

Institute of Biophysics

Department of Biophysical Chemistry and Molecular Oncology Centre of Biophysical Chemistry, Bioelectrochemistry and Bioanalysis



Chemical reactivity of nucleic acids Chemical methods in DNA studies DNA damage

Miroslav Fojta

Olsztyn-Lańsk, September 19th, 2007

Chemical reactivity of DNA

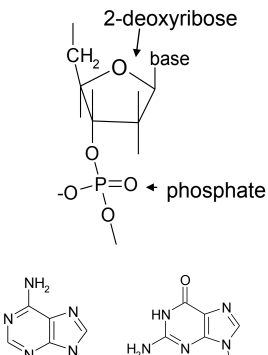
 NH_{2}

 DNA chemistry is derived from chemistry of its costituents

CH₃

ΗN

- phosphodiester bonds
- N-glycosidic bonds
- deoxyribose
- nitrogenous bases

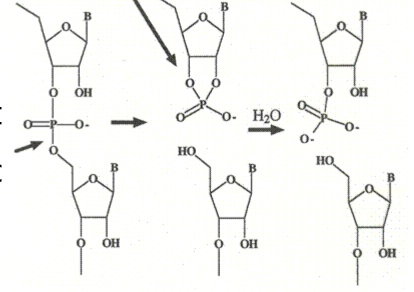


Chemical modification of DNA:

- damage to the genetic material
- analytical use

DNA hydrolysis

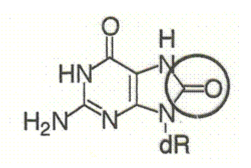
- both phosphodiester and N-glycosidic bonds susceptible to acid hydrolysis
- N-glycosidic bond more stable toward hydrolysis in pyrimidine than in purine nucleosides (and more in ribothan in deoxynucleosides)
- stable in alkali (unlike RNA)
- alkali-labile sites: upon DNA c
- enzymatic hydrolysis (N-glycc phosphodiesterases)



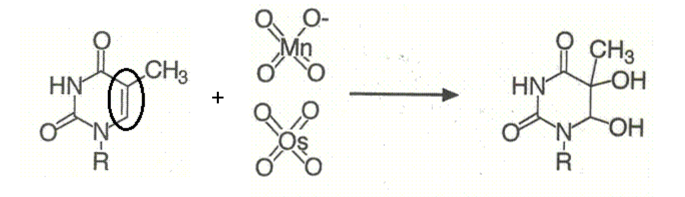
Oxidation

• two main sites susceptible to oxidation attacks:

– C8 of purines (ROS)

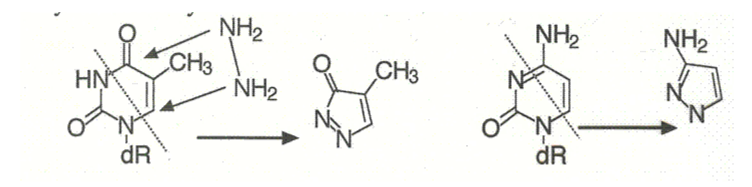


- C5-C6 of pyrimidines



reactions with nucleophiles

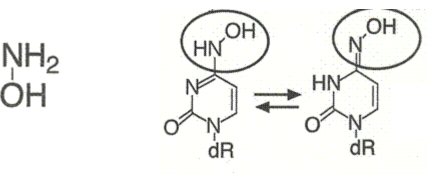
C4 and C6 are centres of electron deficit in pyrimidine moieties



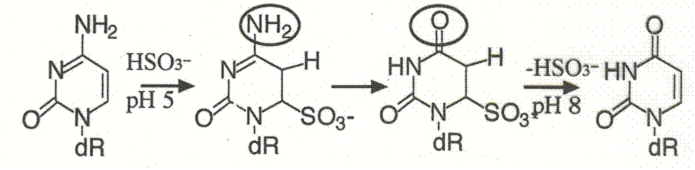
- reaction with hydrazine: pyrazole derivative and urea residue bound to the sugar
- with T the reaction is disfavored in high salt: Maxam-Gilbert sequencing technique

reactions with nucleophiles

- hydroxylamine: cytosine modification
- the products' preferred tautomer pairs with adenine \rightarrow mutagenic

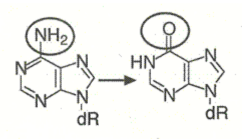


- bisulphite: cytosine modification inducing its deamination to uracil →mutagenic
- 5-methyl cytosine does not give this reaction: genomic sequencing of 5^mC

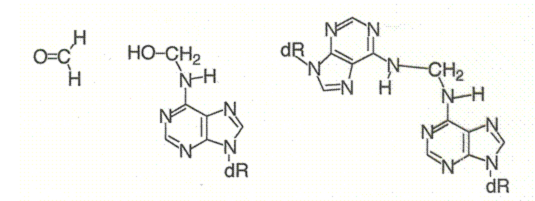


reactions with electrophiles

- attacking N and/or O atoms
- nitrous acid (HNO₂) causes base deamination (C→U, A→I) affecting base pairing, mutagenic

