IVI U N I PHARM

Molecular Biology -2

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The structure and function of biopolymers during the transitions of genetic information

Department of Molecular Biology and Pharmaceutical Biotechnology (45-308)

Mgr. Marie Brázdová, Ph.D. (FAF, BFU AVČR) Mgr. Denis Šubert (BFU AVČR) Mgr. Daniel Renčiuk (BFU AVČR) PharmDr. Jakub Treml, Ph.D. (FAF)

[brazdovam@pharm.muni.cz,](mailto:brazdovam@vfu.cz)

Genetic information is coded by DNA

The experiment combining two strains of Streptococcus pneumoniae bacteria.

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Molecular Biology of the Cell (© Garland Science 2008)

Nuclein

- Nuclein acidic substance rich in nitrogen and phosphorus
- J. F. Miescher 1864
- Isolated from blood of wounded patients and cleaved by pepsin (proteolytic enzyme)

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Roles of genetic material

Genotype – storage of genetic information and its transition to the offspring

Phenotype – expression of genetic information to particular properties of an individual

Evolutionary role – adaptation of an organism/species to the environment through the changes in genetic information

Terminology

Gene – several "definitions" depending on the point of view:

classic genetics (Mendel) – elementary unit of hereditary genetic information molecular genetics – part of DNA coding for RNA (and as a consequence coding for some property of the individual)

structural genes coding for mRNA/protein (+ regulatory regions)

genes coding for functional RNA (miRNA, …)

strict – structural gene – part of DNA that codes for protein sequence

Allele – particular variant of the respective gene

Genome – complete DNA of organism (molecular) x complete genetic information of organism x sum of genes (classic)

Genotype – the combination of particular alleles of all genes in individual

Phenotype – the sum of actual individual properties (as a result of expression of particular genotype in the respective environment)

Genophore – the carrier of genetic information, usually a molecule of DNA (often used for bacteria)

Information biopolymers

Deoxyribonucleic acid (DNA)

• linear heteropolymer composed from 2-deoxyribonucleotides connected by phosphodiester bonds

• usually as a stable and resistant double helix

• serves as a storage of genetic information, as a template for its reproduction (replication) and as a template for the expression of genetic information to the phenotype (transcription)

Ribonucleic acid (RNA)

• linear heteropolymer composed from ribonucleotides connected by phosphodiester bonds

• usually as a single-stranded structure of variable length, structure and reactivity

• many functions depending on type of RNA

Protein

• linear heteropolymer composed from 20 (21) amino acids connected by peptide bonds

• highly variable structures, properties and functions

Central dogma of molecular biology

Gene

Genes

Most **eukaryotic** genes contain **introns**, that are transcribed into primary RNA transcript and introns are consecutively removed by **splicing** process on **spliceosome** to form final **mRNA**.

Prokaryotic genes do not contain **introns** and they are directly transcribed into mRNA that serves as a template for translation into protein.

Exon = coding region

Nucleic acids

5-carbon sugar - pentose

Base

Nucleic acid nomenclature

Modified bases

5-hydroxymethyl cytosine14

N

O[.]

N´

R

 $NH₂$

OH

 CH_2

Conformation of N-glycosidic bond

range of angles that depend on sugar pucker and base. The syn MB conformations are usually found with $\chi = 45^\circ \pm 45^\circ$; anti-
conformations are usually found with $\chi = -135^\circ \pm 45^\circ$.

Formation of sugar-phosphate backbone

REPLICATION DNA-dependent DNA polymerase

TRANSCRIPTION DNA-dependent RNA polymerase

REVERSE TRANSCRIPTION RNA-dependent DNA polymerase

Base reactivity – hydrogen bonds

Hydrogen bond – weak electrostatic interaction of two polar groups – one covalently binds hydrogen (DONOR – usually -N-H a -O-H); the second (ACCEPTOR) is usually N or O Length: \approx 2.8 A (2 - 3,4 A) Energy: < 1 kcal/mol both depend on particular atoms and on the environment Hoogsteen Hogsteen Watson Crick **DONOR** H. . . . H H. . . . H N N **ACCEPTOR** N N N Hoogsteen N N O N Matson Crick R Watson.Crick R **Cytosine Adenine** O H O N N $CH₃$ H H N N N N R H O N R **Guanine Tymine** MB_2_202317

