 1994 1-Mb or larger Probes- unknown target DNA-*l*-mer composition Universal DNA array contains all 4l probes of length l.



### **Topic-104 Sequencing by Hybridization**

Unknown DNA sequence- length l, String s of length n, the l-mer composition is the multiset of n - l + 1 l-mers in s and is written Spectrum(s, l).

If l = 3 and s = TATGGTGC, then

 $Spectrum(s, l) = \{TAT, ATG, TGG, GGT, GTG, TGC\}$ 

Lexicographic order, like ATG,GGT,GTG, TAT, TGC, TGG

Sequencing by Hybridization (SBH) Problem:

Reconstruct a string from its l-mer composition

Input: A set, S, representing all *l*-mers from an (unknown) string s

**Output**: String *s* such that Spectrum(s, l) = S

### **Topic-105-** Fragment Assembly in DNA Sequencing

500- to 700-bp DNA reads Reconstruct the entire genomic DNA sequence Fragment assembly Error rate in DNA reads Sequencing errors, Assignments to read Major problem-repeats in DNA-

20% of human genomeAlu sequence-million times Repeats occur at many scales Human T-cellreceptor locus-Trypsinogen gene (4 kb).1 million Alu repeats ( 300 bp) and 200,000 LINE repeats (<br/>1000 bp)3 billion-letter sequence500-letter reads, large number of repeats.

# **106. Strategy for Sequencing**

#### **Strategy for Sequencing**

Clone into BAC, Sequenced each one-minigenome, Simplifies the computational assembly problem, Human Genome project, Whole-genome assembly paradigm (James Weber and Gene Myers)Celera Genomics Mate-pair reads sequencing technique.

Inserts of length L- both ends are sequenced, Mates at distance L- length is larger than most repeats

#### Most fragment assembly algorithms consist of the following three steps

Overlap: Finding potentially overlapping reads

Layout: Finding the order of reads along DNA

*Consensus*: Deriving the DNA sequence from layout Best match between the suffix of one read and the prefix of another. Sequencing errors-Dynamic programming algorithm. constructing the layout is the hardest step in fragment assembly. Whole-genome shotgun sequencing. Large number of reads to ensure that experimental errors are reduced to minor noise.

### **Topic-107- Protein Sequencing and Identification**

Routinely sequenced proteins-Frederick Sanger-Nobel prize, Computational problem *Edman degradation reaction* to chop off one terminal a.a, Sanger digested insulin with proteases DNA sequencing "break-read the fragments-assemble"

Edman degradation reaction, 1960s protein sequencing machines were on the market DNA sequencing technology Protein-Obtaining reads-problem Assembly- easy

## **108. Computational Protein Sequencing**

#### **Computational Protein Sequencing**

Proteins produced in cell Sequence of previously unknown proteins. Proteins -biological system and Range-Brain cells-Liver cells

*De novo protein sequencing Protein identification* Gedanken experiment Proteins constitute the DNA polymerase

Protein sequencing and identification, *Spliceosome*, Matthias Mann and colleagues purified the spliceosome complex

## **109. Mass Spectrophotometry**

Programmed cell death, Survival factors, Developing nematode DNA sequence data Nervous system- mutation in several genes, Mass spectrometry.

Mass spectrum of a peptide is a collection of masses of these fragments- Derive the sequence of a peptide given its mass spectrum. For an ideal fragmentation process the peptide sequencing problem is simple. The fragmentation process is not ideal, and mass spectrometers measure mass with some imprecision.

Protein Trypsin ↓ Peptides Tandem Mass Spectrometer↓ Small fragments

-

# **110.111 Peptide Sequencing Problem**

Amass spectrometer breaks a peptide  $p_1 p_2 \cdots p_n$ GPFNA GPFNA GPFNA G, GP, GPF, GPFN PFNA, FNA, NA, A GPFNA into GP and FNA, lose some small parts of GP and FNA, fragments of a lower mass GP might lose a water (H<sub>2</sub>O), and the peptide FNA might lose an ammonia (NH<sub>3</sub>). Mass of GP minus the mass of water (1 + 1 + 16 = 18 daltons), and the mass of FNA minus the mass of ammonia (1 + 1 + 1 + 14 = 17 daltons). Two different ion types

Peptide fragmentation is characterized by a set of numbers  $\Delta = \{\delta 1, \ldots, \delta k\}$  types of ions The set of ion types. A  $\delta$ -ion of an N-terminal partial peptide  $P_i$  is a modification of  $P_i$  that has mass  $m_i - \delta$ , The most frequent N-terminal ions are called *b*-ions (ion  $b_i$  corresponds to  $P_i$  with  $\delta = -1$ ) and the most frequent C-terminal ions are called y-ions (ion  $y_i$  corresponds to  $P_i$  with  $\delta = 19$ )

### **112.Protein identification via Database Search**

De novo protein sequencing algorithms- invaluable for known, unknown proteins. Useful for complete spectra Spectra far from complete, De novo peptide sequencing algorithms. If we had access to a database of all proteins from a genome, then we would no longer need to consider all  $20^l$  peptide sequences to interpret an MS/MS spectrum, but could instead limit our search to peptides present in this database . Database search "the back of a book", Experimental spectrum , Sequence of the experimental peptide, SEQUEST algorithm – John, Yates and colleagues.

#### **Protein Identification Problem:**

Find a protein from a database that best matches the experimental spectrum.

**Input**: A database of proteins, an experimental spectrum S, a set of ion types, and a parent mass m. **Output**: A protein of mass m from the database with the best match to spectrum S

### **Topic-113 Modified Protein Identification Problem**

#### **SEQUEST Algorithm**

Exhaustive search approach

Virtual database

Potential modifications

Combinatorial problem

#### **Modified Protein Identification Problem:**

Find a peptide from the database that best matches the experimental spectrum with up to k modifications.

**Input**: A database of proteins, an experimental spectrum S, a set of ion types  $\Delta$ , a parent mass m, and a parameter k capping the number of modifications

**Output:** A protein of mass m with the best match to spectrum S that is at most k modifications away from an entry in the database

Modified Protein Identification problem  $P_1$  and  $P_2$ -  $S_1$  and  $S_2$ , Notion of spectral similarity Shared peaks count, Limitations in detecting similarities by database search.

## **Topic 114-Protein Structures**

#### Background

Complex protein structures enable proteins to perform complex functions. We know over a million protein sequences but only about 100,000 protein structures.

#### Why only 100,000 proteins for over million protein sequences

Estimating exact protein structures is very difficult. Its difficult to crystallize proteins. Even if we manage to get protein's X-Ray, to reconstruct the structure is extremely complex

#### Introduction

What if we could somehow predict protein structures?

• Since we know so many sequences, they can be used for predicting protein structures. This indeed is possible and helpful.

#### The Basic Idea

- 1. Amino acids determine the protein structure
- 2. We have a large protein sequence dataset (uniprot)

Hence, we can fold protein sequences and predict their structures!

#### Why predict and why not exact solutions?

A deterministic solution of protein folding is a major unsolved problem in molecular biology! Proteins fold spontaneously or with the help of enzymes or chaperones.

#### To predict we must first learn!

To computationally predict protein structures, we need to copy or mimic the natural folding! What are the steps in protein folding and structure formation?