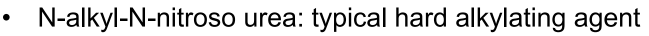


- aldehydes: reactions with primary amino groups
- formaldehyde: two step reaction

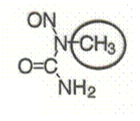


DNA alkylation

- hard or soft alkylating agents
- hard ones attack both N and O atoms, soft only N
- dimethyl sulfate: typical soft alkylating agent



- modifies all N + O in bases as well as phosphate groups (forming phosphotriesters)
- analytical use (sequencing, foorprinting)



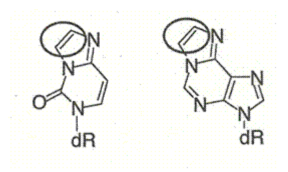
G-N7 >A-N1>C-N3>T-N3

1

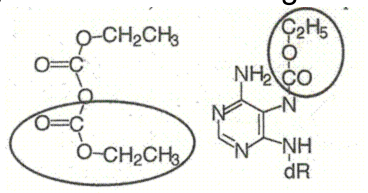
Biological consequences of base alkylation

- N-alkylation: the primary site = N7 of guanine (accessible in both ss and dsDNA)
 - does not change base pairing; easily repairable
- N3 of adenine or guanine: located in minor groove
 cytotoxic modification (DNA/RNA polymerization blocked)
- N1 of guanine: interferes with base pairing
- O-alkylation (G-O6, T-O6) the bases "locked" in enol forms → improper base pairing → mutagenic

- chloro- (bromo-) acetaldehyde: two reactive centres (aldehyde and alkylhalogenide)
- reaction with C or A
- chemical probes (react only with unpaired bases)



- diethyl pyrocarbonate: acylation of purines (primarily A) at N7
- modification leads to opening of the imidazole ring
- chemical DNA probing

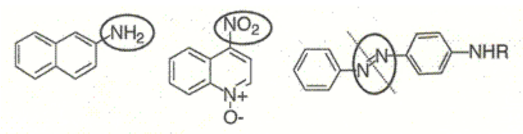


Metabolically activated carcinogens

- some substances became toxic after their metabolic conversion
- detoxifying machinery of the organism acts here as a bad fellow
- microsomal hydroxylase complex, cytochrome P450
- the role of this system is to introduce suitable reactive groups into xenobiotics enabling their conjugation with other molecules followed by removal from the organism
- but....

Metabolically activated carcinogens

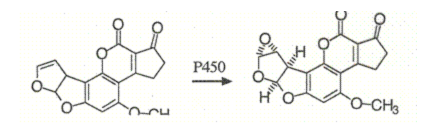
 aromatic nitrogenous compounds (amines, nitroor azo- compounds):

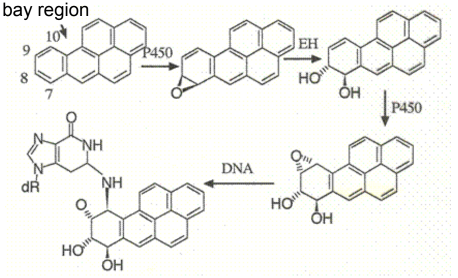


- aromatic amines are converted into either (safe) phenols, or (dangerous) hydroxylamine derivatives
- azo- compounds: "cleaved" into amines
- nitro- compounds: reduced into hydroxylamines

Metabolically activated carcinogens

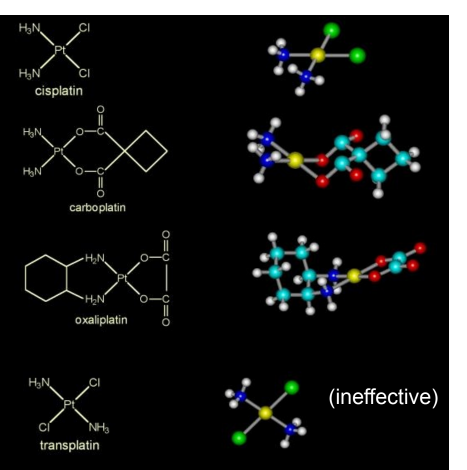
- polycyclic aromatic hydrocarbons like benzo[α]pyrene: three-step activation
 - P450 introduces epoxy group
 - epoxide hydrolase opens the epoxide circle
 - P450 introduces second epoxy group
- DNA adduct formation (primarily -NH₂ of guanine, then G-N7, G-O6 and A-N6)
- similar pathway of aflatoxin activation

