

# CAP 5510: Introduction to Bioinformatics

# CGS 5166: Bioinformatics Tools

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[www.cis.fiu.edu/~giri/teach/BioinfS08.html](http://www.cis.fiu.edu/~giri/teach/BioinfS08.html)

# Overview of Courses

- Sequence Alignment; Multiple Sequence Alignment
- Sequence Analysis
- Sequencing and Mapping
- Phylogenetic Analysis
- Gene prediction techniques
- Pattern discovery techniques
- Protein structure alignment and analysis
- Genomics, Functional Genomics, Proteomics
- Gene Expression Data Analysis
- RNA Secondary structure
- RNA interference and small RNA
- Ribozymes and Riboswitches
- Databases & Software Packages
- Statistics for Bioinformatics
- Computational Learning & Predictive Methods
- Biomedical Image Analysis
- Emerging Biotechnologies

# Software Packages

- Databases (**GenBank, SwissPROT**)
- Programming Environments (**BioPerl**)
- Sequence Alignment (**BLAST, CLUSTALW**)
- Phylogenetic Analysis (**CLUSTALW, Phylip, PAML**)
- Learning Methods (**HMMPro, GeneCluster, ASOM**)
- Pattern Discovery Techniques (**GYM, TEIRESIAS, APRIORI**)
- Molecular Structure Analysis (**DALI, RASMOL, SPDBV**)
- Microarray Analysis (**CLUSTER, GeneCluster, TreeView**)
- Statistical Software Packages (**SAS, R**)

# Genomic Databases

- Entrez Portal at National Center for Biotechnology Information ([NCBI](#)) gives access to:
  - Nucleotide ([GenBank](#), [EMBL](#), [DDBJ](#))
  - Protein ([PIR](#), [SwissPROT](#), [PRF](#), and Protein Data Bank or [PDB](#))
  - Genome
  - Structure
  - 3D Domains
  - Conserved Domains
  - Gene; UniGene; HomoloGene; SNP
  - GEO Profiles & Datasets
  - Cancer Chromosomes
  - PubMed Central; Journals; Books
  - OMIM
  - Database Neighbors and Interlinking

# Evaluation

- Semester Project (50 %)
- Homework Assignments (20 %)
- Exams (25 %)
- Class Participation (5 %)

## Course Homepage

[www.cis.fiu.edu/~giri/teach/BioinfS08.html](http://www.cis.fiu.edu/~giri/teach/BioinfS08.html)

- Lecture notes, required reading material, homework, announcements, etc.

# Introduction

## 1. What is Bioinformatics?

- Analysis of biological data with computing & statistical tools.