#### To fold we must learn the steps

Step 1: "Collapse"- leading to burial of hydrophobic AA's

Step 2: Fluid globule - helices & sheets form, but are unorganized

Step 3: Compaction, and rearrangement of 2' structures

#### Conclusion

- Protein structure prediction involves learning how the amino acids in primary sequence fold.
  - Using this information, upon getting a protein sequence, we can try to predict how it folds!

### **115. Predicting Secondary Structures**

#### Background

Since the first step in protein folding is the formation of secondary structures, we must evaluate which amino acids in the primary sequence prefer which secondary structures?

By looking at the structures in PDB, we know that Alanine mostly found in Alpha Helices. So if we have several Alanines in the sequence, then we can anticipate that a helix may be formed by them

#### Introduction

What if we survey the entire PDB and check the presence of each amino in each type of secondary structure

• If we know which amino acid is found in which specific secondary structure, then we can use it for prediction!

Amino Acid	Pa	P	P,
Glu	1.51	0.37	0.74
Met	1.45	1.05	0.60
Ala	1.42	0.83	0.66
Val	1.06	1.70	0.50
Ile	1.08	1.60	0.50
Tyr	0.69	1.47	1.14
Pro	0.57	0.55	1.52
Gly	0.57	0.75	1.56

#### Conclusion

- Several algorithms have been designed to predict 2' given an amino acid sequence
- The first such algorithm was the Chou-Fasman Algorithm!
- We will see it in the upcoming modules!

### 116. 2' Structures in Chou Fasman Algorithm

#### Background

- For a primary sequence, and a tentative 2' structure, propensity table can help us compute the overall propensity
- Product of propensity values is computed for overall propensity for each 2' structure

#### Introduction

- An important point to note here is that 2' structures are formed due to hydrogen bonding between amino acids
- So, we need to consider the neighboring amino acids as well!



#### Conclusion

- You only need to compute propensities for a small number 2' structures
- The highest net propensity will be the most probably secondary structure that will be formed!

# 117.121 Chou Fasman Algorithm -

Only a small number of combinations of secondary structures are possible due to their individual properties. Such as 4 amino acids are needed to start an Alpha Helix and 5 amino acids for Beta Sheet **Note that besides the alpha helix and beta sheets, LOOPS are an other secondary structure.** 

How can loops be integrated into predicting 2' structures?

• Loops are small ~ 3-4 amino acids

- 1. Scan through the sequence : E M A V I Y P G
- 2. Identify sequence regions where:
   4 out of 6 contiguous residues give a P(α) > 1.0
  - That region is declared as alpha-helix
  - Extend helix to both sides until 4 out of 6 contiguous residues give a P(α) < 1.0</li>
  - That is declared end of the helix

#### Conclusion

- For Alpha Helices, 4 contiguous amino acids are required
- Their Alpha-Helix propensity should be more than 1.0
- Once this propensity falls below 1.0, Alpha-Helix stops

### **Chou Fasman Algorithm - II**

#### Background

Alpha Helices are formed from 4 contiguous amino acids having an Alpha-Helix propensity over 1.0. The Alpha-Helix stops if this propensity falls below 1.0! Introduction

- Once Alpha Helices are constructed, and concluded, the remaining amino acids can be evaluated for Beta sheets and turns etc
- Let's see how Beta sheets are evaluated using Chou Fasman Algorithm
- Compute P(β) for <u>contiguous regions</u> of 5 Amino Acids
- 2. From these regions, identify regions where:
   5 contiguous residues have P(α) > P(β)
  - That region is finalized as alpha-helix

Repeat this step for the full amino acid sequence to finalize all possible alpha helical regions in the sequence.

#### Conclusion

Alpha Helices can be finalized if their propensity is higher than the propensity for Beta Sheets in regions of 5 amino acids

For those regions where that is not the case, further evaluation is required

#### **Chou Fasman Algorithm - III**

Alpha Helices were finalized if their propensity was higher than the propensity for Beta Sheets in regions of 5 amino acids. For those regions where that are not the case, what should be done?

We can evaluate such regions for Beta Sheets.

Let us see step by stop how to find a beta sheet and how to differentiate them from alpha helices

Scan the sequence to identify regions where: • 3 out of 5 amino acids have P(β) > 1.0

- That region is declared as beta sheet
- Extend beta sheet to both sides until 4 <u>contiguous residues</u> average P(β) < 1.0</li>
- · That is declared end of the beta sheet
- Those regions are finalized as beta-sheets which have average  $P(\beta) > 1.05$  and the average  $P(\beta) > P(\alpha)$  for that region.

#### Conclusion

- Regions where overlapping alphahelices and beta-sheets occur are declared helices if
  - the average P(a-helix) > P(bsheet) for that region
- Else, a beta sheet is declared if • average P(b-sheet) > P(a-helix) for that region
- Using the strategy of higher propensity, alpha helices and beta sheets can be completely resolved
- Assignments for each beta sheet and alpha helix can be finalized
- But what about the loops?

#### **Chou Fasman Algorithm - IV**

#### Background

- After computing the propensity of alpha helices and beta sheets, we need to settle for loops
- Let's see how can we find out the loops using Chou Fasman Algorithm
- For any jth residue in sequence, we calculate f(Total) = f(j) f(j+1) f(j+2) f(j+3) (tetrapeptide)
- If 1. f(Total) > 0.000075
  - the average value for P(turn) > 1.00 in the tetrapeptide.
  - the averages for the tetrapeptide are such P(a-helix) < P(turn) > P(b-sheet),

#### Conclusion

- Chou Fasman Algorithm helps predict Alpha Helices, Beta Sheets and Turns
- The algorithm is based on statistical occurrence of Amino Acids in known structures

### 122.125. Chou Fasman Algorithm – Flowchart I, II & III.

Chou Fasman Algorithm helps predict secondary structures such as Alpha Helices, Beta Sheets and Turns. Step by step flowchart of the entire algorithm. Beta sheets can be predicted from primary amino acid sequences





- Chou Fasman Algorithm helps predict secondary structures from amino acid sequences. Step by step flowchart of the algorithm for extracting Alpha Helices
- Alpha helices, beta sheets and turns can be predicted using Chou Fasman Algorithm. This algorithm is based on statistical analysis of amino acid occurrences in proteins.

#### Chou Fasman Algorithm – Improvements

• Secondary structure propensity values of alpha helix, beta sheet and turns should be recalculated with the latest protein data sets.

#### Special consideration for:

- Nucleation regions
- Membrane proteins
- Hydrophobic domains
- Consider variable coil and loop sizes besides the from tetra peptide turns
- Consider local protein folding environments
- Solvent accessibility of residues
- Protein structural class
- Protein's organism

#### Conclusion

Chou Fasman can be improved to better predict secondary structures by incorporating biochemical factors and updated statistics!

### 126. Summary of Visualization, Classification & Prediction

- A. Why do we need to visualize proteins?
- B. Which atoms are used to reconstruct proteins?
- C. Where are the positions of these atoms stored?

#### **Structure Classification**

- A. What is the relationship between protein structure and function?
- B. What is the need to classify proteins
- C. Hierarchy of classification

#### **Structure Prediction**

- A. Why structure of proteins are important?
- B. Why are so few structures reported till date?
- C. Benefits of predicting structures

#### Conclusion

Structure visualization, classification and prediction equip us to perform functional evaluation of proteins! This is important for understanding disease and designing drugs for treating them

### **Topic-127- Introduction to Homology Modelling**

Proteins are 3D molecules with their own unique structures. Protein structure is reflective of the protein function. Protein structure includes 1', 2', 3' and 4' structures. 1' structure of proteins is the sequence of proteins and can be obtained by mass spectrometry. 2' structures formed by proteins are the helices, beta sheets, loops and coils. 3' structure of proteins is the combination of 2' structures such that the overall protein structure is formed. 4' protein structure is formed when two or more proteins complex together. **X**-Ray Crystallography and NMR Spectroscopy are used to find the structures of proteins. However, these methods are difficult and expensive. Solution: Prediction of structures. Protein sequence gives rise to its structure. If another protein can be predicted based on that similar protein . So, it is then possible to identify unknown protein structures by just examining the homologous protein sequences.

