

-old homepage-

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http://mesguerra.net
All class materials at:
https://github.com/esguerra/ki2430

Why Study RNA?

- 1. To understand what RNA does.
- 2. To **predict** what RNA can do.
- 3. To **control** RNA.

Borrowed from:

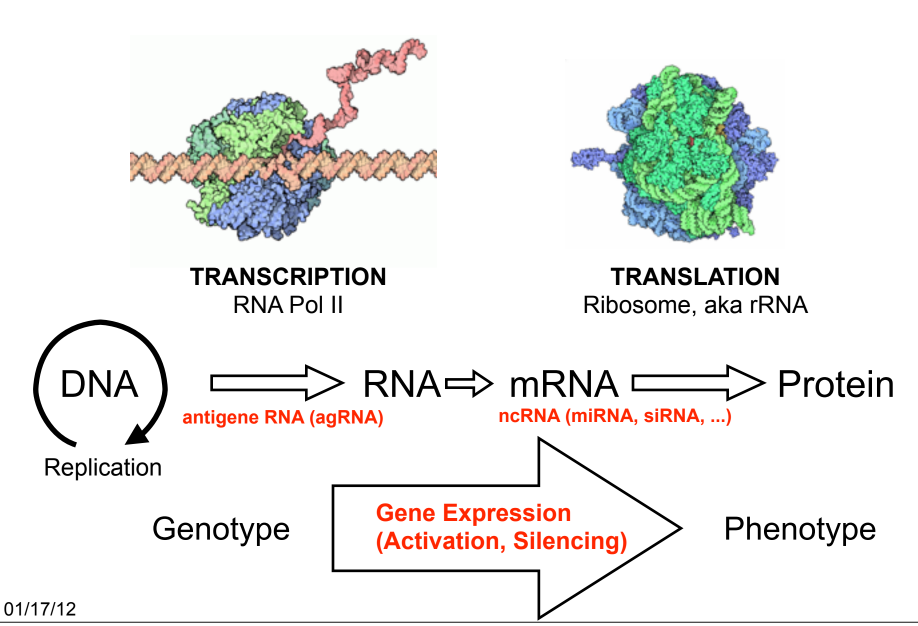
Ignacio Tinoco @ MSRI June 2000

http://www.msri.org/realvideo/ln/msri/2000/molbio/

tinoco/1/banner/01.html

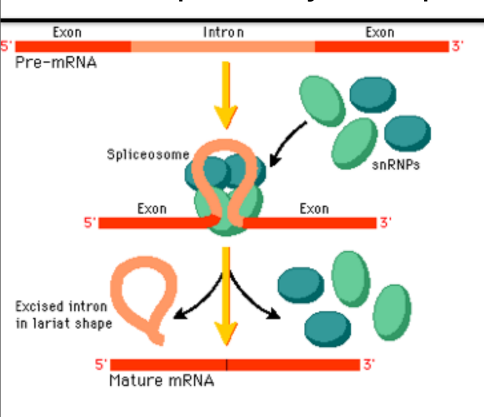
Highly Recommended!

Biology's Central Dogma and Molecular Machines



Monday, January 30, 2012

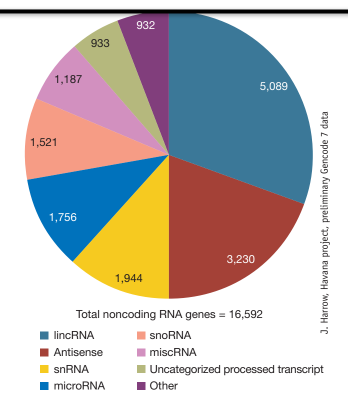
RNA is Spliced by the Spliceosome After Transcription.



Protein GENES RNA GENES

http://genome.ucsc.edu/ENCODE/

http://www.gencodegenes.org/



Long noncoding RNAs are just one of many noncoding transcripts being annotated. lincRNA, long intergenic noncoding RNA; snRNA, small nuclear RNA; snoRNA, small nucleolar RNA; and miscRNA, miscellaneous RNA.

- ~3200 Million base-pairs (Humans)
- < 2% makes mature mRNA (protein)
- ~50% transcribed ncRNA.