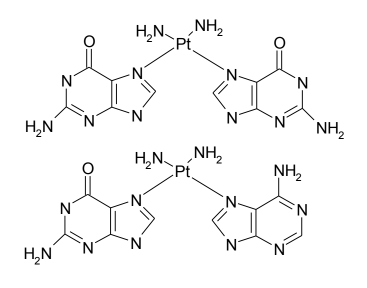




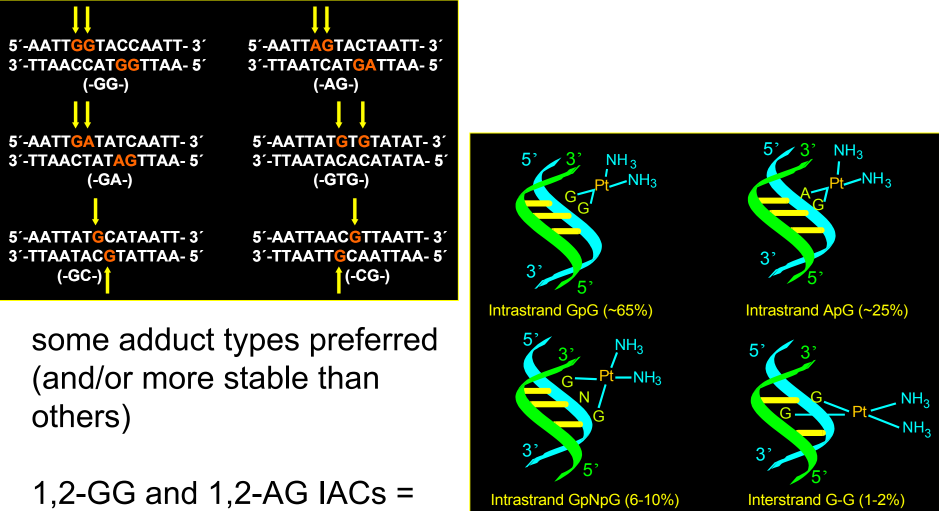
anticancer drugs

- some types of antineoplastic agents act via formation of DNA adducts
- metallodrugs: mainly platinum complexes



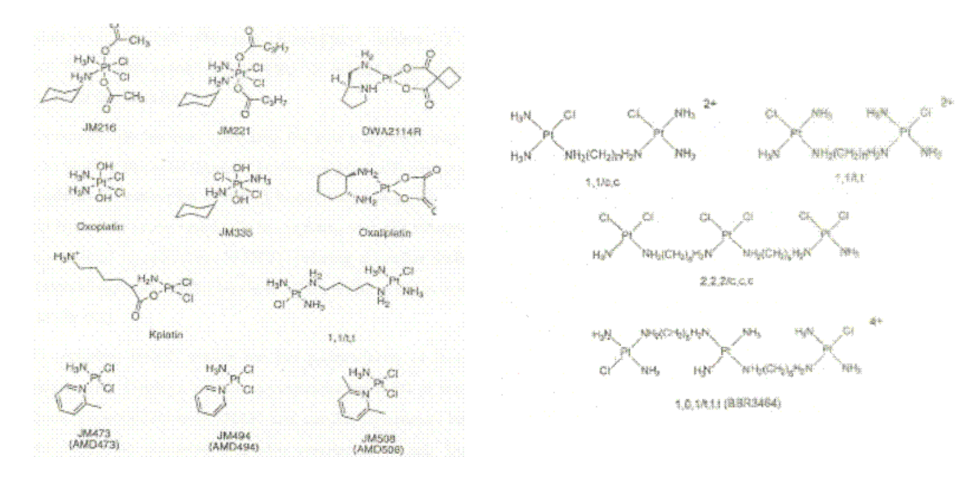


cisplatin: reaction with DNA in certain sequence motifs



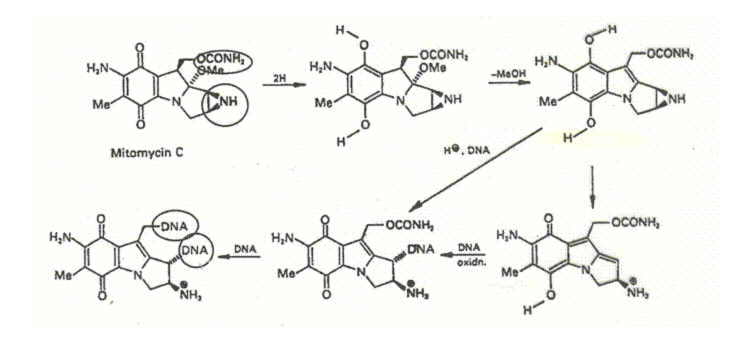
the main cytotoxic lesions

other platinum complexes tested as cytostatics



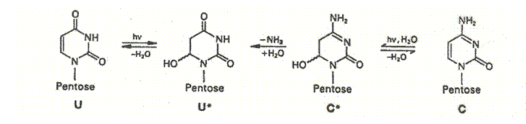
mitomycin C

- reactive aziridine group, quinone group
- reductive activation
- bifunctional adducts

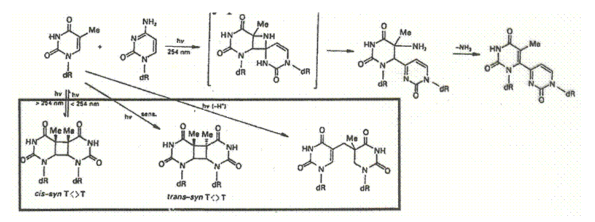


Photochemical DNA modifications

- mainly pyrimidines
- excitation at 240-280 nm: reactive singlet state
- water addition at C5-C6