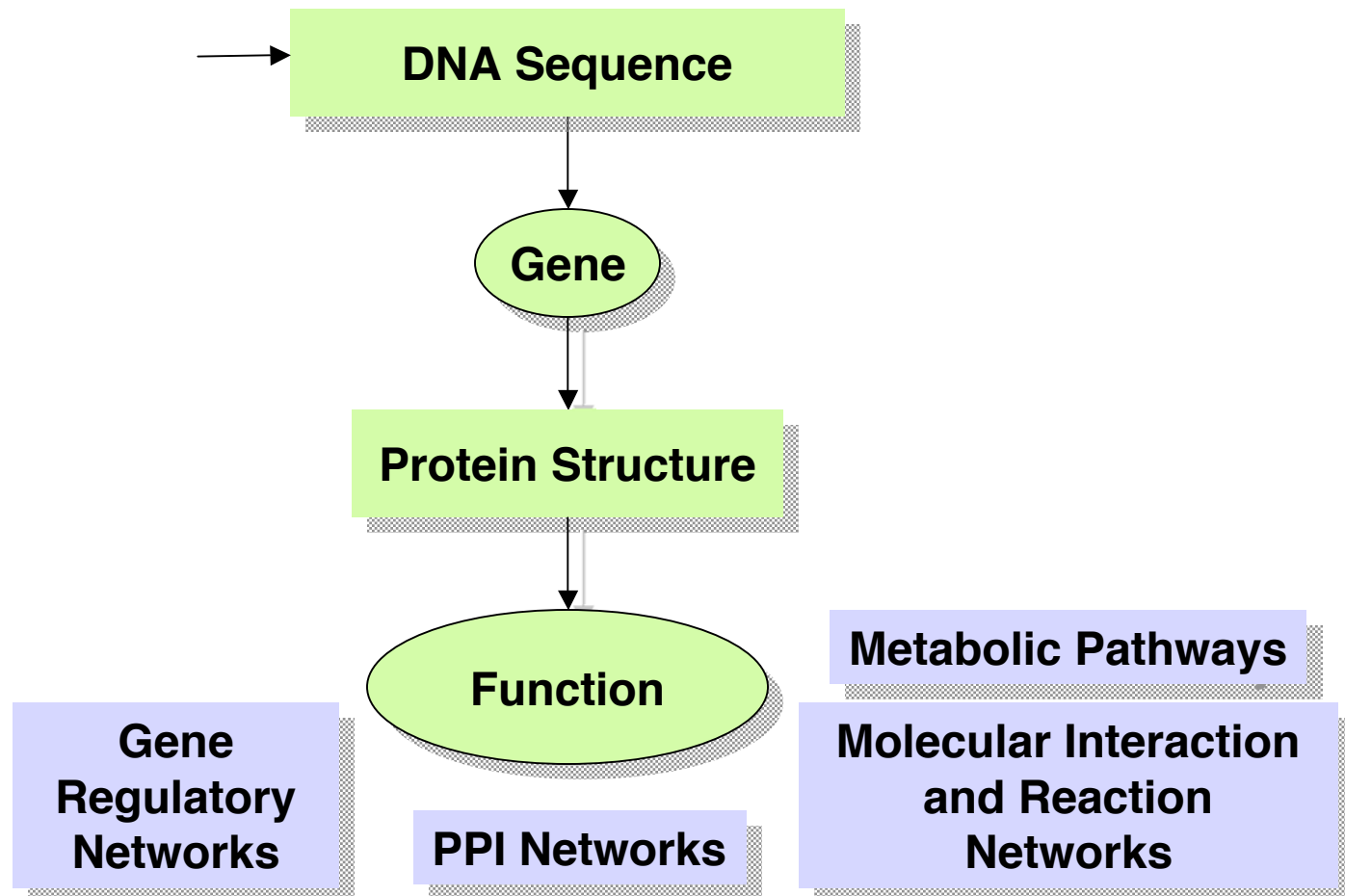
## 2. The different aspects of Informatics?

- Data Management (Database Technology, Internet Programming)
- Analysis/Interpretation of Data (Data Mining, Modeling, Statistical Tools)
- Development of Algorithms/ Data Structures
- Visualization and Interface Design (HCI, Graphics)

## 3. How to assist biological research?

- propose new models or correlations based on data from experiments
- verify a proposed model using known data
- propose new experiments based on model or analysis
- use predicted information to narrow down search in a biological investigation

# Overall Goals



# General Information

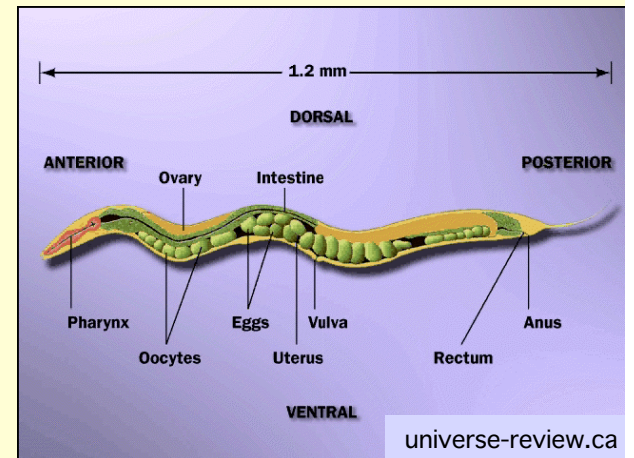
- GenBank Release 157/163 (Dec 2006/7) contains over 64/80 million sequence entries totaling over 83 Gb from over 2,500 organisms  
[<http://www.ncbi.nlm.nih.gov>] (Storage: ~150 GB uncompressed)
- Human Genome has ~3 billion bp with 32,000+ genes.
- 435/624 complete microbial genomes sequenced (684/914 more in progress)
- 2540 Viral genomes (300bp - 300Kb) (1<sup>st</sup> 1978: Simian virus; 5Kb).
- 22 complete eukaryotic genomes sequenced (175 more in progress):  
*Caenorhabditis elegans, Arabidopsis thaliana, Saccharomyces cerevisiae, Mus musculus, Homo sapiens, Oryza sativa, Plasmodium falciparum, Drosophila melanogaster*
- 131 organisms have assemblies and chromosomal maps including:  
*Anopheles gambiae, Macaca mulatta, Bos taurus, Felis catus, Gallus gallus*
- Swiss-Prot Release 51.3/54.7 (Dec'06/Jan'08): 250K/333K entries; 91/120 million amino acids.

