Introduction to evolutionary concepts and VMD/MultiSeq - Part I

Characterizing molecular systems

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VMD/MultiSeq - "A Tool to Think"

Carl Woese - "VMD is far from a simple visualization tool for a biologist, it is a true thinking tool. Without it a whole class of biological hypotheses would simply not exist."



Why Look at More Than One Sequence?

1. Multiple Sequence Alignment shows patterns of conservation

Sequence Name	800										810										820							830									, i
SYN_THEAC SYN_357	s	Q	R	Т	w	Ν	Υ	D	Е	L	М	Q	R	Т	R	Е	А	Ν	L	D									Е	S		А	Υ	Υ	w	Y	V
SYNC_CAEEL 1473	s	М	R	Т	w	к	Е	D	Q	L	L	А	А	F	Е	к	G	G	L	D								S	к	Ν			Y	Y	w	Y	м
SYNC_MOUSE 1475	s	М	R	s	w	D	s	Е	Е	1	L	Е	G	Y	к	R	Е	G	Т	D								Р	А	Р			Y	Y	w	Y	т
SYNC_DEBHA 1480	s	М	R	т	Υ	D	Ν	D	Е	L	۷	А	А	Т	к	R	Е	G	L	D								L	D	s			Y	Y	w	F	т
SYNC_YEAST 1482	s	М	R	Т	D	D	м	D	Е	L	М	А	G	F	к	R	Е	G	Т	D										т	D	А	Y	Y	w	F	I.
SYNC_HUMAN 1476	s	М	R	Т	F	D	s	Е	Е	1	L	А	G	Y	к	R	Е	G	T.	D								Р	т	Р			Y	Y	w	Y	т
SYK2_METMA 133	Y	s	Е	L	Ν	D	Р	L	Е	Q	Е	к	R	F	Е	Е	Q	D	К	к	R	к	L (G		 D	L	Е	А	Q	т	V	D	Y	D	F	I.
SYK_HUMAN 1499	Y	т	Е	L	Ν	D	Р	м	R	Q	R	Q	L	F	Е	Е	Q	А	к	А	к	А	A	G		 D	D	Е	А	м	F	1	D	Е	Ν	F	С
SYK2_METAC 133	Y	s	Е	L	Ν	D	Р	L	Е	Q	Е	к	R	F	Е	Е	Q	D	к	к	R	к	L	G		 D	L	Е	А	Q	т	v	D	Y	D	F	1
SYK_MOUSE 1497	Y	т	Е	L	Ν	D	Р	V	R	Q	R	Q	L	F	Е	Е	Q	А	к	А	к	А	A	G		 D	D	Е	А	м	F	1	D	Е	Ν	F	С
SYK_CRIGR 1499	Υ	т	Е	L	Ν	D	Р	м	R	Q	R	Q	L	F	Е	Е	Q	А	к	А	к	A	A	G		 D	D	Е	А	м	F	1	D	Е	Ν	F	С
SYK_ORYSA 524	Y	т	Е	L	Ν	D	Ρ	۷	۷	Q	R	Q	R	F	Е	Е	Q	L	к	D	R	Q	S	G		 D	D	Е	А	М	А	L	D	Е	т	F	С

2. Are these positions functionally important? Active sites, folding,...

- 3. What and how many sequences should be included?
- 4. Where do I find the sequences and structures for MS alignment?

5. How to generate pairwise and multiple sequence alignments?



New Tools in VMD/MultiSeq

View sequence or structure phylogenies and

Sequence Editor: Manually adjust alignments or sequences

File Fdit Search

Protein / RNA Sequence Data

SwissProt DB (400K), Greengenes RNA (100K) Signatures, Zoom

Metadata Information, Clustal, MAFFT & Phylogenetic Trees

RAXml Trees, Genomic Content, Temperature DB

Blast & PsiBlast

Sequence Editor



Sequence /Structure Alignment

Protein & RNA secondary structure



QR non-redundant seq / str sets

Cluster analysis / Bioinformatics scripting Tutorials MultiSeq/ AARS EF-Tu/Ribosome

J. Eargle, D. Wright, Z. Luthey-Schulten, *Bioinformatics*, 22:504 (2006) E. Roberts, J. Eargle, D. Wright, Z. Luthey-Schulten, *BMC Bioinformatics*, 7:382 (2006)