#### Conclusions

- Sequence Identity
- Alignment Length

Which combination of identity and alignment length is suitable for best for structure prediction?

## **128.Homology, Paralogy and Orthology**

In homology modelling, proteins with similar 1' sequences are considered. Given that one of them has its 3' structure known, then the 3' structure of other protein can be predicted





#### Conclusions

• Good sequence alignment and identity ensures that homology modelling will give accurate results

## **129. Workflow of Structural Modelling**

Homology modelling is used to predict structures of proteins having high sequence similarity with other proteins with known structures! **Overall, there are three different strategies for structure prediction** 

- . Homology Modelling
- Threading/Fold Recognition
- Ab Initio Modelling

### 130.135 Seven Steps to Homology Modelling – I

Protein structure can be predicted by 3 methods:

- 1. Homology Modelling
- 2. Fold Recognition / Threading
- 3. Ab Initio Modelling

Let's start by looking at Homology Modelling. There are seven salient steps in any Homology Modelling pipeline. Definition of Template (known) & Target (unknown). Homology modeling of the target structure can be done as follows:

- 1. Template recognition and initial alignment
- 2. Alignment correction
- 3. Backbone generation
- 4. Loop modeling
- 5. Side-chain modeling
- 6. Model optimization
- 7. Model validation



#### Seven Steps to Homology Modelling - II







So, Homology modelling works in seven steps. It is a repetitive process

### **136.MODELLER for Homology Modelling**

- Template recognition and initial alignment
- Alignment correction
- Backbone generation
- Loop modeling
- Side-chain modeling
- Model optimization
- Model validation

Modeller is a software for homology modelling

#### salilab.org/modeller

#### **Inputs:**

Python script file, Sequence alignment & Template (PDB)

<u>log</u> : log output from the run.

- $.B^{\ast}\,$  : model generated in the PDB format.
- .D\* : progress of optimisation.
- .V\* : violation profile.
- .ini : initial model that is generated.

.rsr : restraints in user format.

.sch : schedule file for the optimisation process.

Automated Modelling Servers

Swiss Model http://swissmodel.expasy.org//SWISS-MODEL.html

Robetta

http://robetta.bakerlab.org/

**3D Jigsaw** 

http://www.bmm.icnet.uk/servers/3djigsaw/

Phyre

http://www.sbg.bio.ic.ac.uk/phyre/

Homology modelling helps predict protein structures by using prior structural informationSeveral tools are available to perform homology modelling in a programmatic or automated way!

## **137.139.Fold Recognition – Threading**





Introduction

- A protein fold is defined by the way the secondary structure elements of the structure are arranged relative to each other in space.
- Common folds include 4-helix bundle and the TIM barrel.
- 5,000 stable folds in nature
- Fold recognition: Finding the best fit of a sequence to a set of candidate folds

Fold recognition is also called Threading. Technique for predicting protein structures. Employed when homology modelling cannot predict quality structures. A protein fold is defined by the way the secondary structure elements of the structure are arranged relative to each other in space. Common folds include 4-helix bundle and the TIM barrel. 5,000 stable folds in nature. Fold recognition: Finding the best fit of a sequence to a set of candidate folds.

Fold recognition or Threading is a technique for predicting protein structures. It is useful in cases where homology modelling fails to predict quality structures

The process of threading

- In the process of "Threading", we mount an amino acid sequence on to the backbone of template structures in a folds library
- Each step is "drag" along the sequence (MQVKLFTY...) through each location of each template fold
- Then, for each fold, we must compute the fitness of sequence matching that fold!

Threading involves "passing" the amino acid sequence through each fold in the database. The best match is computed using a scoring function. Combinations of secondary structures come together to form the best prediction. Scoring typically involves using a Z-Score function based on energy of a molecule.



### 140 Online Tools for Threading – iTasser

iTASSER helps thread amino acid sequences on fold and secondary structure databases. It also helps predict function of structures output.

Iterative threading assembly refinement (I-TASSER) server

- Software for automated protein structure & function prediction
  - based on the sequence-to-structure-to-function.
- Steps:
  - · Starts from amino acid sequence
  - i-TASSER first generates 3D atomic models from multiple threading alignments and iterative structural assembly simulations.
  - The function of the protein is then inferred by structurally matching the 3D models with other known proteins.
  - Outputs full-length secondary & tertiary structures and functional annotations on ligand-binding sites
  - An estimate of accuracy of the predictions is provided based on the confidence score of the modeling

## 141. Advantages and Disadvantages of Threading

Fold recognition or Threading is a technique for predicting protein structures. It is useful in cases where homology modelling fails to predict quality structures.

#### <u>Advantages</u>

Threading helps predict secondary structures of proteins towards tertiary structure prediction. For the "Twilight Zone" with low alignment quality and identity, threading is use.

#### **Disadvantages**

Novel proteins cannot be predicted using threading. Fewer than 30% of the predicted first hits are true remote homologues. Validation of each result is necessary.

# 142. 3D-1D Bowie Algorithm

Homology employed high alignment scores. Threading worked by creating combinations of primary sequences and corresponding secondary structures. Proposed **by Bowie et al in 1991**. It converts 3D structure into a 1-D string profile for each structure in the fold library. Align the target sequence to these profiles. 3D-1D methods convert structure and environment information into "profiles". Score for each amino acid is computed for each profile. <u>Inputs and outputs of 3D-1D</u>

- Identify amino acids based on: protein core, side chain positioning, solubility etc. (6 in all)
- Part of secondary structure including  $\Box$ -helix,  $\Box$ -sheet etc (3 in all)
- Total of  $3 \ge 6 = 18$  distinct states
- P<sub>a: j</sub>= prob. of finding amino acid (a) in environment (j)
- P<sub>a</sub>=probability of finding (a) anywhere
- Maximize sum of scores for the fold:

# **143. Introduction to Ab Initio Modelling**

Ab initio methods have Anfinsen's thermodynamic hypothesis at the center. These methods attempt to identify the structure with minimum free energy. Ab initio methods rely on computing the energies of folded proteins. The protein structures with the lowest energy are deemed as plausible predictions.

#### Need for Ab Initio Modelling

- Applicable to any sequence
- Not very accurate biologically
- Accuracy and applicability are limited by our understanding of the protein folding problem

#### Limitation

Computationally expensive

Suitable for proteins with less than 100 residues

## **144.Rationale of Ab Initio Modelling**

Ab initio methods rely on computing the energies of folded proteins. The protein structures with the lowest energy are declared as plausible predictions

#### **Rationale**

Sometimes it so happens that even slightly homologous proteins may not be available. This renders homology modelling and threading/fold recognition as futile . Also, newer protein structures continue to be discovered every day. These could not have been identified by methods which only rely on matching with available structures. Lastly, homology / fold recognition predict protein structures without computing fundamental physical/chemical properties of the mechanisms and driving forces in structure formation. Ab initio methods, in contrast, base their predictions on physical models for these mechanisms. Energy released during the folding process is computed for predicting structure.