Watson-Crick base pairing

DNA double helix

DNA double helix

- **two** molecules (**strands**) of DNA
- the helix is **right-handed**
- the strands are **antiparallel** their 5´-3´ direction is opposite in the context of double helix direction
- similar content of purines and pyrimidines; content of $A = T$, $G = C$ (Chargaff rules)
- result the strands are **complementary** i.e. according to the Watson-Crick base pairing rules we can predict the sequence of one strand according to the sequence of the other
- on average the double helix contains **10,5 base pairs** per turn of the helix, which is about **3,4 nm** in length

DNA structure –Watson and Crick model

M. Wilkins R. Franklin E. Chargaff

MB_2_2023

No. 4356 April 25, 1953

NATURE

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(1920). (1920). M. S., Mou, Not. Roy. Astro. Soc., Goophys. Supp., 5, 285 (1949). $\frac{1}{2}$ and $\frac{1}{2}$ a

. Eleman, V. W., Arkin, Mat. Astron, Fysik, (Stockholm), 2 (11) (1905).

MOLECULAR STRUCTURE OF NUCLEIC, ACIDS

A Structure for Deoxyribose Nucleic Acid

 W^E wish to suggest a structure for the salt of deoxyribose nucleic acid (D.N.A.). This structure has novel features which are of considerable hiological interest.

A structure for nucleic acid has already been proposed by Pauling and Corey'. They kindly made their manuscript available to us in advance of publication. Their model consists of three intertwined chains, with the phosphates near the fibre axis, and the bases on the outside. In our opinion, this structure is unsatisfactory for two reasons: (1) We believe that the material which gives the X-ray diagrams is the salt, not the free acid. Without the acidic hydrogen atoms it is not clear what forces would hold the structure together, especially as the negatively charged phosphates near the axis will repel each other. (2) Some of the van der Waals distances appear to be too small.

Another three-chain structure has also been suggested by Fraser (in the press). In his model the phosphates are on the outside and the bases on the mside, linked together by hydrogen bonds. This structure as described is rather ill-defined, and for this reason we shall not comment

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 3-n-deoxyribofuranose 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 737

equipment, and to Dr. G. E. R. Deacon and the is a residue on each chain every 3-4 A. in the z-direction. We have assumed an angle of 36° between adiacent residues in the same chain, so that the structure repeats after 10 residues on each chain, that is, after 34 A. The distance of a phosphorus atom from the fibre axis is 10 A. As the phosphates are on

The structure is an ope is rather high. At low expect the bases to tilt become more compact. The novel feature of in which the two chair purine and pyrimidine b are perpendicular to the together in pairs, a sing hydrogen-bonded to a chain, so that the two z-co-ordinates. One of the other a pyrimidine hydrogen bonds are ma I to pyrimidine positi pyrimidine position 6. If it is assumed that structure in the most (that is, with the ket figurations) it is found

bases can bond together Franklin's X-ray photograph shows
(purine) with thymine (DNA's 'B'-form (1952)

In other words, if an a a pair, on either chain, then on these assumptions
the other member must be thymine; similarly for guanine and evtosine. The sequence of bases on a single chain does not appear to be restricted in any way. However, if only specific pairs of bases can be formed, it follows that if the sequence of bases on one chain is given, then the sequence on the other chain is automatically determined

It has been found experimentally^{2,4} that the ratio of the amounts of adenine to thymine, and the ratio of guanine to cytosine, are always very close to unity for deoxyribose nucleic acid.

It is probably impossible to build this structure with a ribose sugar in place of the deoxyribose, as
the extra oxygen atom would make too close a van der Waals contact.

The previously published X-ray data^{5,6} on deoxyribose nucleic acid are insufficient for a rigorous test
of our structure. So far as we can tell, it is roughly compatible with the experimental data, but it must be regarded as unproved until it has been checked against more exact results. Some of these are given. in the following communications. We were not aware of the details of the results presented there when we devised our structure, which rests mainly though not entirely on published experimental data and stereochemical arguments.