Protein: RNA Complexes in Translation Evolutionary Analysis & Dynamics





r-Proteins/r-RNA "Signatures ribosomate Volution"

"Signatures Pibosomal evolution" **PNAS** 2008, **BMC** 2009, **BJ** 2010 "Motion L1 Stalk:tRNA" **JMB** 2010, "Ribosome Biogenesis" **JPC** 2012,3 "Whole cell simulations on GPUs" **IEEE** 2009,**Plos CB** 2011,**PRL**2011, *JCC* 2013, **PNAS** 2013, *SS* **PRL** 2013, **CSB** 2013 **C** 2012 **Nature** 2014, **BJ** 2015

Basic principles of evolutionary analysis for proteins & RNAs

- Comparative analysis of sequences and structures
- Multiple sequence alignments (gaps and editing)
- •Sequence and structure phylogenetic trees*
- •Reference to 16S rRNA tree
- •Horizontal or lateral gene transfer events
- Genomic context
- •Evolutionary profiles representing diversity
- Conservation analysis of evolutionary profiles

*Various models of evolutionary change

Alignment of ~200 EF-Tu sequences in VMD/MultiSeq



"G" scattered around gaps

http://www.clustal.org/clustal2/

* MAFFT v7.221, Katoh and Standley, Mol.Biol and Evol. 2015

~ 5 minutes

~ 30 seconds

More sequences!

Sequence Alignment & Dynamic Programming

number of possible alignments:

Seq. 1:
$$a_1 a_2 a_3 - - a_4 a_5 \dots a_n$$

Seq. 2: $c_1 - c_2 c_3 c_4 c_5 - \dots c_m$

S : substitution matrix

GHIL

-2 -1 -2

0 -3 -2

1 -2 -3

-4 -4 -2 -3 -3

2 -2 0 -3 -2 1 -1 -4 -2

7 -3 0 -4 -2 1 -2 -3

-2 -3 -2 -3 8 -2 -4 -4 -2 -2 -3 -1

0 -2 13 -3 -2 -1

1

1 1 -2 -1 -3 -2 6

-1 1

-1 2 -2

8

-3 -4 -2 -4 -3 -3 -2

-1 -3

-1 0

-2 -4 -2

-2 -2 -5 -2 0 -1 -2 -2 -4 -1 -2

-4 -5 -6 -1 -2 -2 -5 -3 -1 -2 -2

-2 -3 -4 -1 -2 -3 2 0 0 -1

1 0 -1 1 0 0 -1 -2 -3 0 -2 -2

-2 -2 1

-2

-1 -3

2 -2

9 -2 -2 16

2 -2 2

-3 -2

0 -4 0

-3 -3 -1

1 -1 -4

-1

0 0 - 3

-3 -2

1 -1

0 -2

-2 -1

0 -2

$$= \binom{2n}{n} = 2^{2n} \left(\sqrt{n\pi} \right)^{-1}$$

Needleman-Wunsch alignment algorithm

-1 -1 -3 -2

3 -1 -2 -3

0 -2 -3 -2

1 -2 -2 -1 -2 -5

-3 -3 -2 -3 -3 -4 -4 4 2 -2 1 0 -3 -1 1 -3 -1 5 -3 -3 -1 **V**

4 6 -2 0 1 -1 0 -3 -3 0 -3 -3 -2 0 0 -4 -3 -3 5 2 -1 **B**

-1 0 0 1 -3 4 5 -2 0 -4 -2 1 -2 -4 -1 0 -1 -2 -2 -3 2 5 -1 **Z** 0 -1 -1 -1 -2 -1 -1 -1 -1 -1 -1 -1 0 -1 -2 0 0 -2 -1 -1 -1 -1 -1 **X**

-1 -3 -1

0

9 - 4

-4 11

-1

0 -4 -3 0 -3 -4 -2

-2

2 - 3 0

3 -1

0 -1 -1 -1 -1 -2 -2 -1 -1 0 -1 -1 0 2 6 -4 -1

$$H(i, j) = MAX \begin{cases} H(i-1, j-1) + S[a(i), b(j)] \\ H(i, j-k) - W(k), \\ H(i-m, j) - W(m) \end{cases}$$