Monya Baker, Nature Methods, 8, 379-383, 2011

Non-coding RNA's

- RNApol I rRNA not 5S (~ 50% cell RNA)
- RNApol II precursors of <u>mRNA</u> and most <u>snRNA</u> and microRNA
- RNApol III 5S rRNA, tRNA and other small RNAs
- piwiRNA, siRNA, etc.
- http://rfam.sanger.ac.uk/family/browse/ with structure#A

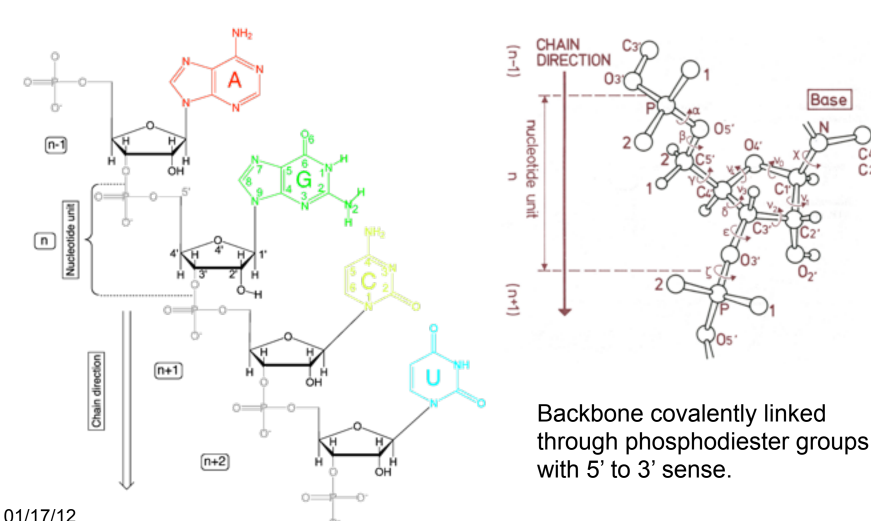
Structure Determines Function

RNA Structure Can Be Understood at Levels.

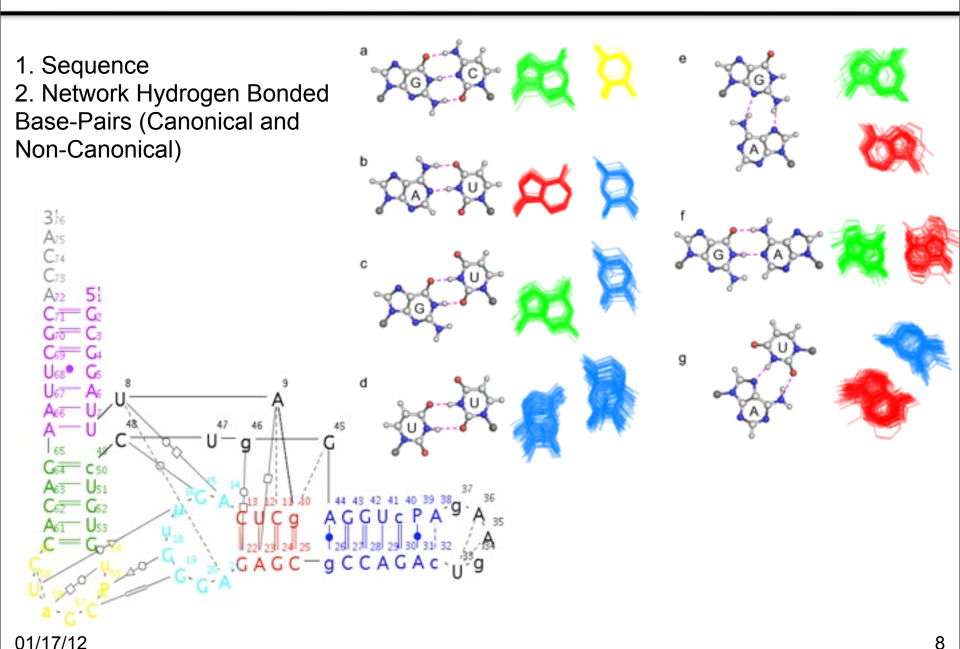
Primary Structure (Sequence)