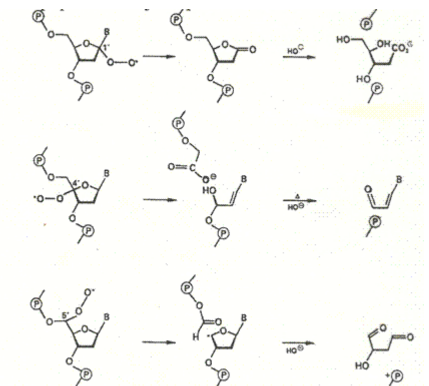
• excitation at 260-280 nm: photodimerization of pyrimidines



photoproducts of C can deaminate to U (mutagenic effects)

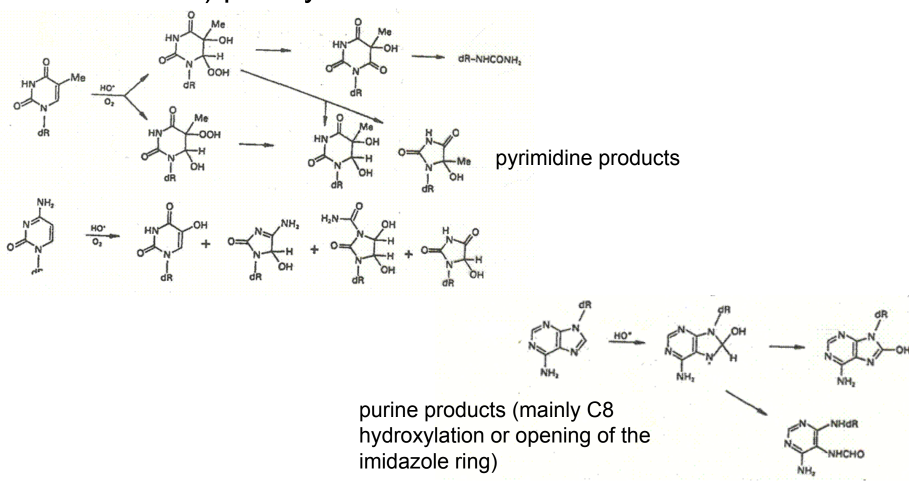
effects of ionizing radiation

- mostly indirect through water radiolysis
- each 1,000 eV produces ~27 •OH radicals that attack DNA
- sugar damage:abstraction of hydrogen atoms from C-H bonds
- a series of steps resulting in strand breakage



effects of ionizing radiation

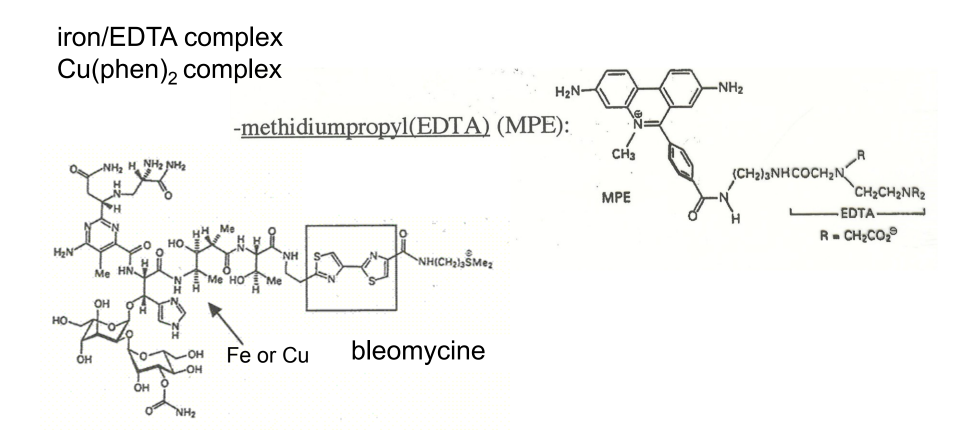
base damage: hydroxylation and/or (under aerobic conditions) peroxylation



chemical nucleases

species containing redox active metal ions mediating production of hydroxyl radicals (or othe reactive oxygen species) via Fenton and/or Haber-Weiss processes

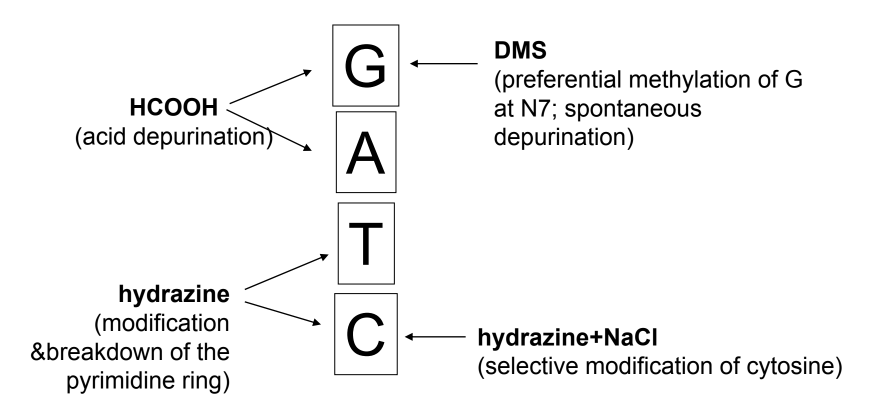
$$Me^n + H_2O_2 \rightarrow Me^{n+1} + \bullet OH + OH^-$$