1 0 - 3

0

1 -4 -5 -4 19

1 4 -3 -2 -1 3

-2 -1

0

-2

-1 -1

-1

-2

-2

2 -5 -2

-2

-1

0

9 -1 -3 -2 -1 Y

0 - 1 - 2

0

0 -2 -2 -3

-1 0 **A**

0 -1 **R**

-1 K

-1 F

0 0 0 5

0 -1 0 **T**

 $\begin{array}{c|c} H(i-1, j-1) & H(i-1, j) \\ +S[a(i), b(j)] & -W(1) \\ \hline H(i, j-1) & \\ -W(1) & H(i, j) \end{array}$

Score Matrix H: Traceback

gap penalty W = -6

Reference: "Biological Sequence Analysis - Probabilistic Models of Proteins and Nucleic Acids" R. Durbin, S. Eddy, A. Krogh, and G. Mitchison, Cambridge U. P.London, 1998; pp. 19-22 (see also other sections)

Needleman-Wunsch Global Alignment



Similarity Values

Initialization of Gap Penalties



http://genome.dkfz-heidelberg.de/husar/fileadmin/handouts/02pairwise_method.pdf

Filling out the Score Matrix H





Traceback and Alignment



The Alignment

м	G	-	к	-	Р
:			:		:
м	G	Р	к	к	Р

Traceback (blue) from optimal score

STAMP - Multiple Structural Alignments

- 1. Initial Alignment Inputs
- Multiple Sequence alignment
- Ridged Body "Scan"
- Pairwise Alignments and Hierarchical Clustering
- 2. Refine Initial Alignment & Produce Multiple Structural Alignment

$$P_{ij} = \left\{ e^{-d_{ij}^2/2E_1} \right\} \left\{ e^{-s_{ij}^2/2E_2} \right\}$$

probability that residue ion structure A is equivalent to residue jon structure B.

 S_{ij} — conform ational similarity; function of rms bew teen i-1, i, i+1 and j-1, j, j+1.

•Dynamic Programming (Smith-Waterman) through P matrix gives optimal set of equivalent residues.

•This set is used to re-superpose the two chains. Then iterate until alignment score is unchanged.

•This procedure is performed for all pairs with no gap penalty

R. Russell, G. Barton (1992) *Proteins* 14: 309 R.B. Russel, T. Walsh, G. Barton, STAMP version 4.4: User Guide, 2010.

Multiple Structural Alignments

STAMP – cont' d

2. Refine Initial Alignment & Produce Multiple Structural Alignment

Alignment score: $S_p L_p - i_A L_p - i_B$

$$S_{c} = \frac{S_{p}}{L_{p}} \frac{L_{p} - L_{A}}{L_{A}} \frac{L_{p} - L_{B}}{L_{B}}$$
$$S_{p} = \sum_{\text{aln.path}} P_{ij}$$

 \mathbf{L}_{p} , \mathbf{L}_{A} , \mathbf{L}_{B} — length of alignment, sequence A, sequence B \mathbf{j}_{A} , \mathbf{j}_{B} — length of gaps in A and B.

Multiple Alignment:

- •Create a dendrogram using the alignment score.
- •Successively align groups of proteins (from branch tips to root).
- •When 2 or more sequences are in a group, then average coordinates are used.

Initial Pairwise Superposition - Single Linkage Cluster

-	_	-		
			ι.	
2			Ŀ	
			L	
-		-		

ABCD



C - - - 3.4

D - - - - -

N = 4 proteins, N(N-1)/2 pairs Table of RMS



Dendogram: Initial step starts by fitting B to D (0.5 A), then A to C (1.0A), single-linkage is a nearest neighbor method in which the distance between a pair of clusters is equal to the shortest distance between any two members - hence furthest pt is 3.0 (AB) and not 4.0 (AD)

Structural Overlaps - STAMP

Ribosome large subunit showing ribosomal proteins L2 and L3 180,000 atoms in 4 rRNAs and 58 proteins