# Genome Sizes

Organism	Size	Date	Est. # genes
<i>HIV type 1</i>	9.2 Kb	1997	9
<i>H. influenzae</i>	1.8 Mb	1995	1,740
<i>M. genitalium</i>	0.58 Mb	1998	525
<i>E. coli</i>	4.7 Mb	1997	4,000
<i>S. cerevisiae</i>	12.1 Mb	1996	6,034
<i>C. elegans</i>	97 Mb	1998	19,099
<i>A. thaliana</i>	100 Mb	2000	25,000
<i>D. melanogaster</i>	180 Mb	2000	13,061
<i>M. musculus</i>	3 Gb	2002	~30,000
<i>H. sapiens</i>	3 Gb	2001	32,000+

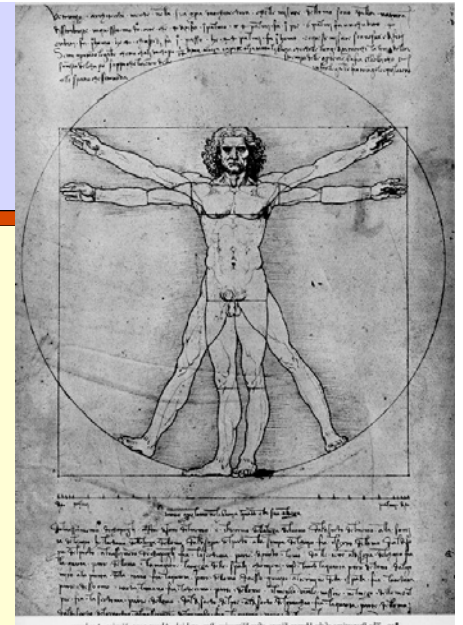
# *Caenorhabditis Elegans*

- Entire genome - 1998; 8 year effort
- 1<sup>st</sup> animal; 2<sup>nd</sup> eukaryote (after yeast)
- Nematode (phylum)
- Easy to experiment with; Easily observable
- 97 million bases; 20,000 genes;
- 12,000 with known function; 6 Chromosomes;
- GC content 36%
- 959 cells; 302-cell nervous system
- 36% of proteins common with human
- 15 Kb mitochondrial genome
- Results in **ACeDB**
- 25% of genes in operons
- Important for HGP: technology, software, scale/efficiency
- 182 genes with alternative splice variants



# *Homo sapiens*

- Sequenced - 2001; 15 year effort
- 3 billion bases, 500 gaps
- Variable density of **Genes, SNPs, CpG islands**
- ~ 1.1% of genome codes for proteins; **99%?**
- ~ 40-48% of the genome consists of repeat sequences
- ~ 10 % of the genome consists of repeats called ALUs
- ~ 5 % of the genome consists of long repeats (>1 Kb)
- 223 genes common with bacteria that are missing from worm, fly or yeast.