Chemical approaches in DNA studies

(several examples)

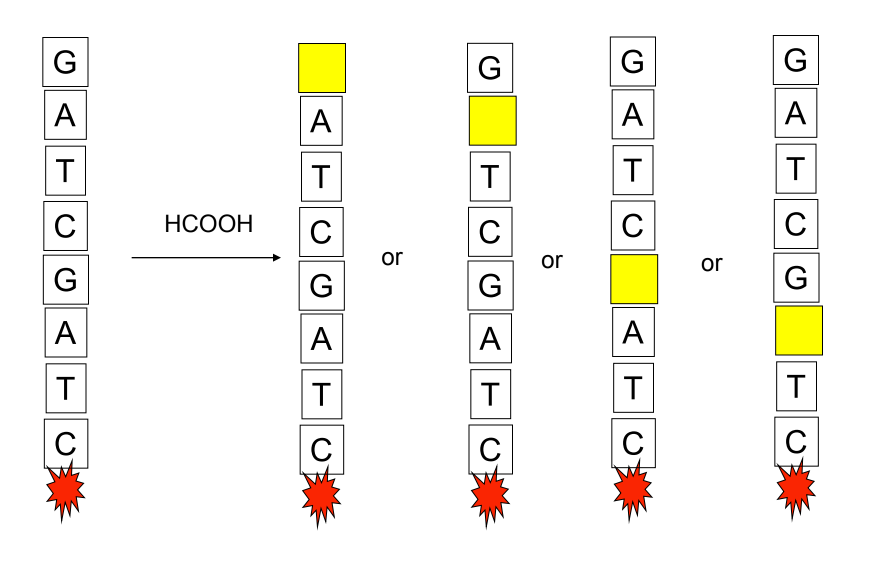
Maxam and Gilbert method of DNA sequencing

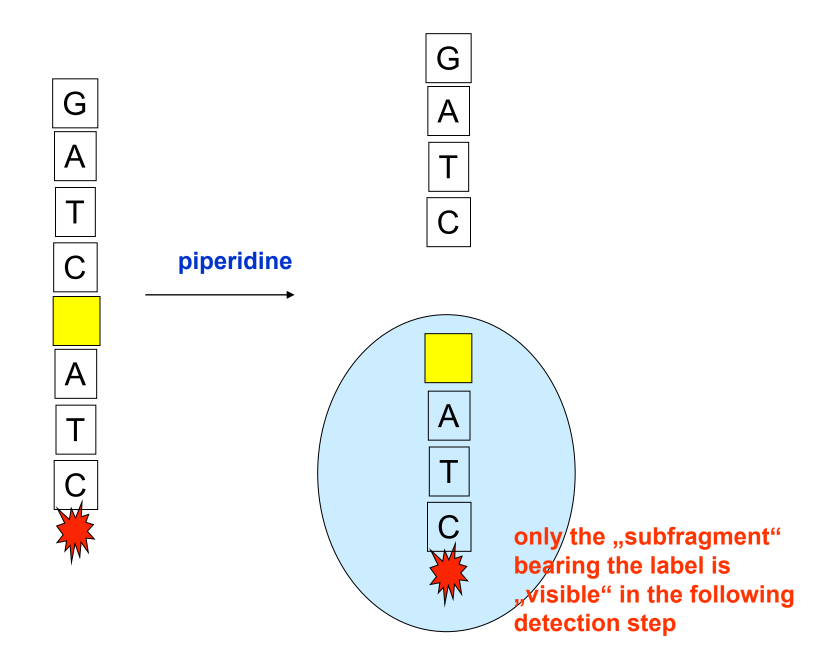


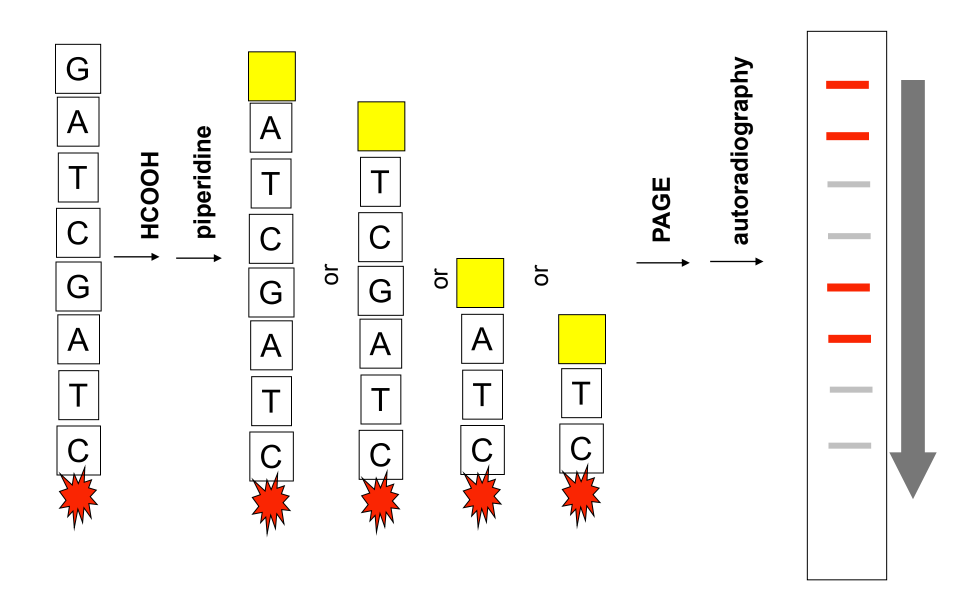
at sites of base modification (removal) the sugar-phosphate backobone is labile towards alkali