Ribosomal Pr	rotein L2																																											
2aw4_C	V r i	41				G	R	Ν	N	Ν	G	R	Т	Т	Т	R	Н	1	G	G	G	н	к	Q	А	Υ	R	1	V		D	F	к	R	Ν	К		D			G	1	Ρ	А
□ 1s72_A	Vri	11	R	G	т	s	т	F											R	А			Р	s	н	R	Y	к	А	D	L	Е	н	R	к	۷	Е	D	G	D	v	1	А	G
Ribosomal Pr	rotein L3																																											
2aw4_D	Vri	11	м	т	R	1	F	т												Е	D	G	V	s	1	Ρ	V	т	V	Т	Е	V	Е	А	Ν	R	۷	Т	Q	V	к			
□ 1s72_B	vr i	49	т	н	V	v	L	V	N	D	Е	Р	Ν	S	Р	R	Е	G	м	Е	Е	т		v	Р	v	т	V	1	Е	т	Р	Р	М	R	А	V	А	L	R	А	Y	E	D

Universal Phylogenetic Tree 3 domains of life



Reference 16S rRNA tree

Leucyl-tRNA synthetase displays the full canonical phylogenetic distribution.

Look for horizontal gene transfer events



After W. Doolittle, modified by G. Olsen

Phylogenetic Distributions

Full Canonical

Basal Canonical

Non-canonical



increasing inter-domain of life Horizontal Gene Transfer

"HGT erodes the historical trace, but does not completely erase it...." G. Olsen

Woese, Olsen, Ibba, Soll MMBR 2000

Protein Structure Similarity Measure

Q_H Structural Homology

fraction of native contacts for aligned residues + presence and perturbation of gaps

 $Q_H = \aleph \left[q_{aln} + q_{gap} \right]$

$$q_{aln} = \sum_{i < j-2} \exp\left[-\frac{(r_{ij} - r_{i'j'})^2}{2\sigma_{ij}^2}\right]$$



O'Donoghue & Luthey-Schulten MMBR 2003.

Structural Similarity Measure: The effect of insertions

"Gaps should count as a character but not dominate" C. Woese





Structure encodes evolutionary information!

Structure reveals distant evolutionary events Class I AARSS

structure-based phylogenetics

sequence-structure overlap









Sequences define more recent evolutionary



Conformational changes in the same protein.

ThrRS T-AMP analog, 1.55 A. T, 2.00 A.

 $Q_{\rm H} = 0.80$ Sequence identity = 1.00



Structures for two different species.

ProRS

M. jannaschii, 2.55 A. *M. thermoautotrophicus*, 3.20 A.

 $Q_{\rm H} = 0.89$ Sequence identity = 0.69

Relationship Between Sequence & Structure



Structural alignment & visualization software MultiSeq/VMD

Non-redundant Representative Profiles



QR computes a set of maximal linearly independent structures.

P. O'Donoghue and Z. Luthey-Schulten (2003) MMBR 67:550-571.

P. O'Donoghue and Z. Luthey-Schulten (2005) J. Mol. Biol., 346, 875-894.

Numerical Encoding of Proteins in a Multiple Alignment

Encoding Structure

Rotated Cartesian + Gap = 4-space

 $(x_{C_{\alpha}}, y_{C_{\alpha}}, z_{C_{\alpha}}, 0)$ Aligned position

Gapped position

(0,0,0,g)

Gap Scaling

 $\frac{\|X\|_{F_4} + \|Y\|_{F_4} + \|Z\|_{F_4}}{\|G\|_{F_4}}$ adjustable parameter

Sequence Space Orthogonal Encoding = 24-space

23 amino acids (20 + B, X, Z) + gap

Alignment is a Matrix with Linearly Dependent Columns



A maximal linearly independent subset can be determined with respect to a threshold, e.g., similarity measure threshold.



Evolution of Structure and Function in AspRS

Summary Structural Evolutionary Profiles

1.Structures often more conserved than sequences!! Similar structures at the Family and Superfamily levels. Add more structural information to identify core and variable regions 2. Which structures and sequences to include? Use evolution and eliminate redundancy with QR factorization

New Tools in VMD/MultiSeq

Protein / RNA Sequence Data

SwissProt DB (400K), Greengenes RNA (100K) Signatures, Zoom

Metadata Information, Clustal & Phylogenetic Trees

RAXml Trees, Genomic Content, Temperature DB

Blast & PsiBlast

Sequence Editor



View sequence or structure phylogenies and Group data by taxonomic classification eliminate redundancy with QR Eukaryota:Fung V r i □ 1asy_A □ 1eov A VII SYDC YEAS 2 1 r 1 2 1 V r i V r i SYD METMA r i r i SYD PYRHO r i Vr i Vri TEYCG Align sequences with Clustal