# Sequence Alignment – Why?

```
>gi|12643549|sp|O18381|PAX6_DROME Paired box protein Pax-6 (Eyeless protein)
MRNLPCLGTAGGSGLGGIAGKPSPTMEAVEASTASHRHSTSSYFATTYYHLTDECHSGVNQLGGVFVGG
RPLPDSTRQKIVELAHS GARPCDISRILQVSNGCVSKILGRYYETGSIRPRAIGGSKPRVATAEVVSKIS
QYKRECP SIF AWEIRD RLLQENVCTNDNIPSVSSINRVLRNLAAQKEQQSTGSGSSSTSAGNSISAKVSV
SIGGNVSNVASGSRGT LSSSTDLMQTATPLNSSES GGASNSGE GSEQEAIYEKLRLLN TQHAAGPGPLEP
ARAAPILVGQSPNHLGTRSSH PQLVHGNHQALQQHQQQSWPPRHYS GS WYPTSLSEIPISSAPNIASVTAY
ASGPSLAHSLSPN DIESLSIGHQRNC PVATEDIH LKKELDGHQSDETGS GEGEN SNGGAS NIGNTEDD
QARLILKRKLQRNRTSFTNDQIDSLEKEFERTHYPDV FARERLAG KIGLPEAR I QVWF SNRR AWRREEK
LRNQR RTPNSTGASATSS TSATAS LTDSPNSL SACSSLLSGSAGGP SV STINGLSSP TLSTNVNAPTL
GAGIDSSE SPTPIPHIRPSCTSDNDNGRQSEDCRRVCSPCPLGVGGHQ NT HIIQ SNGH A QGH ALVPAI SP
RLNFNSGSFGAM YSNMHTALSMSD SYGAVT PIP SFNHS AVGPL APPS PI PQ QGD LTP SLYP CHMT LRP
PPMAPAHHHIVPGD GGRPAGVGLGSGQ SANLGASC SGSGY EVL SAYAL PPPPMASS SAAD SSF SAASS AS
ANVTPHHTIAQ ESCP SP C SASHFGVAHSSGFSSDPISP AVSSYAHMS NYASSANT MTPSSA GTSAHV
APGKQQFFASC FYSPWV
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```
>gi|6174889|PAX6_HUMAN Paired box protein (Oculorhombin) (Aniridia, type II protein)
MQNSHGVNQLGGVFVN GRPLPDSTRQKIVELAHS GARPCDISRILQVSNGCVSKILGRYYETGSIRPRA
IGGSKPRVATPEVV SKIAQYKRECP SIF AWEIRD RLLSEG VCTNDNIPSVSSINRVLRNLASEKQQMGAD
GMYDKLRLMLNGQTGSW GTRPGW YPGT SVP GQPT QDGCQ QQEGGENT NSI S NGED SDEAQM RLQL KRKL
QRNRTSFTQE QIEALEKEFERTHYPDV FARERLA AKIDLPEAR I QVWF SNRR AWRREEK LRNQ RRQ ASN
TP SHIPI SSSF STSVYQ PIP QPT TPVSSFTSGSMI GRTDTALTNTY SALPP MPSFTM ANN LPMQ PPVPSQ
TSSYSCMLPTSPSVN GRSYDTYTPPHM QT H MNSQ PMGTSGTTSTGLI SPGV SVPVQV PGSEP DM SQY WPR
LQ
```

# Drosophila Eyeless vs. Human Aniridia

Query: 57 HSGVNQLGGVFVGGRPLPDSTRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETG 116  
HSGVNQLGGVFV GRPLPDSTRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETG

Sbjct: 5 HSGVNQLGGVFVNGRPLPDSTRQKIVELAHSGARPCDISRILQVSNGCVSKILGRYYETG 64

Query: 117 SIRPRAIGGSKPRVATAEVVSKISQYKRECP SIF AWEI RDRL QENVCTNDNIPSVSSIN 176  
SIRPRAIGGSKPRVAT EVVSKI+QYKRECP SIF AWEI RDRL E VCTNDNIPSVSSIN

Sbjct: 65 SIRPRAIGGSKPRVATPEVVSKIAQYKRECP SIF AWEI RDRL SEGVCTNDNIPSVSSIN 124

Query: 177 RVLRNLAQKEQ 188  
RVLRNLA++K+Q

Sbjct: 125 RVLRNLAQKEQ 136

Query: 417 TEDDQARLILKRKLQRNRTSFTNDQIDSLEKEFERTHY PDVFARERLAGKIGLPEARIQV 476  
+++ Q RL LKRKLQRNRTSFT +QI++LEKEFERTHY PDVFARERLA KI LPEARIQV