## 145.Strategies for Ab Initio Modelling

Ab initio methods base their predictions on physical models of folding mechanisms. Stabilization is measured by energy released during the folding process. Start with an energy function. Fold structures in order to obtain the most stable structure. This structure will have the minimum energy.

#### **Energy Optimization in Ab Initio Modelling**

- 1. Start with a rough initial model.
- 2. Define an energy function mapping structures to energy values. We have to minimize this later!!
- 3. Solve the computational problem of finding the global minimum.

#### Simulation of the Folding Process

- 1. Build an accurate initial model (including energy and forces).
- 2. Accurately simulate the dynamics of the protein folding process.
- 3. The native structure will steadily emerge.

## **146.Energy States of Folded Proteins**

Ab initio methods predict protein structures by folding proteins based on each constituent atom's volume, charge, mass etc. The protein structure reporting lowest energy is selected to be the optimal structure.

#### Energies of Bonded Atoms vs. Nonbonded Atoms

 $V(R) = E_{bonded} + E_{non-bonded}$ 

# **147.Local versus Global Minima**

The protein structure reporting lowest energy is selected to be the optimal structure

#### **Best Case Energy Function**

- Clear energy minimum in the native structure
- Viable path towards this minimum
- Global optimization finds the most stable structure

#### **Optimal Energy Function**

- Easier to design and compute
- Native structure not always at the global minimum
- No clear way of choosing among alternative structures that are generated

### 148. Pros and Cons of Ab Initio Modelling

Native structure not always at the global minimum. No clear way of choosing among alternative structures that are generated <u>Advantages</u>

- Ab Initio methods can fold any target sequence using only physical atomic properties
- Predictions are mostly accurate and correctly describe the natural folding process

#### Disadvantages

- Ab initio methods are the very difficult to design (energy function)
- These methods are slow due to the huge possibilities

## **149.151 Summary of Structural Modelling**

#### **Strategies for Structural Modelling**

- Homology Modelling
- Fold Recognition
- Ab Initio Modelling

#### Homology modeling of the target structure can be done as follows:

- 1. Template recognition and initial alignment
- 2. Alignment correction
- 3. Backbone generation
- 4. Loop modeling
- 5. Side-chain modeling
- 6. Model optimization

7. Model validation

#### **Energy Optimization in Ab Initio Modelling**

- 1. Start with a rough initial model.
- 2. Define an energy function mapping structures to energy values. We have to minimize this later!!
- 3. Solve the computational problem of finding the global minimum.

#### **Simulation of the Folding Process**

- 1. Build an accurate initial model (including energy and forces).
- 2. Accurately simulate the dynamics of the protein folding process.
- 3. The native structure will steadily emerge.

#### Conclusion

- · Homology modelling is performed in cases of high identity and alignment score
- For the "Twilight zone", other strategies are employed
- For low identity and alignment scores, a "Twilight zone" for structure prediction exists
- Fold recognition / threading is useful in such cases
- For cases where even the fold libraries do not give any high scoring matches, Ab Initio strategies can help model the structure
- However, this is a complex and computationally expensive process

# **152.Review of Sequence Analysis**

#### **Important Concepts**

How do we sequence:

Genomes, Proteomes

#### How do we compare sequences:

Pair-wise Sequence Alignment, Multiple Sequence Alignment

#### **Types of Alignments:**

Global Alignment, (Needle Wunsch), Local Alignment, (Smith Waterman)

Advanced Tools:

Fast Alignment (FASTA), Basic Local Alignment Search Tool,

(BLAST) Databases: GenBank, UniProt Online Portals:

Ensemble, Expasy, UniProtKB

# **153.Review of Phylogenetics**

#### Important Concepts

#### **Molecular Evolution**

Insertions, Deletions, Substitutions

#### **Phylogenetic Trees**

Scaled Trees, Unscaled Trees

#### **Phylogenetic Trees**

Rooted Trees, Unrooted Trees

#### UPGMA: Unweighted Pair – Group Method using arithmetic Averages

Two sequences with with the shortest evolutionary distance between them are considered

These sequences will be the last to diverge, and represented by the most recent internal node.

#### **Clustering Vs. Non-clustering Methods:**

UPGMA is a clustering method

Maximum Parsimony etc are non-clustering methods (not included in this course).

## **154.Review of Protein Sequencing**

#### **Important Concepts**

Techniques of protein sequencing, Edman Degradation, Mass Spectrometry, Protein Ionization, Mass Analysis, Protein Fragmentation, MS1, MS2, Estimating and scoring whole protein mass, Extracting & Scoring Peptide Sequence Tags, Searching Post-translational Modifications

#### **Composite Scoring Schemes**

**Online tools:** 

Mascot, Sequest, Prosight PC

### **155 Review of RNA Structure Prediction**

## **156.Review of Protein Structures**

#### **Important Concepts**

#### Protein Structures are generally of four types:

Primary, Secondary, Tertiary, Quaternary

#### Techniques for determining protein structures

X-Ray Crystallography, NMR Spectroscopy

#### **Types of Protein Secondary Structures**

Helices, Beta Sheets, Coils, Loops

• Foundation of structure prediction algorithms Propensities of certain amino acids to form specific secondary structures

#### • Algorithm for predicting protein structures

Chou Fasman Algorithm

Protein Structure Database – PDB, Online tools for predicting structures by using proteins sequences

## **157.**Review of Homology Modelling

#### Four Strata of Protein Structures

Primary, Secondary, Tertiary, Quaternary

#### Justification for homology modelling

Number of known protein sequences is much larger as compared to known proteins structures

#### **Three Strategies for Structure Prediction**

Homology Modelling, Fold Recognition, Ab Initio Modelling, Protein Structure Database – PDB, Online tools for predicting structures such as MODELLER and iTASSER

## **158.Conclusions from this Course**

Definition of Bioinformatics, Need for Bioinformatics, Areas within Bioinformatics, Bioinformatics as an interdisciplinary area, Need to store, process and analyze biological data, Requirement of newer faster algorithms **Specific areas focused were:** 

Comparing sequences, Comparing structures, Predicting structures

#### We looked at:

Algorithms, Databases, Online Tools for each topic.

We studied the basic algorithms for each topic, With evolution and growth of Bioinformatics, newer and better algorithms are now also available!

## **159.Advanced Follow-up Courses**

We looked into the foundations of Bioinformatics, However, each topic that was studied has a undergone a lot of development.

For advanced study in Genomics, you may take "Computational Genomics" course **Topics:** 

Genome Assembly, Gene Finding, Annotation, GWAS etc

- For advanced study in Proteomics, you may take "Computational Proteomics" course. **Topics:** Protein Sequencing, PTM search, Structure Modelling and PPI studies
- For advanced study in Integrative Biology, you may take "Systems Biology" course. Topics: Metabolomics, Transcriptomics, Network Biology etc

#### Also, now there are cutting edge courses on:

Nano-Bio-IT, Computational Drug Design, Personalized Medicine

# **160.Careers in Bioinformatics**

Pakistan as an infrastructure-limited country. The onset of digital revolution. Emergence of data as the most precious commodity, globally. Specifically, health data as a key commodity of the future. Health and disease as the primordial challenge of mankind

• Unique opportunity for us in Pakistan

Bioinformatics requires two things

- 1. Smart mind
- 2. Internet connected computer

#### One man company

You can take public databases and design drugs. One man vs. Roche?