GCGGAUU UA gCUC AGUUGGGA GAGC gCCAGAC UgAAg APCUGGA GGUC CUGUG uPCGaUC CACAG AAUUCGC ACCA 1234567 89 0123 45678901 2345 6789012 34567 8901234 5678 90123 4567890 12345 6789012 3456

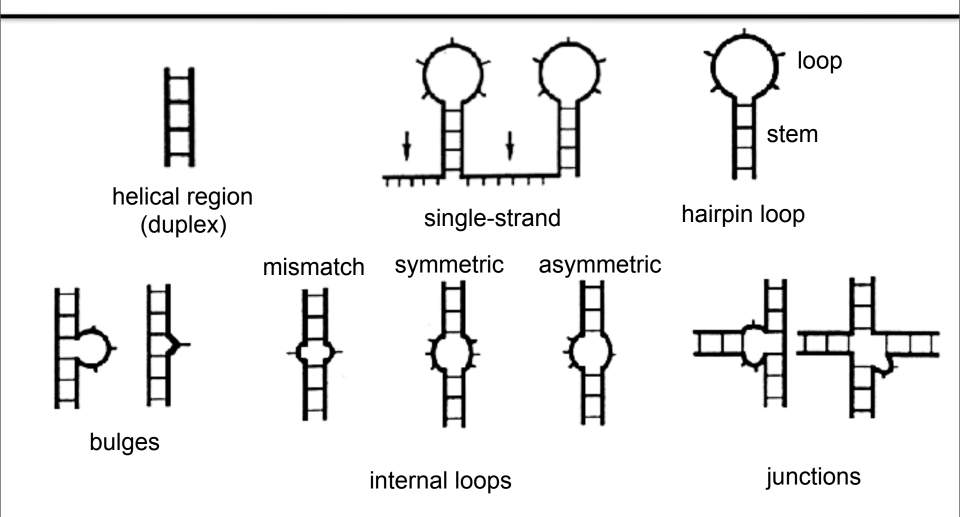
C4 (purines) C2 (pyrimidines)



Secondary Structure (Base-pairing)

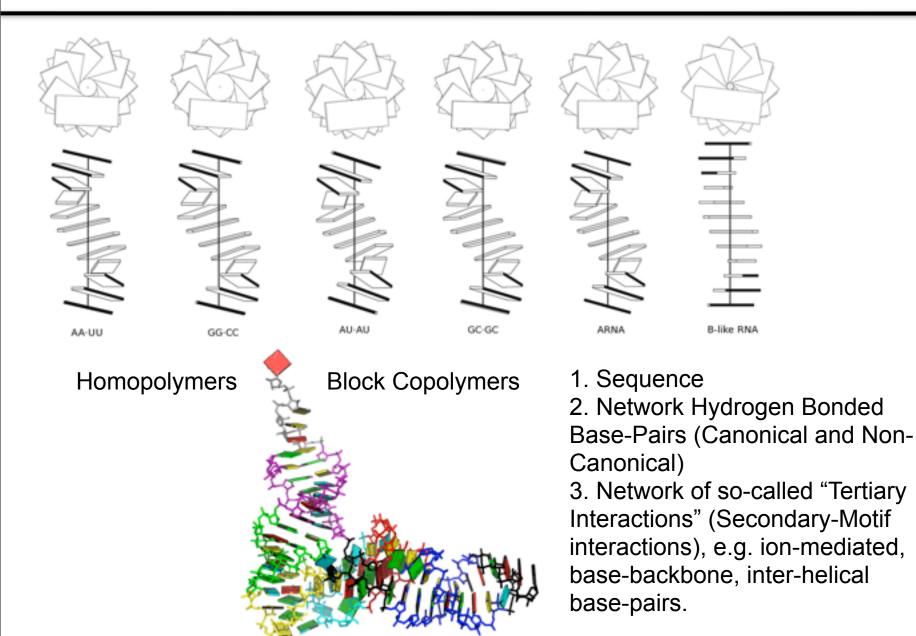


RNA Secondary Structure "Elements"