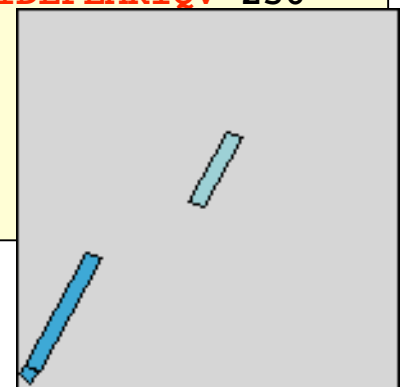
Sbjct: 197 SDEAQMRQLQKRKLQRNRTSFTQE QIEALEKEFERTHY PDVFARERLA KIDLPEARIQV 256

Query: 477 WFSNRRAKWRREEKLRNQRR 496

WFSNRRAKWRREEKLRNQRR

Sbjct: 257 WFSNRRAKWRREEKLRNQRR 276

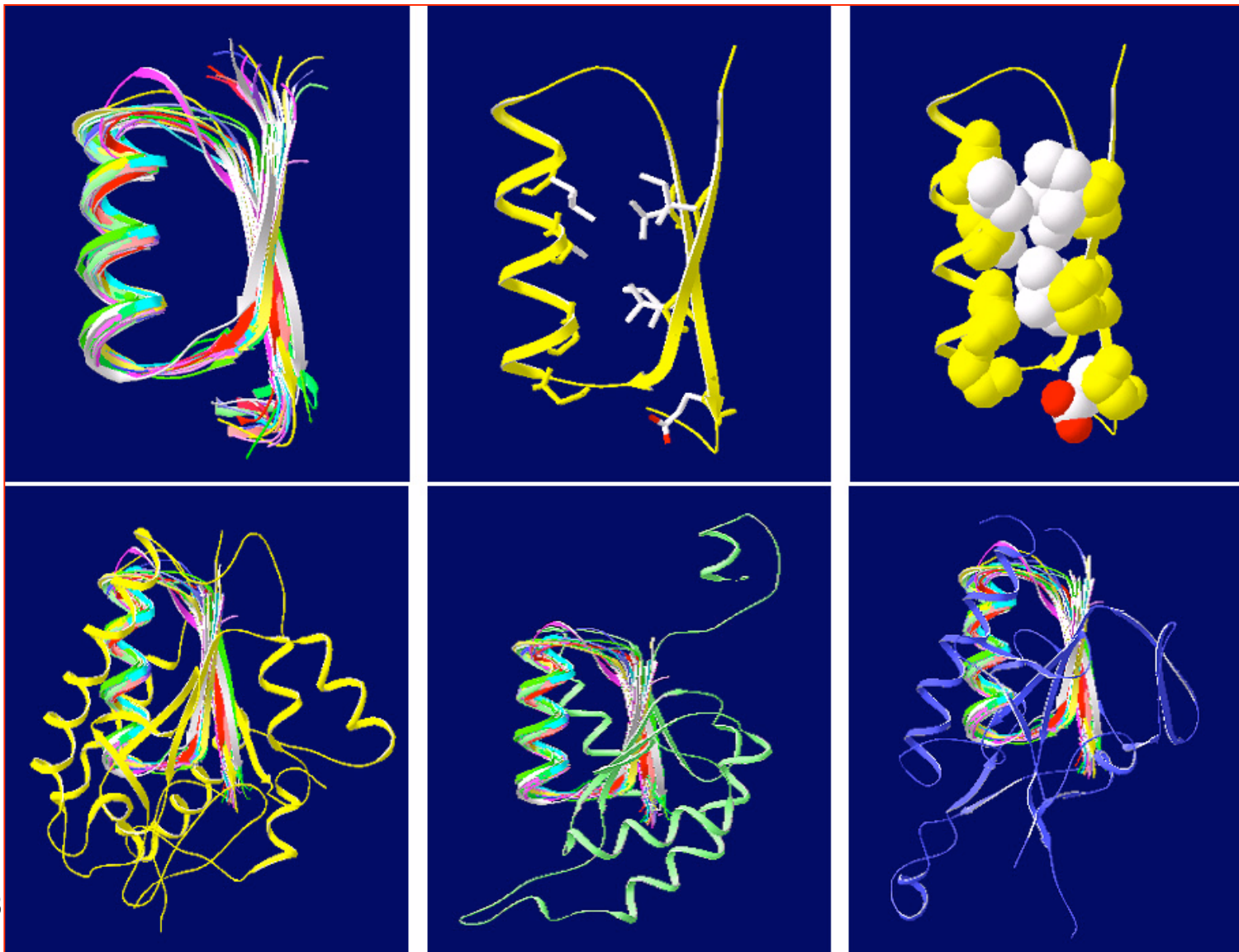
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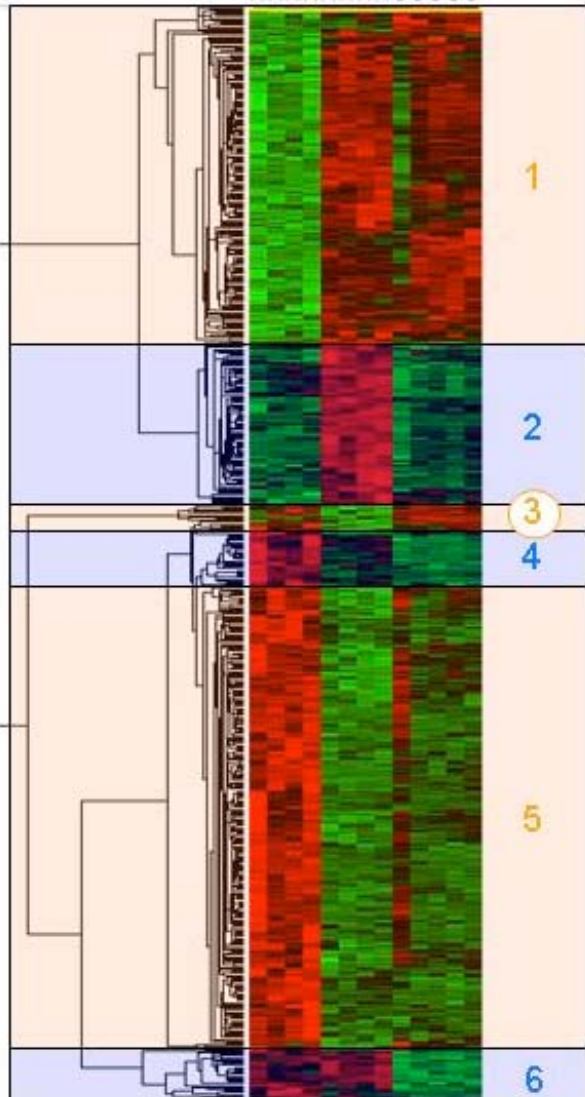
# Motif Detection in Protein Sequences

