N Bases / Nucleosides / Nucleotides / Nucleic Acid Structures (Review)

Goals for this review unit:

- 1. Recognize the common building blocks of nucleic acids: names / 1letter abbrev.
- 2. Nomenclature for nucleosides and nucleotides (structure of ATP)
- 3. Primary structures of RNA and DNA
- 4. Conformations in DNAs
- 5. Characteristics of B-DNA, A-DNA and Z-DNA
- 6. Denaturation of DNA
- 7. Features of RNA / Functions of RNA
- 8. DNA Sequencing (Maxam Gilbert vs. Sanger Dideoxy)

iClicker Question #1

The approximate charge at pH 1 of the oligopeptide "H - A - P - P - Y - D - A - Y - S" is _____.

Answers: A) +3 B) +2 C) +1 D) -1 E) -2

iClicker Question #2

The world at 1,000,000 X (1 nm \rightarrow 1 mm)

At this magnification, double stranded DNA would be approximately _____ in width.

Answers: A) 1 mm B) 1 m C) 2 mm D) 20 m E) 5 mm

The Birth of Molecular Biology: DNA Structure

inside, linked together by hydrogen bonds. This structure as described is rather ill-defined, and for this reason we shall not comment



This figure is purely diagrammatic. The two ribbons symbolize the two phosphate—sugar chains, and the horizontal rods the pairs of bases holding the chains together. The vertical line marks the fibre axis on it.

We wish to put forward a radically different structure for the salt of deoxyribose nucleic acid. This structure has two helical chains each coiled round the same axis (see diagram). We have made the usual chemical assumptions, namely, that each chain consists of phosphate diester groups joining B-D-deoxyribofurances residues with 3',5' linkages. The two chains (but not their bases) are related by a dyad perpendicular to the fibre axis. Both chains follow righthanded helices, but owing to the dyad the sequences of the atoms in the two chains run in opposite directions. Each chain loosely resembles Furberg's² model No. 1; that is, the bases are on the inside of the helix and the phosphates on the outside. The configuration of the sugar and the atoms near it is close to Furberg's 'standard configuration', the sugar being roughly perpendicular to the attached base. There



Nature - 2001

Nature - 1953



The Nobel Prize in Physiology or Medicine 1962

"for their discoveries concerning the molecular structure of nucleic acids and its significance for information transfer in living material"



Francis Harry Compton Crick

(1/3 of the prize

United Kingdom

MRC Laboratory of Molecular Biology Cambridge, United Kingdom

b. 1916 d. 2004



James Dewey Watson

3 1/3 of the prize

USA

Harvard University Cambridge, MA, USA

b. 1928



Maurice Hugh Frederick Wilkins

(1/3 of the prize

United Kingdom and New Zealand

London University London, United Kingdom

b. 1916 (in Pongaroa, New Zealand) d. 2004



Left to right: Maurice Wilkins, John Steinbeck, John Kendrew, Max Perutz, Francis Crick and Jim Watson after the Nobel Ceremony in Stockholm in December 1962.



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A) GuanosineD) deoxyThymine MP

B) ThymineE) dexoyThymidine MP

C) Cytidine MP



dGuanosine MP

dCytidine MP

DNA primary structure



Rotational angles of phosphodiester chain



	B-DNA	A-DNA	Z-DNA C	Z-DNA G
$\alpha(\omega)$	-41	-90	138	100
$\beta(\phi)$	136	211	-94	- 108
$\gamma(\psi)$	38	47	80	-70
$\delta(\psi')$	139	83	48	- 130
$\varepsilon(\phi')$	- 133	- 185	180	- 140
$\zeta(\omega')$	-57	-45	- 170	56
x	78	27	20	- 100





Sugar pucker in DNA



Sugar pucker in DNA



Double stranded DNA



- Two single stranded DNA paired by Hydrogen bonds.
- Helical structure





Watson-Crick base pairs



G-C base pair



A-T base pair

HHMI: http://www.hhmi.org/biointeractive/dna/DNAi watson basepairing video.html

A and B Double Helices



(a) B-DNA, end-on view



(b) B-DNA, side view



(c) A-DNA, end-on view



(d) A-DNA, side view





Structure of double stranded DNA (B-DNA)



Right handed helix 10.5 residue per turn Helix pitch = 34Å Base pair tilt-helical axis = -6° Diameter = 20Å

Sugar pucker dA, dT, dC, dA: C2' endo

Glycosidic bond dA, dT, dC, dA: anti

Minor grove show base diversity

A-DNA vs. B. DNA



A - DNA 11 residue per turn Helix pitch = 28Å Major groove Base pair tilt = 20° $Diameter = 23\text{\AA}$ Sugar pucker C3' endo Glycosidic bond anti





Z - DNA

left handed helix 12 residue per turn Helix pitch = 45Å Base pair tilt = 7° Diameter = 18ÅSugar pucker dA, dT, dC: C2' endo dG: C3' endo Glycosidic bond dA, dT, dC: anti dG: syn

Denatuted DNA:

Heat denaturation of DNA is called "melting," The purine and pyrimidine bases exhibit very strong p-p* transitions around 260 nm. *E. coli* DNA absorption is only about 60% of that predicted from the weighted average spectrum based on its composition, this loss of intensity is called *hypochromism*. Since the absorpance goes up as DNA "unwinds", it can be used to monitor the unstacking of DNA.