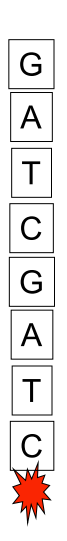
treatment with hot piperidine \rightarrow cleavage at such sites

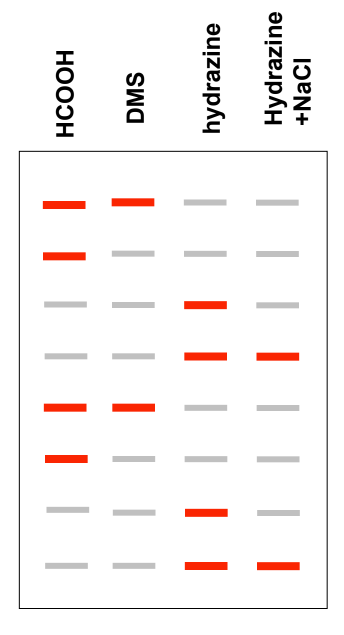
- DNA fragment is end-labeled (radionuclide, fluorophore)
- the sample is divided into four reactions (HCOOH, DMS, hydrazine, hydrazine + NaCI)
- the conditions are chosen to reach only one modification event per DNA molecule











Biochemistry: Maxam and Gilbert

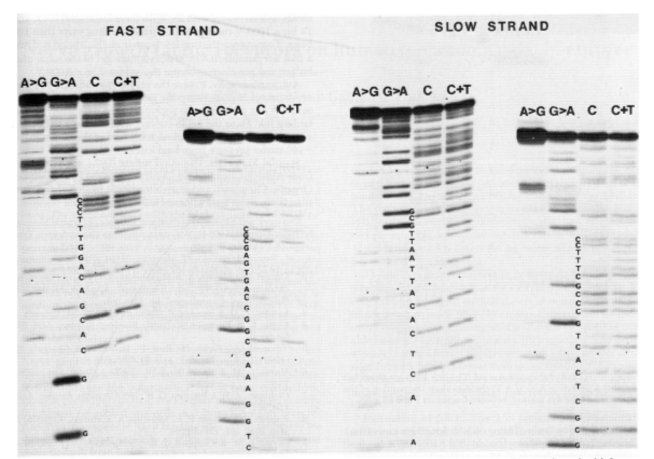


FIG. 2. Autoradiograph of a sequencing gel of the complementary strands of a 64-base-pair DNA fragment. Two panels, each with four reactions, are shown for each strand; cleavages proximal to the 5' end are at the bottom on the left. A strong band in the first column with a weaker band in the second arises from an A; a strong band in the second column with a weaker band in the first is a G; a band appearing in both the third and fourth columns is a C; and a band only in the fourth column is a T. To derive the sequence of each strand, begin at the bottom of the left panel and read upward until the bands are not resolved; then, pick up the pattern at the bottom of the right panel and continue upward. One-tenth of each strand, isolated from the gel of Fig. 1, was used for each of the base-modification reactions. The dimethyl sulfate treatment was 50 mM for 30 min to react with A and G; hydrazine treatment was 18 M for 30 min to react with C and T and 18 M with 2 M NaCl for 40 min to cleave C. After strand breakage, half of the products from the four reactions were layered on a $1.5 \times 330 \times 400$ mm denaturing 20% polyacrylamide slab gel, pre-electrophoresed at 1000 V for 2 hr. Electrophoresis at 20 W (constant power), 800 V (average), and 25 mA (average) proceeded until the xylene cyanol dye had migrated halfway down. Autoradiography of the gel for 8 hr produced the pattern shown.