- ❑ MTDKMQLALAPVGNLDSYIRAAANAWPMLSADEERALAEKLHYHGDLEAA  
KTLILSHLRFVVHIARNYAGYGLPQADLIQEGRNIGLMKAVRRFNPEVGVR  
LVSFAVHWIKAEIHEYVLRNWRIVKVATTKAQRKLFFNLRKTQRLGWFN  
QDEVEMVARELGVTSKDVREMESRMAAQDMTFDLSSDDSDSQPMAPVLY  
LQDKSSNFADGIEDDNWEEQAANRLTDAMQGLDERSQDIIRARWLDEDNK  
STLQELADRYGVSAERVRQLEKNAMKKLRAAIEA
  
- ❑ MTDKMQLALAPVGNLDSYIRAAANAWPMLSADEERALAEKLHYHGDLEAA  
KTLILSHLRFVVHIARNYAGYGLPQADLIQEGRNIGLMKAVRRFNPEVGVR  
LVSFAVHWIKAEIHEYVLRNWRIVKVATTKAQRKLFFNLRKTQRLGWFN  
QDEVEMVARELGVTSKDVREMESRMAAQDMTFDLSSDDSDSQPMAPVLY  
LQDKSSNFADGIEDDNWEEQAANRLTDAMQGLDERSQDIIRARWLDEDNK  
STLQELADRYGVSAERVRQLEKNAMKKLRAAIEA