#### BIGDATA

You can make a startup company which manages and process health BIGDATA. All it needs is basic software development skills coupled with Bioinformatics

#### The next disruption

The next Google, Facebook and Uber is going to emerge from Health and Bioinformatics. Pharmaceutical companies are investing into bioinformatics human resource development

#### Jobs Market

Pharmaceutical Giants, Research Centers & Universities, Hospital & Diagnostic IT departments, Your own startup company

# **Topic-161 RNA Structure**

Outline

- RNA folding
- Dynamic programming for RNA secondary structure prediction
- Covariance model for RNA structure prediction

#### Base Pairing

- RNA bases A,C,G,U
- Bases can only pair with one other base
- Canonical Base Pairs
  - A-U
  - G-C
- "wobble" pairing
  - G-U
  - I-U



**Canonical Base Pairs** 

A-U, G-C

• "wobble" pairing

G-U, I-U, I-A & I-C

RNA Types messenger RNA

(mRNA) Non-coding RNA

transfer RNA (tRNA), ribosomal RNA (rRNA), small interfering RNA (siRNA), micro RNA (miRNA), small nucleolar RNA (snoRNA)

Types of RNAs	Primary Function(s)	Types of RNAs	Primary Function(s)
mRNA - messenger	translation (protein synthesis), Coding, regulatory	scRNA - small cytoplasmic	signal recognition particle (SRP), <u>tRNA</u> processing
<u>rRNA</u> - ribosomal	translation (protein synthesis)	snRNA - small nuclear	mRNA processing, poly A
t-RNA - transfer	translation (protein synthesis)	snoRNA - small nucleolar	processing/maturation/me
hnRNA -	precursors &		uryiduon
heterogeneous nuclear	intermediates of mature mRNAs & other RNAs	regulatory RNAs (siRNA, miRNA, etc.)	regulation of transcription and translation, other??



## **162.RNA Secondary Structure**

Some form of RNA can form secondary structures by "pairing up" with itself. This can change its properties dramatically.

#### **Base Pairing**

Pairing of bases helps in determining the secondary structure

Aligning bases, based on pairing with each other gives an algorithmic approach to determining the optimal structure

#### **RNA Folding**

RNA is produced as a single stranded molecule (unlike DNA)

- Strand folds upon itself to form base pairs & secondary structures
- RNA sequence analysis is different from DNA sequence

#### **RNA Structure**

Structures are more conserved than sequences •

Covariation

#### Secondary Structure representation

2D, Circle plot, Dot plot, Mountain, Parentheses, Tree model

#### **Tertiary Structures**



#### Pseudoknots

Pseudoknots cause a breakdown in the Dynamic Programming Algorithm. In order to form a pseudoknot, checks must be made to ensure base is not already paired – this breaks down the recurrence relations

#### Conclusions

Some forms of RNA can form secondary structures by "pairing up" with itself. This can change its properties dramatically.

## **163.RNA Secondary Structure Prediction**

There are different approaches to predict secondary structure of RNA

Energy minimization

Comparative sequence analysis

- Folding and alignment
- Base-Pair Maximization

#### **1. Energy minimization**

#### Dynamic programming approach

Does not require prior sequence alignment. Require estimation of energy terms contributing to secondary structure

#### Assumptions;

Energetically most stable structure is more likely structure. Energy associated with any position is only influenced by local sequence and structure. Neglect pseudoknots

#### Approach

Energy minimization algorithm predicts secondary structure by minimizing the free energy ( $\Box G$ ).  $\Box G$  calculated as sum of individual contributions of:

- Loops
  - stacking

#### **Energy minimization**

Thermodynamic Stability

Estimated using experimental techniques

- Theory : Most Stable is the Most likely
- No Pseudknots due to algorithm limitations
- Uses Dynamic Programming alignment technique
- Attempts to maximize the score taking into account thermodynamics
- MFOLD and ViennaRNA

#### Drawbacks

Compute only one optimal structure. Usual drawbacks of purely mathematical approaches



#### 2. Comparative sequence Analysis

#### Need a multiple sequence alignment as input.

Requires sequences be similar enough (so that they can be initially aligned), Sequences should be dissimilar enough for covarying substitutions to be detected, comparative analysis produces accurate structure predictions.

# 164.RNASeq

Calculating transcript abundance and prevalence by Ultra high throughput cDNA sequencing (Mortazwi et al, 2008)

The sequence reads are individually mapped to the source genome and counted to obtain the number and density of reads corresponding to RNA from each

- known exon
- splice event
- new candidate gene

#### Procedures

Isolation of all mRNA, Convert to cDNA using reverse transcriptase, Sequence the cDNA, Map sequences to the genome.

The more times a given sequence is detected, the more abundantly transcribed it is.

If enough sequences are generated (> 40 Million), *a comprehensive and quantitative view* of the entire transcriptome of an organism or tissue can be obtained (Mortazvi et al, 2008)

#### Data analysis

Mapping reads, Visualization Genome browser, De novo assembly, Quantification, Differential Gene Expression, Functional Analysis, Gene Networks

In RNASeq, transcript abundance and prevalence is calculated using Ultra high throughput cDNA sequencing

## **165.RNASeq Normalization**

Sequencing reactions may vary across different sequencing plate-forms as well as within different lanes of the same sequencer. Transcript lengths also vary

Raw read counts may vary

#### **RNASeq challenges**

Uniformity of sequence coverage, Quantity of sequence required to reliably detect RNAs of lower abundance classes, Quantification and conversion of relative quantification to absolute RNA concentrations, Transcriptomes of organisms with large genomes, containing genes with more complicated structure, present some special challenges

#### **Mapping Biases**

Read counts will be higher if sequencer produces more reads. Longer genes will have the probability of mapping more reads than smaller ones

#### **RPKM** (reads per kilobase of transcript (or exon model) per million mapped reads)

Method for quantification of transcript levels. **RPKM** measure of read density reflects the molar concentration of a transcript in the starting sample by normalizing for RNA length and for the total read

number in the measurement. This facilitates transparent comparison of transcript levels both within and between samples

#### RPKM

Number of reads mapped per gene length in KB per total reads in that sample in millions

C = Count of Mapped Reads L= Length of transcript M = Mapped reads of sample-RPKM = C L/1000 \* M/1000000 RPKM RPKM = C L/1000 XM/1000000 RPKM = CX 10<sup>9</sup> LM Example: What is the RPKM for a transcript of length 2500KB, with 900 alignments in a sample of 10000000 reads out of which 8000000 reads mapped? RPKM = 900X 10<sup>9</sup> 4.5 2500X8X10<sup>6</sup> **RPKM** How many reads are required to map at 1 RPKM with a transcript of 2Kb length from a total of 40 Million Mapped reads? RPKM= X 109  $\times \mathbf{M}$ CX 109 2000X40X10<sup>6</sup> L 80 C= 2000X40X106 109 FPKM (Fragments per kilobase of transcript per million mapped reads) Paired end RNASeq experiments produce two reads per fragment FPKM counts fragments not the reads Both reads might not map Counting reads might doubble count some fragmnets (Trapnell et al 2010) **RPM** While comparing the same genes expression across different samples (treatments), normalizing for gene may not be necessary C RPM = M/1000000 Can compare relatively bigger numbers and get more DEG RPKM reflects the molar concentration of transcript in starting sample normalized for

• Length of RNA

• Total reads in the sample

It facilitates transcript comparison within and across samples Topic

### **166.Neural Network**

The human brain can be described as a biological neural network an interconnected web of neurons transmitting elaborate patterns of electrical signals A neural network is a "connectionist" computational system. Information is processed collectively in parallel throughout a network of nodes. Complex adaptive system.