It has not escaped our notice that the specific pairing we have postulated immediately suggests a possible copying mechanism for the genetic material.
Full details of the structure, including the conditions assumed in building it, together with a set of co-ordinates for the atoms, will be published elsewhere.

We are much indebted to Dr. Jerry Donohue for constant advice and criticism, especially on interatomic distances. We have also been stimulated by a knowledge of the general nature of the unpublished experimental results and ideas of Dr. M. H. F. Wilkins, Dr. R. E. Franklin and their co-workers at

DNA structure – Pauling model

Linus Pauling

A PROPOSED STRUCTURE FOR THE NUCLEIC ACIDS

BY LINUS PAULING AND ROBERT B. COREY

GATES AND CRELLIN LABORATORIES OF CHEMISTRY, * CALIFORNIA INSTITUTE OF TECHNOLOGY

Communicated December 31, 1952

The nucleic acids, as constituents of living organisms, are comparable in importance to the proteins. There is evidence that they are involved in the processes of cell division and growth, that they participate in the transmission of hereditary characters, and that they are important constituents of viruses. An understanding of the molecular structure of the nucleic acids should be of value in the effort to understand the fundamental phenomena of life

CHEMISTRY: PAULING AND COREY PROC. N. A. S.

which are involved in ester linkages. This distortion of the phosphate group from the regular tetrahedral configuration is not supported by direct experimental evidence; unfortunately no precise structure determinations have been made of any phosphate di-esters. The distortion, which corresponds to a larger amount of double bond character for the inner oxygen atoms than for the oxygen atoms involved in the ester linkages, is a reason-

Plan of the nucleic acid structure, showing several nucleotide residues.

able one, and the assumed distances are those indicated by the observed values for somewhat similar substances, especially the ring compound S₃O₉, in which each sulfur atom is surrounded by a tetrahedron of four oxygen atoms, two of which are shared with adjacent tetrahedra, and two unshared. The O-O distances within the phosphate tetrahedron are 2.32 \AA (between the two inner oxygen atoms), 2.46 \AA , 2.55 \AA , and 2.60 \AA . The

Various types of double helix

B-DNA

• DNA in water/salt soulutions

A-DNA

Z-DNA

• DNA in crowding solutions • CpG sequences in crowding conditions

• RNA

Left handed Zig-zag step

Reversed Watson-Crick pairing

Base protonation

Cytidine

•base protonation might alter the base reactivity

• free bases have pK far from physiological

•pK of bases in DNA might be closer to pH 7.4

• cytosine in C_n sequences has pK~7 – cytosine i-motif

DNA double helix x ions / water

- phosphates in DNA backbone are negatively charged repulsion
- this is compensated by interaction with ions (Na⁺, K⁺, Mg²⁺, ...) or water (H-H bonds)

Stability of DNA double helix

-
-

• base stacking various various Tm increases with length and ions

Base-pair parameters in double helix

Types of nucleic acids

- **linear** (human chromosome) **x circular** (bacterial genome)
- **single-stranded** (most RNAs) **x double-stranded** (human DNA)

Superhelicity

have a decompacted large-scale structure

Superhelicity happen mostly as a result of transition of polymerase complex and unwinding of DNA (helicase, …) during replication and transcription.

Topoisomerases

- Enzymes that relax the superhelicity
- Topo I works on 1 DNA strand
- Topo II works on 2-strand DNA

Reactivity of bases with amino acids

Double-stranded NA: Interaction of Hoogsteen side with amino acid in major groove.

Reactivity of bases with amino acids

Single-stranded NA: Interaction of Watson-Crick side with amino acid.

Figure 2-17

Interactions involving two hydrogen bonds between amino acids and bases that take the place of Watson-Crick base pairing.

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Genome composition

Repetitive sequences - repeats

Some sequences in genome are **unique**, usually the genomic sequences (both coding and non-coding). In contrast, other sequences exist in many copies – **repetitive sequences (repeats)**. The length of repeat (microsatelites 2-6 bp x LINE 6-7000 bp), as well as the number of copies (several – 1.5M SINE in human) is highly variable.

Inverted repeats

⁵´ A--T ³´

Inverted repeats

Special types of repetitions - transposons

Interspersed repetitions with various lengths and number of copies.