Sangerovo ("dideoxy") sekvenování



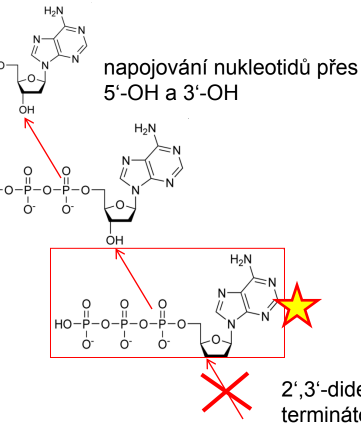
značení a sekvenování DNA

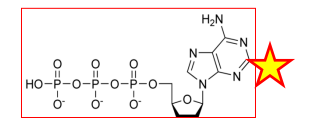


0 -0-P-0

0 || HO-P

F. Sanger

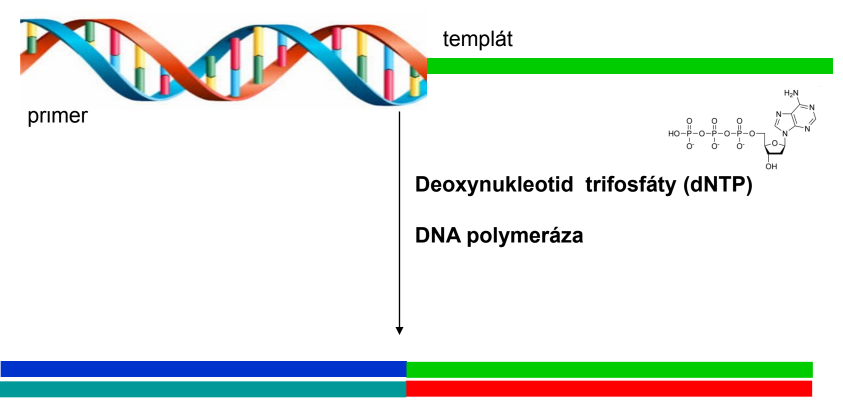




značené dideoxy: navázání značky podle komplementární báze

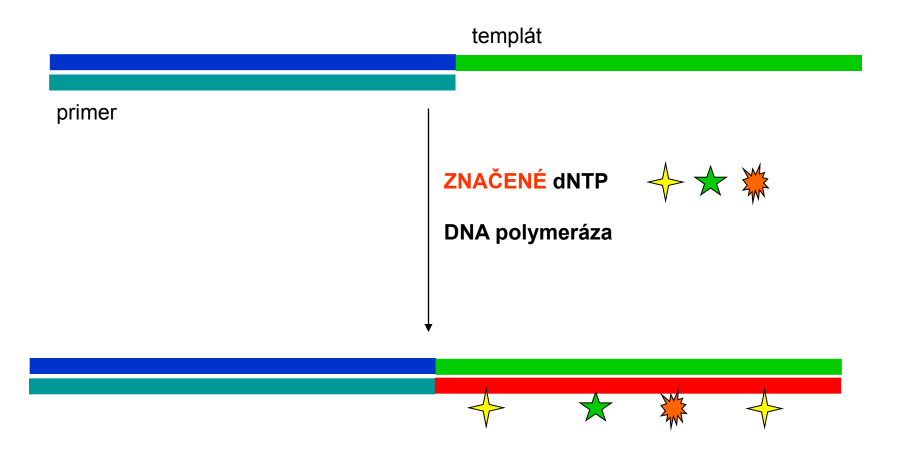
2',3'-dideoxynukleotidy: terminátory syntézy DNA (není kam napojit další nukleotid)

Syntéza DNA in vitro ("primer extension")

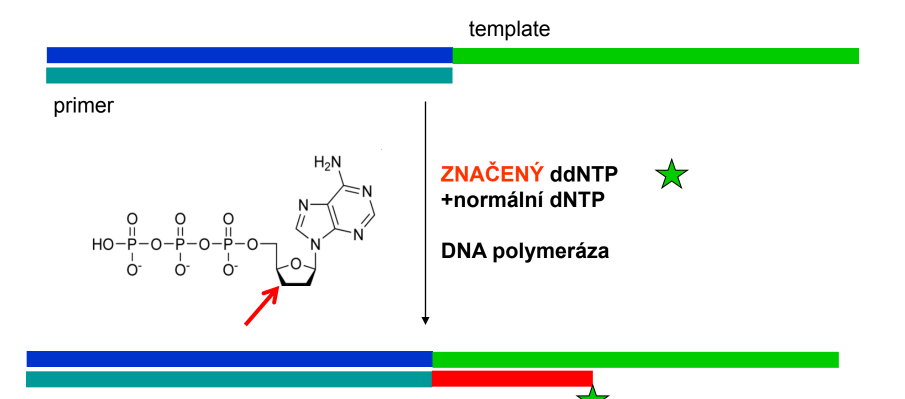


syntetizovaný úsek

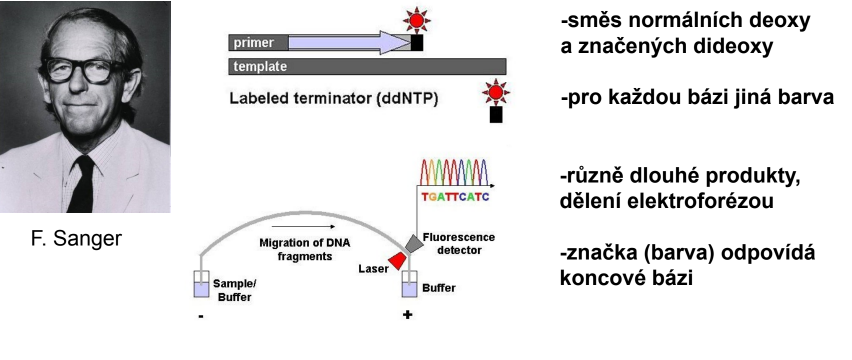
vložení značek do DNA pomocí DNA polymeráz