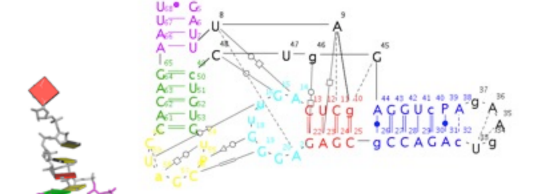
SCOR: Structural Classification of RNA http://scor.berkeley.edu

3D Structure (Subtle Conformational Variation of Helical Regions)





Quick Tour of RNA Secondary Structure Prediction



Mauricio Esguerra Neira Lennart Nilsson Group

http://mesguerra.net

Principles of Nucleic Acid Structure
KI Doctoral Course 2430
Torsdag, Februari Två, 2012
13:00 - 16:00

Two Kinda ThreeTypes of RNA Secondary Structure Prediction

RNA Sequence Covariation (Gutell)

RNA Free Energy Minimization (Tinoco-Uhlenbeck)

RNA Base-Pair Maximization (Nussinov)

Sequence Covariation

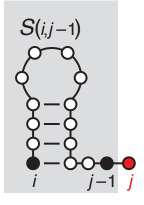
- Main Idea is:
- Gutell quite successful in getting rRNA.

Delta G Minimization

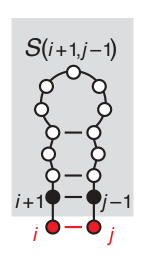
Tinoco-Uhlenbeck Postulate
 "Base-pair free energies are additive"

Dynamic Programming Nussinov Algorithm

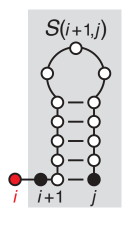
Nussinov Algorithm (Maximum Number of Base-Pairs)



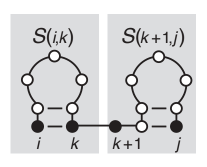
3. *j* unpaired



1. *i,j* pair



2. i unpaired



4. Bifurcation

Initialization

$$D(i,j) = \max \begin{cases} D(i,i) = 0 & \forall i = 1..L \\ D(i,i-1) = 0 & \forall i = 2..L \end{cases}$$

Recursion

$$D(i,j) = \max \begin{cases} D(i,k) + D(k+1,j) & where \ i \le k < j \\ D(i+1,j-1) + w(i,j) \end{cases}$$

Nussinov and Jacobson PNAS 77, 6309-6313 (1980)

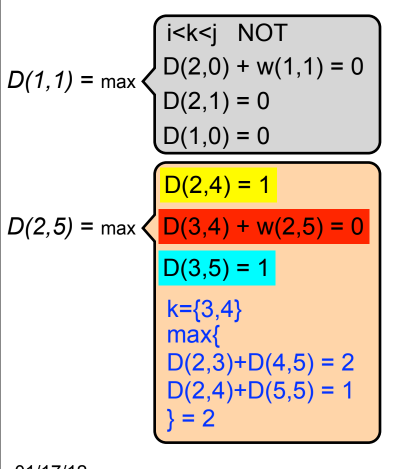
Nussinov et al. SIAM J. Appl. Math. 35, 68-82 (1978)

 $D(i,j) = \max \begin{cases} D(i,j-1) \\ D(i+1,j-1) + w(i,j) \\ D(i+1,j) \\ \max_{i < k < j} D(i,k) + D(k+1,j) \end{cases}$

The Algorithm in Action (4 MNBP)

$$D(i,j) = \max \left\{ \begin{array}{ll} D(i,j-1) \\ D(i+1,j-1) + w(i,j) \\ D(i+1,j) \\ \max_{i < k < j} D(i,k) + D(k+1,j) \end{array} \right. \begin{array}{ll} S(i,j-1) \\ O(i+1,j) \\ \max_{i < k < j} D(i,k) + D(k+1,j) \end{array} \right. \begin{array}{ll} S(i,j-1) \\ O(i+1,j) \\ \max_{i < k < j} D(i,k) + D(k+1,j) \end{array} \\ \begin{array}{ll} S(i,j-1) \\ O(i+1,j-1) \\ O(i$$