- Learning processes in biological systems.
- Learning as an optimization process.
- Learning by modification of synaptic strength.

### **167.Association Rule Mining**

It is an important data mining model studied extensively by the database and data mining community. Assume all data are categorical. No good algorithm for numeric data. An association rule has two parts

- an antecedent (if)
- a consequent (then)

Antecedent is an item found in the data. A consequent is an item that is found in

combination with the antecedent Market basket transactions: t1: {bread, cheese,

milk} t2: {apple, eggs, salt, yogurt}

tn: {biscuit, eggs, milk} Concepts:

An item: an item/article in a basket I:

The set of all items sold in the store A

transaction:

Items purchased in a basket; it may have TID (transaction ID) A

transactional dataset:

A set of transactions

## **168.Clustering**

Clustering is "a process of organizing objects into groups whose members are similar in some way" A cluster is therefore a collection of objects which are "similar" between them and are "dissimilar" to the objects belonging to other clusters.

- Simplifications
- Pattern detection
- Useful in data concept construction
- Unsupervised learning process
  - Hierarchical agglomerative general algorithm
- Find the 2 closest objects and merge them into a cluster

- Find and merge the next two closest points, where a point is either an individual object or a cluster of objects
- If more than one cluster remains, return to step 2

#### Applications

- For administrative purposes
- Hospital activity and performance
- Used for researchers Data used by physician
- For laboratory use etc.

## **169.Machine Learning**

Programming computers to optimize performance criterion using example data or past experience

When to learn? Calculate payroll, Solution needs to be adapted to particular cases (user biometrics)

#### When To Learn

Human expertise does not exist (navigating on Mars), Humans are unable to explain their expertise (speech recognition), Solution changes in time (routing on a computer network)

#### Model

Build a model that is a good and useful approximation to the Data

**KDD** is the non-trivial process of identifying valid, novel, potentially useful, & ultimately understandable patterns in data

#### Applications

Retail, Finance, Manufacturing, Medicine, Telecommunications, Bioinformatics, Web mining

Retail: Market basket analysis, Customer relationship management (CRM)

Finance: Credit scoring, fraud detection

Manufacturing: Optimization, troubleshooting

Medicine: Medical diagnosis

Telecommunications: Quality of service optimization

Bioinformatics: Motifs, alignment

Web mining: Search engines

#### **Machine Learning**

Study of algorithms that Improve performance at some task with exp. Role of Statistics, Role of CS

#### **Applications of ML**

Speech recognition, NLP, Computer vision, Medical outcomes analysis/Computational biology, Robot control

## **170.ML Concepts**

- Association Analysis
- Supervised Learning
  - Classification
  - Regression/Prediction
- Unsupervised Learning

#### Reinforcement Learning

#### **Learning Associations**

Basket analysis:

 $P(Y \mid X)$  probability that somebody who buys X also buys Y where X and Y are products/services.

Example: P ( chips | Beer) = 0.7

TID	Items
1	Bread, Milk
2	Bread, Diaper, Beer, Eggs
3	Milk, Diaper, Beer, Coke
4	Bread, Milk, Diaper, Beer
5	Bread, Milk, Diaper, Coke

### Machine Learning

#### Classification

Samings

θ,

- Example: Credit scoring
- Differentiating between low-risk and high-risk customers from their *income* and savings Model



Discriminant: IF *income* >  $\theta_1$  AND *savings* >  $\theta_2$ THEN low-risk ELSE high-risk



FR: Pose, lighting, occlusion (glasses, beard), make-up, hair style

Character recognition:

Speech recognition:

Medical diagnosis: From symptoms to illnesses

Web Advertizing

Retail: Market basket analysis, Customer relationship management (CRM)

Finance: Credit scoring, fraud detection

Manufacturing: Optimization, troubleshooting

Medicine: Medical diagnosis

Telecommunications: Quality of service optimization

Bioinformatics: Motifs, alignment
Web mining: Search engines

## **Machine Learning**

**Prediction: Regression** 

- Example: Price of a used car
- x : car attributes
  y : price

 $y = g(x \mid \theta)$ 

 $\theta$  parameters



## **171.ML Applications**

#### **Supervised Learning**

- Prediction of future cases
- Knowledge extraction
- Compression
- Outlier detection

### **Un-supervised Learning**

Learning "what normally happens"

No output

#### Clustering: Grouping

similar instances

### Applications

- Customer segmentation in CRM
- Image compression
- Bioinformatics

#### **Reinforcement Learning**

Policies: what actions should an agent take in a particular situation

Utility estimation: how good is a state ( $\rightarrow$ used by policy)

No supervised output

Delayed reward

Credit assignment problem (what was responsible for the outcome)

Applications:

Game playing Robot in a maze Multiple agents, partial observability, ...

## **172.Forensic Science**

#### Forensic science

Study & application of science to matters of law Methods

Associations between people, places, things & events involved in crimes.

Crime investigation & Criminal Identification.

## Iris Recognition.

Face Recognition.

Thumb printing.

Finger Printing.

#### Limitations of physical analysis

Ever one has unique iris pattern, Finger prints, Thumb prints. We can get the fingerprints and thumb prints from the crime scene Fingerprints can be

changed by cutting or burning of finger/thumb.

## Limitations of Thumb

## Limitations of FR

Using fingerprints require finger or the thumb, no other parts of body can be used and criminal can use gloves and no prints are there for investigation.

Face Recognition can only be used when the that you get a print from the photographs of the crime scene has been taken or suspicious person has been arrested and victim tells the physical appearance of criminal

## 173.Advantages of bioinformatics in forensic

## DNA Finger Printing / DNA Profiling:

It is a form of forensic identification that is used primarily to identify people.

## **Advantages**

Even though all humans share 99.9% of their DNA sequences, the remaining 0.01% of sequences isunique enough to differentiate people. Because DNA is in every cell of a person's body is present and same, any part of a human's body including dead skin skills, hair, saliva, and more contain DNA sequences.

# 174. .Methods used in DNA profiling.

## **DNA finger printing**

A forensic scientist will need two pieces of DNA to be compared. For example, DNA discovered at crime scene should be compared to a DNA sample taken from a suspect.

### Steps

- Comparison of repeating DNA
- sequences
- Match b/w two samples

## **DNA extraction from cell**

## 1. Collecting cell from sample:

Two meters of DNA in cellCollect cells from the sample with buccal swab.Place the swab into Eppendorf tube.

### 2. Burst cells open to release DNA:

Add the lysis solution to the tube to separate the cells.LS breaks Cell membrane& nuclear envelope causing cells to burst open& release DNA. It also removes histones proteins from DNA.

## 3. Separate DNA from proteins and Debris:

Cells have stayed in warm to for such a time that DNA is freed from cells.Salt causes proteins & other cellular debris to clump together. Place tube into micro centrifuge. Inside the centrifuge tube spin around and debris &heavy proteins sink in bottom of tube and DNA strands remains distributed throughout liquid.

#### 4. Isolate concentrated DNA:

Add the liquid containing DNA in separate tube Now add isopropyl alcohol to tube.DNA is not soluble in this alcohol so it comes out and it can be seen with naked eye.DNA is collected at bottom tube after placing in centrifuge.