# Patterns in Protein Structures



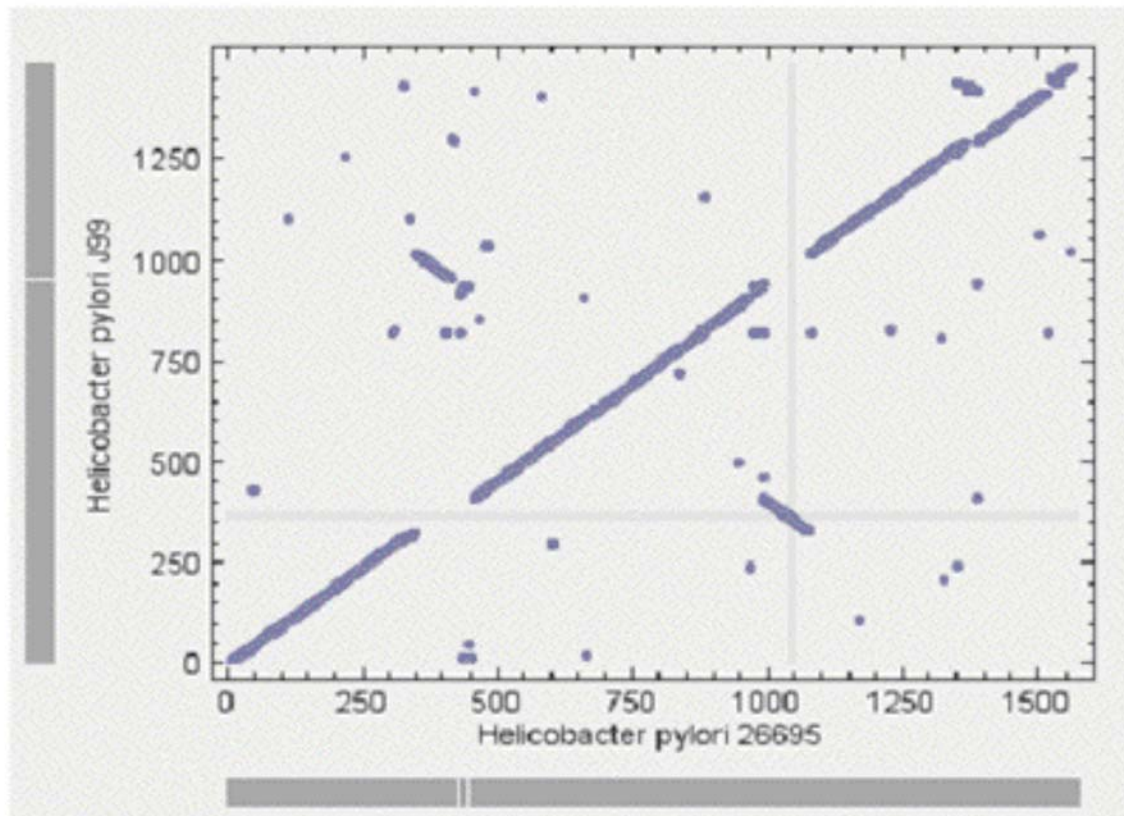
# Microarray Analysis



Different patterns of gene expression of oral epithelial IHGK cells upon co-culture with *A. actinomycetemcomitans* or *P. gingivalis*.

# Tools: GenePlot

1491 proteins total



Comparison of proteins from two strains of *Helicobacter Pylori*, 26695 and J99. Each point represents a pair of proteins from the two organisms showing a symmetrical best BLAST score; the coordinates of each point correspond to the position of the protein genes in the 2 genomes. Note the juxtaposition and inversion of two segments of the genome between the two strains.

# SIDS



- 18000 Amish people in Pennsylvania
- Mostly intermarried due to religious doctrine
- rare recessive diseases occurred with high frequencies.
- SIDS: 3000 deaths/year (US); 21 deaths (Amish community)
- Many research centers failed to identify cause
- Collaboration between Affymetrix, TGEN & Clinic for special children solved the problem in 2 months
- Studied 10000 SNPs using microarray technology
- Their experiments showed that all the sick infants had two mutant copies of a specific gene, and their parents were carriers of the mutant gene.
- Conclusion: **Disease caused by 2 abnormal copies of TSPYL gene**
- Identified genes expressed in key organs (brainstem, testes)
- [http://www.affymetrix.com/community/wayahead/modern\\_miracle.affx](http://www.affymetrix.com/community/wayahead/modern_miracle.affx)