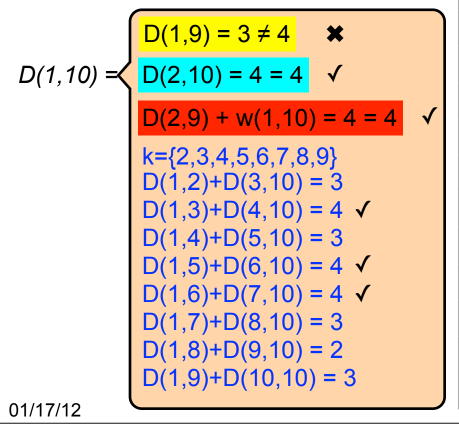
S. Eddy, Nature Biotech. 22, 1457-1458 (2004)



	j	1	2	3	4	5	6	7	8	9	10
i		Α	C	G	G	С	Α	Α	С	G	U
1	Α	0	0	1	1	2	2	2	2	3	4
2	O	0	0	1	F	2	2	2	2	3	4
3	G		0	0	0	7	1	1	2	2	3
4	G			0	0	1	1	1	1	2	3
5	O				0	0	0	0	0	1	2
6	A					0	0	0	0	1	2
7	Α						0	0	0	1	2
8	O							0	0	1	1
9	G								0	0	0
10	J									0	0

Backtracking to Structure (Counterclockwise)

$$D(i,j) = \begin{cases} D(i,j-1) \\ D(i+1,j) \\ D(i+1,j-1) + w(i,j) \\ for_{i < k < j} do D(i,k) + D(k+1,j) \end{cases}$$



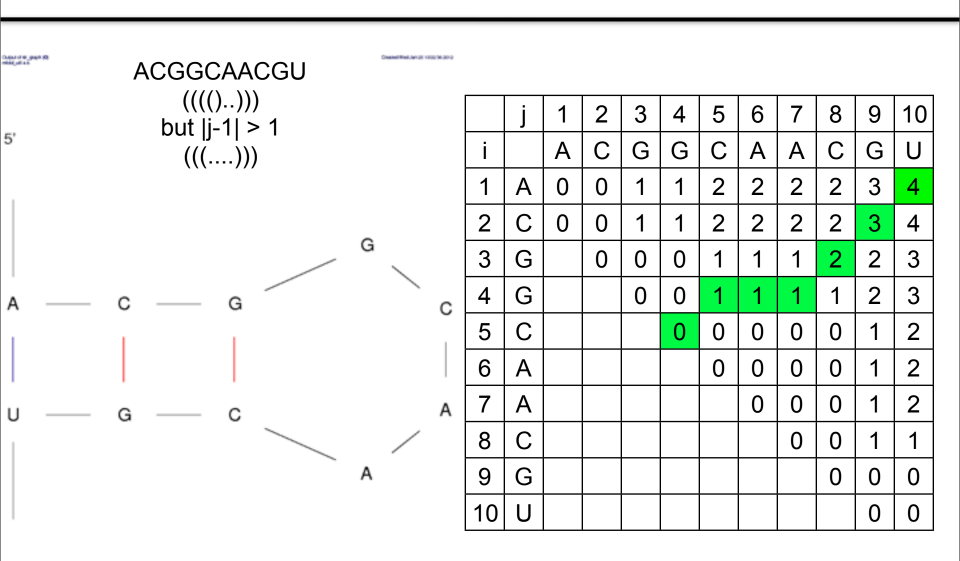
	j	1	2	3	4	5	6	7	8	9	10
i		Α	С	G	G	C	Α	Α	С	G	U
1	Α	0	0	1	1	2	2	2	2	3	4
2	O	0	0	1	1	2	2	2	2	3	4
3	G		0	0	0	1	1	1	2	2	3
4	G			0	0	1	1	1	1	2	3
5	C				0	0	0	0	0	1	2
6	Α					0	0	0	0	1	2
7	Α						0	0	0	1	2
8	C							0	0	1	1
9	G								0	0	0
10	U									0	0

One Possible Structure With 4 Base-Pairs

ACGGCAACGU (((()..))) but |j-1| > 1 (((....)))

	j	1	2	3	4	5	6	7	8	9	10
i		Α	C	G	G	O	Α	Α	O	G	U
1	Α	0	0	1	1	2	2	2	2	3	4
2	O	0	0	1	1	2	2	2	2	3	4
3	G		0	0	0	1	1	1	2	2	3
4	G			0	0	1	1	1	1	2	3
5	O				0	0	0	0	0	1	2
6	A					0	0	0	0	1	2
7	A						0	0	0	1	2
8	O							0	0	1	1
9	G								0	0	0
10	כ									0	0