Sequence /Structure Alignment

Protein & RNA secondary structure

QR non-redundant seq / str sets

Cluster analysis / Bioinformatics scripting Tutorials MultiSeq/ AARS EF-Tu/Ribosome

J. Eargle, D. Wright, Z. Luthey-Schulten, *Bioinformatics*, 22:504 (2006) E. Roberts, J. Eargle, D. Wright, Z. Luthey-Schulten, *BMC Bioinformatics*, 7:382 (2006)

MultiSeq Combines Sequence and Structure

- Align sequences or structures; manually edit alignments
- View data colored by numerous metrics including structural conservation and sequence similarity
- Synchronized coloring between 1D and 3D views



Load large sequence sets*

Swiss-Prot (Proteins) Curated sequences 392,667 sequences Unaligned 177 MB on disk 2 minutes to load 2.4 GB memory used

Greengenes (RNA)*

- **Environmental 16S rRNA**
- 90,654 entries
- Aligned (7682 positions)
- 670 MB on disk
- 2.5 minutes to load *
- 4.0 GB memory used*

*"Signatures of ribosomal evolution" with Carl Woese, PNAS (2008) *Release May 2013 contains 1.2 million sequences – Memory??

Sequence editor

- New sequence API allows editing of large alignments. Align closely related sequences by group, combine groups, and then manually correct.
- Zoom window gives an overview of the alignment, quickly move the editing window to any part of the alignment.

660 sequences

protein S4 from

of ribosomal

all complete

bacterial

genomes^{*}.



* K. Chen, E. Roberts, Z Luthey-Schulten (2009) BMC Bioinformatics

Phylogenetic tree editor

 Automatically add annotations and colors to phylogenetic trees based on taxonomy, enzyme, temperature class, and/or MultiSeq groupings.



Maximum likelihood tree of 660 S4 sequences reconstructed using RAxML.



A cluster of five proteobacterial sequences branch near the cyanobacterial sequences. These are cases of horizontal gene transfer.

Elijah Roberts 2009

Scripting MultiSeq

- All MultiSeq functions can be scripted.
- Scripting an analysis provides benefits:
 - It can be checked for correctness.
 - It can be quickly repeated by anyone.
 - It can be modified later with new functionality.
 - It can be run on a cluster in VMD text mode. (if it can be easily broken into independent chunks)
- Many functions are too user specific and/or too complex to be turned into a GUI.
- Some examples of MultiSeq scripts...

Genome content

- When using sequence from fully sequenced genomes, additional information is available in the genome content.
- Conservation of gene ordering, neighbors, or intergenic regions can provide additional evolutionary information not contained in the sequence.
- Gene names and ordering can be obtained from the genome PTT files, want to organize the information in an evolutionarily meaningful manner.

Location	Strand	Length	n PID	Gene	Synonym	Code COG	Product
34376383438021	-	127	16131173	rplQ	b3294 -	COG0203J	50S ribosomal subunit protein L17
34380623439051	-	329	16131174	rpoA	b3295 -	COG0202K	RNA polymerase, alpha subunit 🛛 🗸
34390773439697	-	206	16131175	rpsD	b3296 -	COG0522J	30S ribosomal subunit protein S4
34397313440120	-	129	16131176	rpsK	b3297 -	COG0100J	30S ribosomal subunit protein S11
34401373440493	-	118	16131177	rpsM	b3298 -	COG0099J	30S ribosomal subunit protein S13
34406403440756	-	38	16131178	rpmJ	b3299 -	COG0257J	50S ribosomal subunit protein L36
34407883442119	-	443	16131179	secY	b3300 -	COG0201U	preprotein translocase membrane subunit
34421273442561	-	144	16131180	rplO	b3301 -	COG0200J	50S ribosomal subunit protein L15
34425653442744	-	59	16131181	rpmD	b3302 -	COG1841J	50S ribosomal subunit protein L30
34427483443251	-	167	16131182	rpsE	b3303 -	COG0098J	30S ribosomal subunit protein S5

Combined genomic context/phylogenetic tree

 Use a script to walk through a phylogenetic tree, find the genome content near the source gene, create a graphical representation of the combined data.

proc draw_genome_context_of_phylogeny {args} {

```
# Load the sequences.
set alignment [::SeqData::Fasta::loadSequences $alignmentFilename]
```

```
# Load the tree
set tree [::PhyloTree::Newick::loadTreeFile $treeFilename]
# Reorder the alignment by the tree.
set treeAlignment {}
set leafNodes [::PhyloTree::Data::getLeafNodes $tree]
foreach node $leafNodes {
    set foundNode 0
    set nodeName [::PhyloTree::Data::getNodeName $tree $node]
    foreach sequence $alignment {
        if {$nodeName == [::SeqData::getName $sequence]} {
            lappend treeAlignment $sequence
            set foundNode 1
            break
        }
    }
}
```

Draw the genomic context.