RNA primary structure









The Proto-Ribosome: An Ancient Nano-machine for **Peptide Bond Formation**

Chen Davidovich,^[a] Matthew Belousoff,^[a] Itai Wekselman,^[a] Tal Shapira,^[a] Miri Krupkin,^[a] Ella Zimmerman,^[a] Anat Bashan,^{*[a]} and Ada Yonath^[a]



Figure 1. Top: the symmetrical region within the large ribosomal subunit. Bottom: Schematic representation of "pocket-like" proto-ribosome formation from a RNA precursor, showing simple catalytic peptidyl transferase activity.

Genetic information



\dots G T A C T G A A C G C A G G T \dots Genetic code

Human being: ~ 3,000,000,000 base-pairs ~ 30,000 – 40,000 Genes (Public Human Genome Project and Celera Genomics)

Chromosome



HHMI: http://www.hhmi.org/biointeractive/dna/DNAi_packaging_vo2.html

Sequencing DNA

Prior to the mid-1970's no method existed by which DNA could be directly sequenced. Knowledge about gene and genome organization was based upon studies of prokaryotic organisms and the primary means of obtaining DNA sequence was so-called reverse genetics in which the amino acid sequence of the gene product of interest is back-translated into a nucleotide sequence based upon the appropriate codons.

Maxam-Gilbert DNA Sequencing

•Sanger (didexoy) DNA Sequencing



"for his work on the structure of proteins, especially that of insulin"



Frederick Sanger

United Kingdom

University of Cambridge Cambridge, United Kingdom

Ь.1918



The Nobel Prize in Chemistry 1980

"for their contributions concerning the

determination of base sequences in

"for his fundamental studies of the biochemistry of nucleic acids, with particular regard to recombinant-DNA"



Paul Berg

1/2 of the prize

USA

b. 1926

Stanford University Stanford, CA, USA

nucleic acids"

Walter Gilbert 9 1/4 of the prize USA Harvard University, Biological Laboratories Cambridge, MA, USA

b.1932



Frederick Sanger

9 1/4 of the prize

United Kingdom

MRC Laboratory of Molecular Biology Cambridge, United Kingdom

b. 1918

Maxam-Gilbert DNA Sequencing



Figure 1. Chemical targets in the Maxam-Gilbert DNA sequencing strategy. Dimethylsulphate or hydrazine will attack the purine or pyrimidine rings respectively and piperidine will cleave the phosphate bond at the 3' carbon.

http://www.idtdna.com/support/technical/TechnicalBulletinPDF/DNA_Sequencing.pdf



Allan Maxam / Walter Gilbert DNA Sequencing

Sequencing single-stranded DNA

Two-step catalytic process:

1) Break glycoside bond between the ribose sugar and the base / displace base

Purines react with dimethyl sulfate

Pyrimidines react with hydrazine

2) Piperidine catalyzes phosphodiester bond cleavage where base displaced

- "G" dimethyl sulfate and piperidine
- "A + G" dimethyl sulfate and piperidine in formic acid
- "C" hydrazine and piperidine in 1.5M NaCI
- "C + T" hydrazine and piperidine





Figure 2. The Maxam-Gilbert manual sequencing scheme. The target DNA is radiolabeled and then split into the four chemical cleavage reactions. Each reaction is loaded onto a polyacrylamide gel and run. Finally, the gel is autoradiographed and base calling proceeds from bottom to top.

Maxam-Gilbert DNA Sequencing

- 200-300 bases of DNA sequence every few days
- Use large amounts of radioactive material, ³⁵S or ³²P
- Constantly pouring large, paper thin acrylamide gels
- Hydrazine is a neurotoxin

Early Benefits -

Discovery that the gene for ovalbumin in chicken and the gene encoding β -globin in rabbit contained non-coding gaps in the coding regions. These gaps< were flanked by the same dinucleotides in the two genes; GT on the 5' end of the gaps and AG on the 3' end of the gaps. Soon, the terms intron and exon were added to the genetic lexicon to describe the coding and non-coding regions of eukaryotic genes (1977).

Fred Sanger (dideoxy) DNA Sequencing

Primer strand

strand

Sanger knew that, whenever a dideoxynucleotide was incorporated into a polynucleotide, the chain would irreversibly stop, or terminate. Thus, the incorporation of specific dideoxynucleotides in vitro would result in selective chain termination.



Ribose Deoxyribose Dideoxyribose







Consider the following nucleic acid sequencing gel experiment using the Sanger dideoxy sequencing method:

What is the expected sequence (5' -> 3') of the original DNA sample assuming the primer was labeled with a 5'-prime fluorescent label?

C)



Advantages of dideoxy DNA Sequencing

- Elimination of dangerous chemicals (hydrazine)
- Greater efficiency

Taq polymerase makes DNA strands off of a template at rate of about 500 bases per minute
Chemical synthesis of a 25-mer oligonucleotide takes more than two hours.

→ High Throughput Methods (Human Genome Project)

Automated Fluorescence Sequencing

In 1986, Leroy Hood and colleagues reported on a DNA sequencing method in which the radioactive labels, autoradiography, and manual base calling were all replaced by fluorescent labels, laser induced fluorescence detection, and computerized base calling.



Figure 5. A. Chemical structure of the four succinylfluorescein dyes developed at DuPont. B. Normalized fluorescence emission spectra for each of the four dyes following excitation at 488nm. Shifts in the spectra were achieved by changing the side groups R₁ and R₂.

Automated DNA sequencing



bands migrate past detector





International sequence databases exceed 100 gigabases

In August 2005, the INSDC announced the DNA sequence database exceeded 100 gigabases. GenBank is proud of its contributions toward this milestone. We thank all the scientists who have worked through the submission process at GenBank and made their sequence data available to the world. See the related press release.

>100,000,000,000 bases



> 200,000 organisms!!