One Possible Structure With 4 Base-Pairs



01/17/12

dG = -3.90 [Initially -3.40] ACGGCAACGU

Online Java Tool Using the Nussinov Algorithm

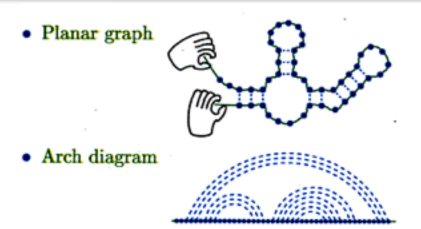
Nussinov algorithm - Ultrastudio.org

Visualization of Secondary Structure

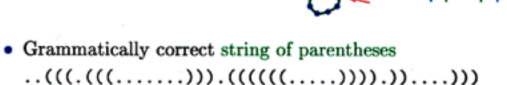
- Standard
- Circular
- Dot Plot

From Ralf Bunschuh at:

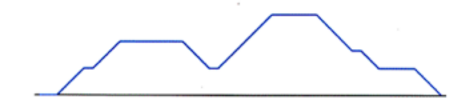
http://online.itp.ucsb.edu/
online/infobio01/bundschuh1/



Pseudo-knots neglected
 ⇒ arches do not cross

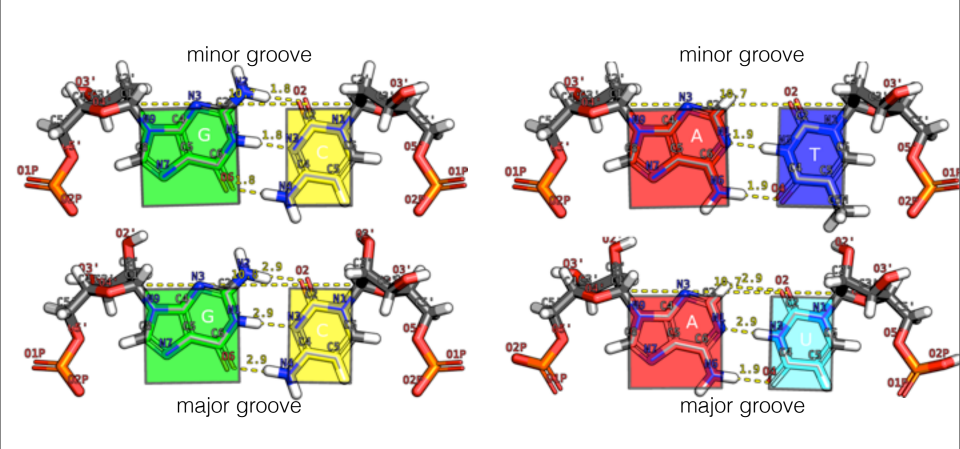


Mountain diagram



RNA 3D STRUCTURE

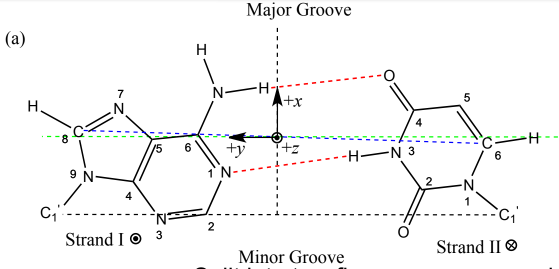
NDB/PBD Color Convention



Purines aka **R** = Guanine (green) and Adenine (red)
Pyrimidines aka **Y** = Citosine (yellow), Thymine (blue) and Uracil (cyan)