#### 175. DNA Profile Encrypted sets of numbers that reflect a person's DNA makeup, Variable number tandem repeats(VNTRS). Short term tandem repeats (STR) in making the DNA profile of a person. These DNA profiles are the basis of a national DNA databases. **DNA profiling processes RFLP** analysis. PCR analysis. STR analysis. Amp FLP. Y-chromosome analysis. **Restriction Fragment length Polymorphism (RFLP)** Process Works RFIP How the ©2008 HowStuffWorks Sponge С Gel Nitrocellulose Paper paper towels

6 A photographic film laid on top of the paper is overall on top obe of the paper is exposed by the radioactivity in the bond probe 4 to form an image corresponding to the DNA bands. aper is exposed to a solution containing radioactively-labeled probe

Alkaline solution is gel to a sheet of nitro-cellulose laid on the top of it, transferring the DNA to the paper.

Alkaline solution

RFLP

Paper blot

в

added

restriction

DNA Samples

with

restriction produce fragments.

It analyzes the length f strands of DNA that include repeating base pairs (VNTRs).

2 Electrophoresis

Plastic bag

fragments. Each sample forms a characteristic pattern of bands.

Repeated sequence of human genome can be same but the number of times it is repeated is unique to everyone.

RFLP analysis requires investigators to dissolve DNA in an enzyme that breaks the strand at specific points. The number of repeats affects the length of each resulting strand of DNA. Investigators compare samples by comparing the lengths of the strands.

Example: CAT is repeated continuously 13 times in a row. In somebody else, it might be 12 times or 14 or whatever.

Limitation: RFLP analysis requires a fairly large sample of DNA that hasn't been contaminated with dirt.

## **176. DNA Profile Methods**

#### **Polymerase Chain**

#### Reaction (PCR)

Replicate a small amount of DNA to create a larger sample for analysis. First, a heat-stable DNA polymerase -- a special enzyme that binds to the DNA and allows it to replicate -- is added. Next, the DNA sample is heated it to 200 degrees F (93 degrees C) to separate the threads. Then the sample is cooled and reheated. Reheating doubles the number of copies. Process is repeated about 30 times, there is enough DNA for further analysis.

#### Analyzing STRs

PCR is the first step in analyzing STRs (Short Tandem Repeats), which are very small, specific alleles in a variable number tandem repeat (VNTR).

#### Short Tandem repeats

Analyzing STRs is more accurate than the RFLP technique because their small size makes them easier to separate. If you want to create a fingerprint, you might look at 20 different STRs at different places in order to create a profile.

#### STRs

It is impossible for two persons to have same number of STR repeated in a given sequence.

#### Y-chromosome Analysis

STRs in Y-chromosome Useful if the sample has mixed DNA. Gender analysis cases. It is processed just like simple STR analysis.

#### AmpFLP

Amplified fragment length polymorphism, is another technique that uses PCR to replicate DNA. Like RFLP, it first uses a restriction enzyme. Then, the fragments are amplified using PCR and sorted using gel electrophoresis. Can be automated, Doesn't cost very much., DNA sample must be high quality otherwise errors may result, which is the case with most DNA analysis techniques.

## **Topic 177- Introduction to Drug Discovery**

Drug Discovery

Primary objective -

design & discovery of new compounds that are suitable for use as drugs

#### A team of workers-

chemistry, biology, biochemistry, pharmacology, mathematics, medicine & computing ...

#### requirement

- (i) Synthesis of the drug
- (ii) Administration method
- (iii) Development of tests
- (iv) Procedures to establish how it operates in the body
- (v) safety assessment
- (vi) research into the biological and chemical nature of diseased state.

#### **Drugs: Definition**

Chemical substances that are used to prevent or cure diseases in humans,

animals and plants

Activity: Pharmaceutical/pharmacological effect on the subject, e.g. Analgesic or  $\beta$ -blocker

Potency: quantitative nature of the effect

**Drugs Properties: ADMET** 



**Drug:** agent used for the psychotic effect by the media or general public. Even the drugs abused have their **activity**. No drug is completely safe. Suitable quantity to cure or excess to be poisonous! E.g. aspirin, paracetamol can be toxic if excesses.

# **178. Drug Discovery Applications:-**

#### **Areas Influencing DD**

- Molecular Biology on Drug Discovery
- High-Throughput Screening
- Combinatorial Chemistry

#### **Molecular Biology Influence**

#### Genetic information

Biochemical and chemical terms. Cloning and expressing genes that encode therapeutically useful protein

#### **High Throughput Screening**

Widely used in the pharmaceutical industry. Automation to quickly assaythe biological or biochemical activity of a large number of drug-like compounds.

#### **Combinatorial Chemistry**

Laboratory technique in which millions of molecular constructions can be synthesized and tested for biological activity.



# **179.Pharmacogenetics**

It is the branch of pharmacology concerned with the effect of genetic factors on reactions to drugs.

How people respond to medicines, Correlating heritable genetic variation to drug response.

#### **Defination:-**

Biotechnological science combines techniques of medicine, pharmacology & genomics which developing drug therapies to compensate for genetic differences in patients which cause varied responses to a single therapeutic regimen.



### Applications:-

- 1. Detection of genetic variability of drug effects on the genome level
- 2. Agent selection
- 3. Analysis of drug
- 4. reactions and drug toxicity on gene expression
- 5. Development of new indications for already approved drugs
- 6. Discovery of new drug targets
- 7. Identification of (non) responders in clinical trials of phase I-IV
- 8. Identification of genotype dependent adverse drug reactions
- 9. Identification of individuals at risk for severe adverse drug effects

## **180.Pharmacogenomic applications**

How genes affect persons response to drugs. Pharmacology (science of drugs). Genomics (the study of genes and their functions). Develop effective, safe medications & doses tailored to a person's genetic makeup.



efficacy trials.

## **Future**

Blessing in research. As a simple example, for nearly a decade the ability to store more information on a hard drive has enabled us to investigate a human genome sequence cheaper.

# 181.182. Drug Discovery – Pipeline

Target Identification, Target Validation, Lead Identification, Lead Optimization, Pre-Clinical

• Pharmacology & Toxicology





#### Methods for DD

- ✓ Random Screening
- Molecular Manipulation
- Molecular Designing
- Drug Metabolites Serendipity

#### **Random Screening**

Higher/crude plants, opium, senna, reserpine, etc. Penicillin microorganism. Antibacterials with improved therapeutic profiles.