 $LTR - long terminal repeat - 100 bp - 5 kbp - variant of retrotransposons$ LINE – long interspersed nuclear elements – up to 6 kbp – human> 500k copies - 3 types (L1, L2, L3) – only some L1 are able to transpose SINE – (Alu, ...) short interspersed nuclear elements – up to 500 bp – human \sim 1,5M copies

Loops and hairpins in RNA

 $m¹A$

G

Functional types of RNA

Hoogsteen pairing - triplexes

 $A^*A^{NP_2_2_2023}$

Hoogsteen pairing – G-quadruplexes

Guanine quadruplexes

$GGGN_nGGGN_nGGGN_nGGG$

(Huppert J.L., Chem Soc Rev, 2008)

(Biffi G., et al., Nat Chem, 2013)

Base reactivity

Hydrofobic bases with high ability to form hydrogen bonds are reluctant to be freely expressed into water environment around – if there is any chance to avoid this and lower the base exposition to the environment by any type of base pairing or base stacking, the bases tend to form a structure. Even the "single-stranded" RNA or DNA forms, in fact, compact structure with number of base pairs.

Packing of DNA into chromosome

44

Binding of DNA to a histone octamer

45

Folding of nucleosomes into 30 nm fiber

30 nm fiber binds to protein scaffold

 (B)

 $20 \mu m$

Chromosome

Centromere – here are the chromosomes connected to the system of cellular microtubules – important for chromosome segregation during cell division

Telomere – terminal part of chromatides that protect the end from being recognised as a double-strand break by a DNA repair machinery

Chromosome

Fully condensed chromosomes are present only during the cell division, otherwise they are more or less decondensed to a lower levels of structure, especially in transcriptionally active sites (**euchromatin**). Transcriptionally inactive parts of DNA, as well as repetitive regions or telomeres are much more condensed (**heterochromatin)**. Various types of chromatin differ in **epigenetic** markers of both DNA (5-methyl cytosine) and histones (methylation a acetylation).

Table 1-1 Some Genomes That Have Been Completely Sequenced

Genome size and gene number vary between strains of a single species, especially for bacteria and archaea. The table shows data for particular strains that have been sequenced. For eucaryotes, many genes can give rise to several alternative variant proteins, so that the total number of proteins specified by the genome is substantially greater than the number of genes.

Levels of structure of biopolymers

Genetic code

Set of rules that assign a sequence of aminoacids in the protein to the sequence of nucleotides in DNA or RNA.

BY PHILIP LEDER AND MARSHALL W. NIRENBERG

NATIONAL HEART INSTITUTE, NATIONAL INSTITUTES OF HEALTH Communicated by Richard B. Roberts, October 1, 1964

Previous studies utilizing randomly ordered synthetic polynucleotides to direct amino acid incorporation into protein in E. coli extracts indicated that RNA codewords corresponding to valine, leucine, and cysteine contain the bases (UUG).¹⁻⁴ The activity of chemically defined trinucleotides in stimulating the binding of a specific C¹⁴-aminoacyl-sRNA to ribosomes, prior to peptide bond formation,⁵ provided a means of investigating base sequence of RNA codewords and showed that the sequence of a valine RNA codeword is GpUpU.6

Properties of genetic code

• genetic code is based on **triplets** – one aminoacid in protein is coded by a sequence of three nucleotides in DNA (RNA)

> CGUGGUACGAUUGGAUGU **Arg Gly Thr Ile Gly Cys** mRNA Protein Triplet = Codon x anticodon = complementary sequence on particular tRNA that carries the respective aminoacid

• genetic code is **universal** – individual triplets code for the same aminoacid in almost all organisms (x mitochondria)

• genetic code is **degenerated** – one aminoacid might be coded by several different triplets (but the opposite is not true)

Genetic code

Reading frames

Genetic code is based on triplets – three possible ways of reading (**reading frames**), but only one is correct.

Genetic code

Although the genetic code is universal, the usage of particular codons, as well as the amount of particular tRNAs and aminoacyl transferases differ

Modification of synthetic genes for recombinant protein production according to the expression system used (Bacteria, human, …) might be highly beneficial.