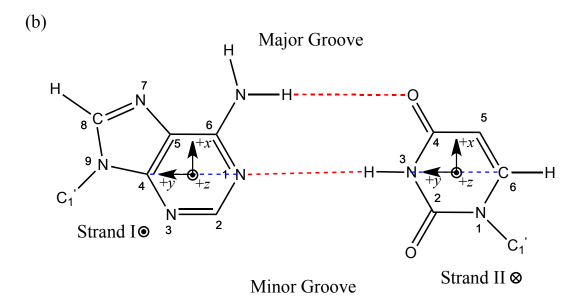
Standard Reference Frame

new **3DNA, Curves+** 1999-2001

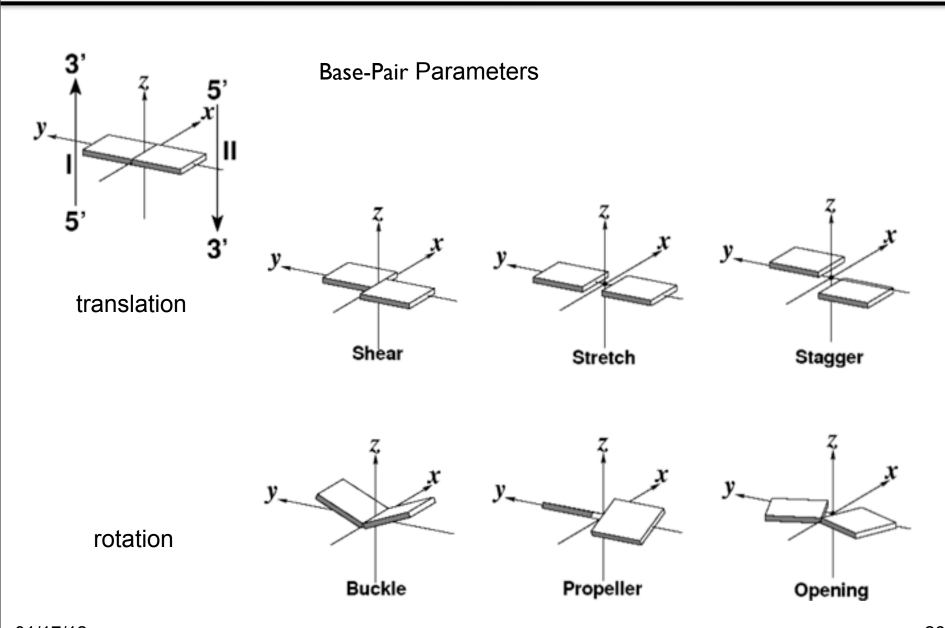


Split into two figures move up and include reference

old SCHNAaP 1997

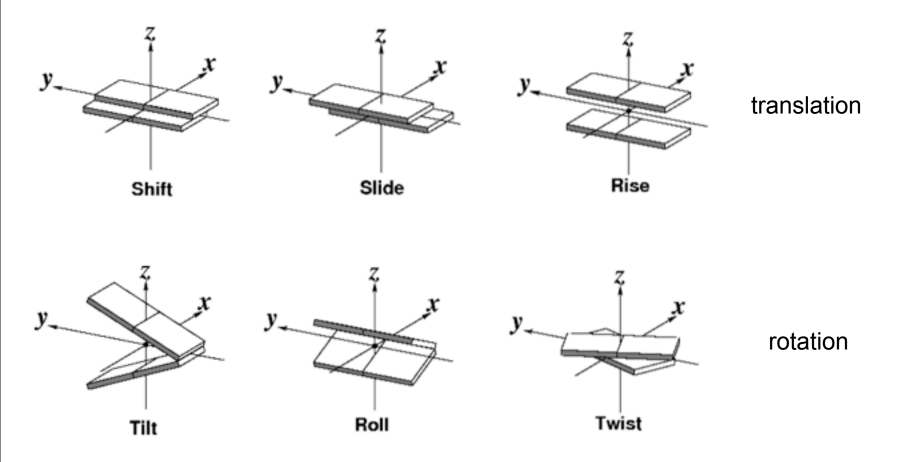


Calladine-Drew Rigid Block Models (Base-Pairs)



Rigid Block Model Base-Pair Steps

Base-Pair-Step Parameters



Software to Compute Rigid-Body Parameters

NEW

- 3DNA (Xiang-Jun Lu @ Columbia, Bussemaker lab.)
 - http://w3dna.rutgers.edu
 - http://rutchem.rutgers.edu/~xiangjun/3DNA
- 3dnaV2, UNIX preferred (linux, OS-X, freebsd, cygwin "yikes!")
- Curves+ (Richard Lavery @ Universite d' Lyon)
 OLD
- FREEHELIX, RNA, SCHNAaP, SCHNArP, compDNA, NUPARM