}

drawGenomicContextOfAlignment \$outputFilename \$treeAlignment \$contextDistance \$scaling \$genomeDirectory

Combined genomic context/phylogenetic tree

proc drawGenomicContextOfAlignment {outputFilename alignment contextDistance scaling genomeDirectory} {

```
fcreach sequence $alignment {
    # Make sure we have the GI number for this sequence.
    set giNumber [::SeqData::getSourceData $sequence "gi"]
    # Make sure we can tell which genome this sequence is from.
    set taxonomy [join [::SeqData::getLineage $sequence 1 0 1] ","]
    if {![info exists genomeTaxonomyMap($taxonomy)]} {
        error "ERROR} Unknown genome for sequence [::SeqData::getName $sequence]: $taxonomy"
    }
    # Go through each of the genome context files for the genome.
    set foundGene 0
    foreach genomeName $genomeTaxonomyMap($taxonomy) {
        ...
    }
}
```

```
# Draw the genomic context.
```

}

drawMultipleGenomicContext \$outputFilename \$alignment \$geneFiles \$genePositions \$geneStrands \$contextDistance

	ſ	Betaproteobacteria, Thiobacillus denitrificans ATCC 25259	+rpsE -rpm +-	+secY	+infA +rpsM +-	+rpsD	+-	+-	+-		
	100	100 Betaproteobacteria, Azoarcus sp. BH72	-rpsE rpmD -rplO	-secY	infA rpmJ -rpsM -rpsK	-rpsD	-rpoA	-rplQ	-galE2	+-	+uvrA2
	67	Betaproteobacteria, Azoarcus sp. EbN1	+rpsE +rpm +rplO	+secY	+rpm +rpsM +rpsK	+rpsD	+rpoA	+rplQ	+galE		+-
	П	⁰⁰ Betaproteobacteria, Dechloromonas aromatica RCB	+rpsE +rpm +rplO	+secY	+infA +- +rpsM +-	+rpsD	+-	+-			
10		Betaproteobacteria, Nitrosospira multiformis ATCC 25196	+rpsE +- +-	+secY	+infA +rpsM +-	+rpsD	+-	+-	+- +-		
	1	Betaproteobacteria, Nitrosomonas eutropha C91	+rpsE +- +-	+secY	+infA +rpsM +-	+rpsD	+-	+-		+smpB	+-
		Betaproteobacteria, Nitrosomonas europaea ATCC 19718	+rpsE -rpm +-	+secY	+infA +rpsM +rpsK	+rpsD	+rpoA	+-		+smpB	+-
4		100 Gammaproteobatteria, Psychrobacter arcticus 273-	+rpsE -rpm +rplO	+secY	+rpmJ +rpsM +rpsK	+rpsD	+rpoA	+rplQ			+-
		100 Gammaproteobacteria, Psychrobacter cryohalolentis K	+rpsE -rpm +rplO	+secY	+rpmJ +rpsM +-	+rpsD	+-	+rplQ			+-
	10	Gammaproteobacteria, Psychrobacter sp. PRwf-1	+rpsE +rpm +rplO	+secY	+rpsM +-	+rpsD		+rplQ			+-
	10	Gammaproteobacteria, Acinetobacter sp. ADP1	-rpIR -rpsE rpmD -rpIO	-secY	rpmJ -rpsM -rpsK	-rpsD	-rpoA	-rplQ			+fadE
		Gammaproteobacteria, Acinetobacter baumannii SDF	+rplF +rpsE +rpm +rplO	+secY	+rpmJ +rpsM +rpsK	+rpsD	+rpoA	+rplQ	+-		-fadE
		Gammaproteobacteria, Acinetobacter baumannii AYE	+rplF +rpsE +rpm +rplO	+secY	+rpmJ +rpsM +rpsK	+rpsD	+rpoA	+rplQ	+-		-fadE
		Gammaproteobacteria, Acinetobacter baumannii ACICU	+rplF +rpsE +rpm +rplO	+secY	+rpmJ +rpsM +rpsK	+rpsD	+rpoA	+rplQ	+-		-fadE

Genome content future directions

- Genome content still a work in progress.
- Good candidate for a GUI: combined phylogenetic tree/ genome content viewer.
- Can also use COG codes to color by gene function.
- Still need API for manipulating PTT files.