#### **Molecular Manipulation**



#### Drug Metabolism

Xenobiotic metabolism Biochemical modification of pharmaceutical substances or xenobiotics by living organisms, usually throughspecialized enzymatic systemsLipophilic chemical compounds into more

readily excreted hydrophilic products.Rate of metabolism determines duration & intensity of a drug's pharmacological action

#### Serendipity

Prototype psychotropic

drugs

Development of psychiatry

Finding of one thing while looking for something else

# **184.Biomedical annotated corpora**

#### Biomedical researchers are interested

- 1. Understanding data
- 2. context information
- 3. background knowledge
- 4. curated databases &
- 5. Literature extensive

#### What is Annotated Corpora?

Dataset for extraction of disease/treatment entities relations. Corpora are usually constructed for training or evaluation purposes during the development of particular system

#### **Annotation Consumers**

The linguistic community typically uses annotation as training data or for specific tasks. An abundance of tools that can *produce* annotations in the specific format of those resources. Biomedical annotation typically used for gene set enrichment analysis



Bio-databases, controlled vocabularies and bio-ontologies encode small fraction of information

#### Linking text to dbs and ontologies

Curators struggling to process scientific literature. Discovery of facts & events crucial for gaining insights in biosciences: need for text mining

## **185.Steps for Creation of Biomedical Corpora**

- ✓ Nature of data
- ✓ Standard datasets
- ✓ Formalism
- ✓ Users of dataset
- Evaluation measures

**Data Mining** 

- ✓ Discovers unsuspected associations
- ✓ Combines & links facts
- ✓ and events
- ✓ Discovers new knowledge, finds new associations

**IR:** yields all relevant . Corpora; gathers, selects, filters documents that may prove useful , Finds what is known

**IE:** extracts facts & events of interest to user, Finds relevant concepts, facts about concepts. Finds only what we are looking for

## **Annotation & Information Extraction**



- IE systems can be developed by referencing annotated corpus.
- The performance of IE systems can be evaluated by being compared to the annotated corpus.

#### **Text Mining Pipelines**

- ✓ Text documents
- ✓ Retrieval/storage Indexaccess relevant storage
- Text Processing: word Filters, Pattern filters, Lexicon matching, Ontology, NLP parsingetc, ...

#### **Feature Extractions:**

Statistical: Word counts, pattern extraction & counts, etc

Domain-specific: Gene Name counts, etc

NLP-specific: Phrase counts, etc

Data Mining: Classification, Clustering, Association, Statistical Analysis, Visual Analysis,

etc ...

# **186.Target Discovery Strategy**

#### Target discovery to clinical application saga

- Physiology-based approach
- Target-based approach
- 1. Physiology-based approach

Is a disease-centric approach in which target is not identified, multiple targets are involved. In vivo screening is done by using drugs, siRNAor antisense oligonucleotides.

This process relies on disease, not on the target.



#### 2. Target-centric approach

Target based discovery starts with the identification of genes and their protein products. Aim to develop drugs affecting one gene or a molecular mechanism. The identification of diseaserelevant genes in vitro cellular models has been possible due to several tools. Gene-suppression tool used to linked the genes with disease.

Target has two types

- 1. Genetic
- 2. Mechanistic

Genetic targets are represented by genes and genes products. Mechanistic targets include mechanism

based targets such as receptors, enzymes or genes, identified on the basis of the disease state.



## **187.Strategies To Identify Possible Drug Targets**

First step of drug discovery is to identification of disease-associated targets. Genome sequencing and

screening have enhanced opportunities for target identification and lead optimization.

### Structure-based Target Discovery

It helped in defining the contours of the cognate surfaces of ligands and their protein targets, permitting optimization of their potency and selectivity. Some drugs that are originated from structure-based approach:

Dorzolamide

Captopril

- Imatinib
- Zanamivir

The protein structure contributes in the following fields:

- Target identification from sequence structure homolog recognition.
- Structural genomics and drug targets.
- Identification of ligand binding region
- Identification of hits and leads
- Structure-guided design and screening
- Kinase drug discovery & Kinome

#### **Target Discovery through Cell-based Genetics**

- New drugs are based on target identification.
- It wants the thorough knowledge of the disease processes and characterization of genes.
- Combination of three target discovery will provide the desired result.

#### **Target Discovery Strategies**

Target discovery strategies based on

- Expression profiling Proteomic approach to identify disease related genes based on differential EP, homology and post translational modification
- Biochemical and cell biological assays To identify genes and proteins linked with disease pathways
- Cell-based genetics Leads to the discovery of targets by disturbing gene function in whole organisms, correlation with phenotypes.

#### **Cell-based Genetics**

Cell-based assays may lead to the identification of genes involve in cellular transformation, activation, migration and a host of biological processes relevant to a human disease.

#### Genetic-based Target Identification It

has some methods:

- Positional Cloning- Laboratory technique used to locate the position of a disease associated gene along the chromosomes.
- Candidate gene approach

To identify complex disease-linked genes through SNP markers. 10 million in HGP and 3 million identified.

#### Target class genetic approach

- Is applied to drug target gene families such as proteases, ion channels and GPCRs.
- 24000 protein coding genes & 2400 DTs
- Best candidate are selected from gene family for genetic analysis.

## 188. Target Validation

- A potential target is identified in the context of a specific disease.
- A validated target is the one that can be manipulated with drugs to produce positive clinical effects in humans.

#### **Requirements for TV**

→ Genetic Approach

Gene to disease correlation in animal model

- Forward Genetics
- Reverse Genetics

#### **Target Validation Tools**

Helps in assessing their genetic association with disease. The following tools are:

- Antisense agents
- Ribozymes

- Peptide nucleic acids
- Transcription factors
- Gene knockouts

#### Antisense Technology

- Drugs as molecules action cellular proteins (enzymes, receptors).
- Downstream: Block events at nuclear/ribosomal levels.
- Prevent expression of aberrant proteins at earlier stage.
- Oligonucleotide reagents

#### Ribozymes

- Small RNA mol. cleave other RNA mol. at sequence specific sites.
- Hairpin Ribozyme: GUC

HammerHead: NUH, H is AUC

- Mode of Delivery: Exogenously , Endogenously
- Peptide NAs: block protein translation
- Transcription Factors: Zinc finder proteins high Affinity to bind to correct region of DNA. Gene Knockouts:

## **189.Lead Identification**

- Compound for biological activity on target.
- Potency threshold
- Libraries of molecules

#### Lead Identification – Technologies

#### Virtual Screening:

Protein structure , docking Chemical similarity search, Knowledge of compounds against receptor, receptor structure & receptor ligand interactions



#### **Visual Screening**

MLCC: Multilevel chemical Compatibility scoring

- Top selling drugs
- Compounds under biological scrutiny
- Anticancer drugs
- Compounds with poor drug like characters

## Chemoinformatics



#### **Pharmacophore Mapping**

Identify lead compounds against a desired target

Definition: 3-D arrange... Usage:

interaction of receptor & legend

DB concept

• QSAR

Quantitative structure activity relationship

• SAR:

synthesizing & testing a series of structurally related compounds

Least squares & KNNs

### High Throughput Docking

- Ligand & protein
- Docking algos
- Force fields, knowledge based & empirical

#### NMR based screening

- Nuclear Magnetic
- Resonance
- 3-D potential DC & tertiary structure of Proteins
- Need of prior information
- SHAPES
- WaterLogsy



#### **Chemical Genetics**

• Gene-product function in cellular or organismal context using exogenous ligands

- Knockouts
- Cell cycle arresting agents

# **190. Lead Optimization**

- Optimize the desirable traits of the lead
- Lead should be amenable for chemistry optimization
- Methods from Lead Identification

#### Lead Optimization – Technologies

- Medicinal chemists conduct extensive SARs to improve potency and selectivity.
- Improve physicochemical and drug-like properties
- Best molecules are advanced to animal Models & preliminary toxicology

#### LO Methods

De novo drug desing

Charge distribution, liphophilicity or pka of side chains and H-bonds donors and acceptors

### SBDD

Structure based drug Design. Effective: 3-D structure of inhibitor with target known. Large no of medicines. Molecular recognition in protein ligand complexes

### **Drug Like properties**

- ADMET in phase 1
- Filters
- Bioavailability, PK, CNS

### Pre- Clinical Pharma-cology & Toxicity

- Animals testing
- Xenograft models
  - ADME/T testing and validation