What's a base-pair in 3DNA? (In misc_3dna.pra)

- The distance between the origins of the two bases (as defined by their standard reference frames) must be less than certain limit (15.0 Å by default) - otherwise, they would be too far away to be called a pair.
- The vertical separation (i.e., stagger) between the two bases must be less than certain limit (2.5 Å by default) otherwise, they would be stacking instead of pairing.
- The angle between the two base z-axes (i.e., their normal vectors) is less than a cut-off (65.0° by default).
- There is at least one pair of nitrogen/oxygen base atoms that are within a H-bonding cut off distance (4.0 Å by default).

Please Show Us What They Look Like.

Step-Parameters Make Analysis Easier

- Graphs of A-DNA vs. B-DNA Conformations. (Roll vs Twist)
- Graph of DNA-Bending.
- Graph of Nucleosome.

Base-Pair Databases

• http://bps.rutgers.edu for RNA.

http://rnasteps.rutgers.edu for RNA Steps.

http://3dnascapes.rutgers.edu for DNA.

Classic Papers on Secondary Structure

- Classics
 - Tinoco Uhlenbeck Nature 1971
- Books
 - Tinoco's Book
 - Bioinformatics Book Chapter by Mathews
- Online
- Recent Trends
- People

RNA Secondary Structure Prediction Groups

- Doug Turner University of Rochester (Turner Rules)
- David Mathews University of Rochester (Dynalign)
- Michael Zuker Rensselaer Polytechnic Institute (mfold)
- Ivo Hoffacker Wien (vienna)
- Peter Stadler Leipzig (vienna)
- Francois Major Montreal (mc-sym)
- Tamar Schlick NYU (graph-grammars)
- Ruth Nussinov Tel Aviv (1978 dynamic algorithm)
- Nacho Tinoco (1971 free-energy minimization)

Overview

PART I (Secondary Structure)

- Introduction
 - Motivation.
 - Levels of RNA Structure?
 - Secondary Structure Prediction
 - Representation
- Software
 - Prediction (mfold, vienna, dynalign)
 - Rendering (Annotation s2s, others.)
- Everyones Favorite FIKA!

Overview

PART II (Rigid Block Model for Nucleic Acids)

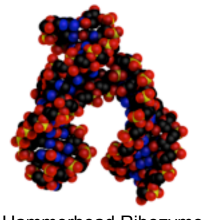
- NDB/PDB Color Conventions.
- 1991 Tsukuba Standard Reference Frame.
- Calladine-Drew Rigid Block Model.
- Software (3DNA(ansi-C), Curves+(fortran))

Overview

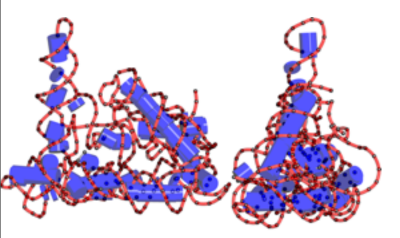
PART III (Practical Workshop-Tutorial)

- Practical Examples.
 - Using paper and pencil alone do Nussinov algorithm to some sequence. This gives you some structure, not necessarily minimal energy I guess.
 - Use computer to go to mfold, or vienna webserver to do prediction for same sequence. (maybe mfold is better because it gives more cartoons)
 - Make B-DNA, A-RNA, Z-DNA.
 - Chromosomal DNA. (download luger's get rid of protein analyze - rebuild.)
 - Download the ribosome of s. cerevisia. latest at 3.0
 - (What do the conformation of helices look like in ribosome?)

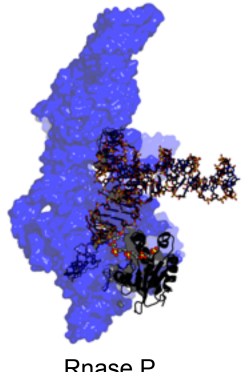
The RNA Zoo



Hammerhead Ribozyme 1hmh.pdb



GroupII Intron 3bwp.pdb



Rnase P