Roberts, Chen, ZLS, **BMC Evol. Bio**. 2009

See also ITEP for microbial genomes, Benedict et al. **BMC Genomics 2014**



Genome content of ribosomal protein S4 by occurrence of the gene in the alpha operon.

Fifteen Clostridia genomes contain two copies of S4: one zinc-binding and one zinc-free.

Tree of Methanogens

Conservation Differentially Expressed Genes



Molecular Signatures of Translation- Drug Targets



E. Roberts, A. Sethi, J. Montoya, C. R. Woese & Z. Luthey-Schulten. *PNAS* "Molecular Signatures of Ribosomal Evolution" (2008)

Kim,... Luthey-Schulten, Ha, and Woodson, *Nature* "Protein-guided RNA dynamics during early ribosome assembly (2014)

Flexible Grouping of Data

- Automatically group data by taxonomic classification to assist in evolutionary analysis (HGT) or create custom groups
- Apply metrics to groups independently, e.g bacterial signal

Sequence Name									90							
Eukaryota:Fungi																
🗖 1asy_A 🛛 🔽 🖬 83	s	R	D	S	D	R	т	G	Q	К	R	V	Κ	F	V	D
☐ 1eov_A ▼ 1 83	s	R	D	S	D	R	т	G	Q	к	R	V	K	F	V	D
SYDC_YEAST SYDC_82	s	R	D	S	D	R	Т	G	Q	К	R	V	Κ	F	V	D
Eukaryota:Metazoa																
SYD_CAEEL S7	s	к			Е	Κ	к	V	L	Ν	F	L	K	V	к	Е
SYD_HUMAN 🖬 🖬 33	s	Q			Е	Κ	Р	D	R	V	L	V	R	V	R	D
SYD_MOUSE I 33	s	Q			Е	Κ	Р	D	R	V	L	V	R	V	к	D
Archaea:Crenarcha																
SYD_AERPE SYD_1								м	L	к	D	R	F	1	А	D
Archaea:Euryarchaeota																
🗖 1n9w_A 🛛 🔽 🛐 1										м	R	V	L	V	R	D
🗖 1b8a_A 🛛 🔽 🖬 1								М	Y	R	Т	н	Y	S	S	Е
SYD_METMA II 1				М	S	L	А	N	L	R	т	н	Y	т	А	D
SYD_HALN1 II 1								М	Е	Ν	R	т	Y	т	А	D
SYD_THEAC SI 1											М	L	S	1	А	Е
SYD_PYRHO I								М	Т	Е	К	۷	Y	С	Q	Е
Bacteria:Proteobacteria																
🗖 110w_A 🛛 🖬 1							М	R		R	Т	Н	Y	А	G	S
🗖 1il2_A 🛛 🔽 1								М		R	Т	Е	Υ	С	G	Q

MultiSeq: Display and Edit Metadata

- External databases are crossreferenced to display metadata such as taxonomy (lineage), data source (sp, Uniprot #), EC, enzymatic function
- Changes to metadata should periodically be updated!!!
- Electronic Notebook: Notes and annotations about a specific sequence or structure can be added – and saved

Sequence Name:	SYDC_YEAST	
Source Organism:	Saccharomyces cerevisiae	
Common Name:	yeast	
EC Number:	6.1.1.12	
EC Description:	AspartatetRNA ligase.	
Description:	Aspartyl-tRNA synthetase, cytoplasmic (EC 6.1.1.12) (AspartatetRNA ligase) (AspRS) - Saccharomyces cerevisiae (Baker's yeast).	•
Data Sources:	sp=P04802,SYDC_YEAST pdb=1EOV,A	•
Lineage:	1	
Lineage.	Eukaryota	-
	Ascomycota	
	Saccharomycotina	
	Saccharomycetes	-
Notes	,	_
Ther miss	e were ing residues	
	OK Cancel	•