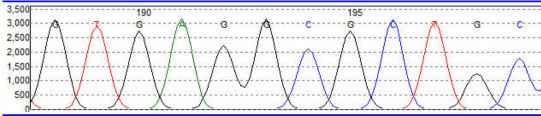


vložení značek do DNA pomocí DNA polymeráz



značení a sekvenování DNA

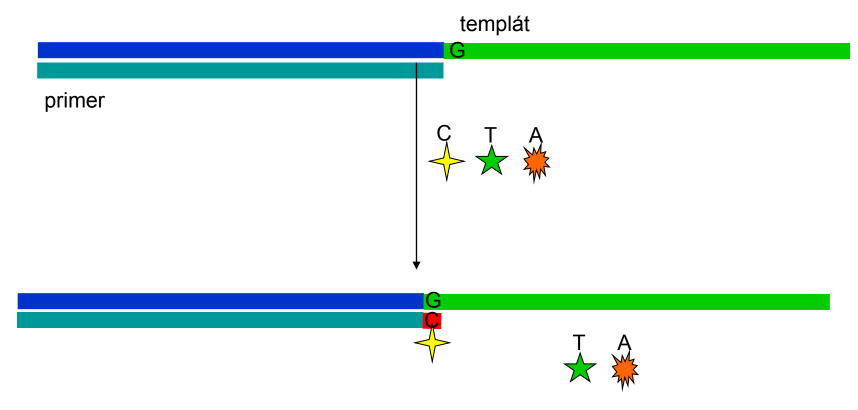




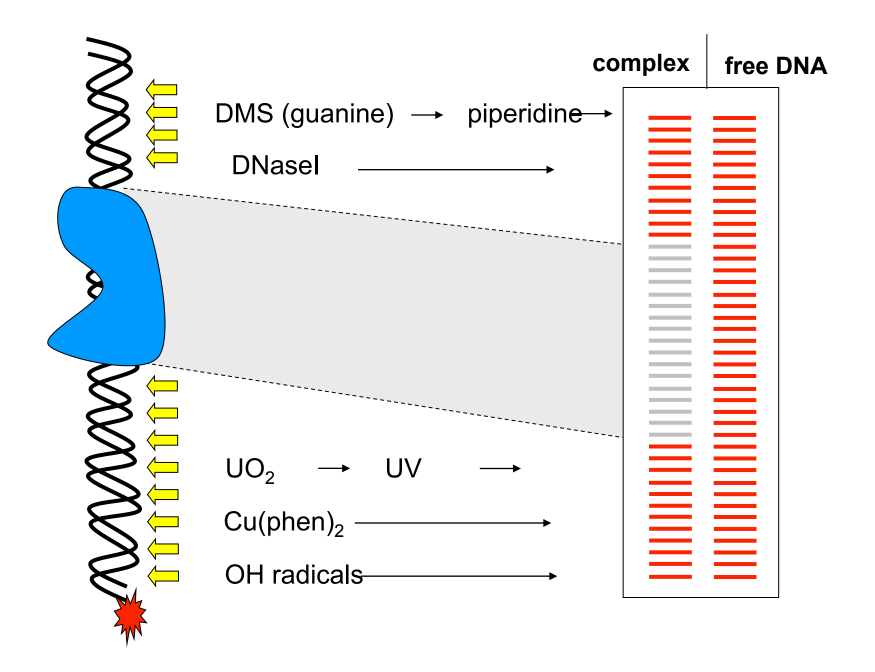
Detekce a identifikace mutací a polymorfismů

-důležité pro diagnostiku (určité mutace v určitých pozicích jsou spojeny s určitým onemocněním)

-zjišťuje se, jaká báze je v konkrétní pozici



DNA "footprinting": determination of binding sites of other molecules (e.g. proteins) at the DNA sequence level



single strand-selective chemical probes

Open local structures in negatively supercoiled DNA

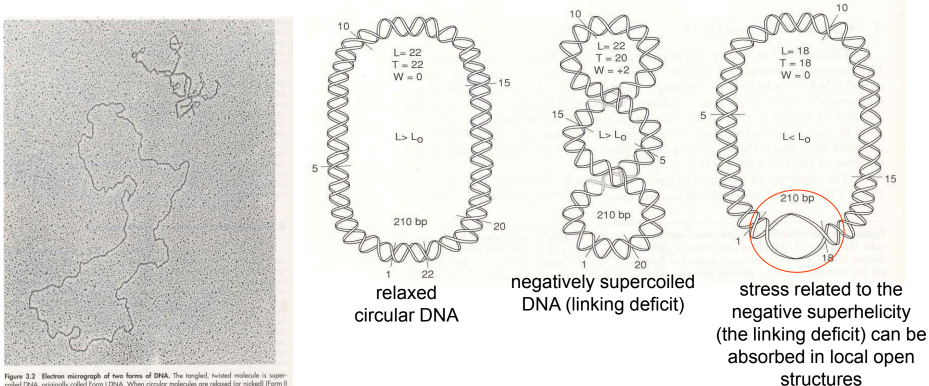
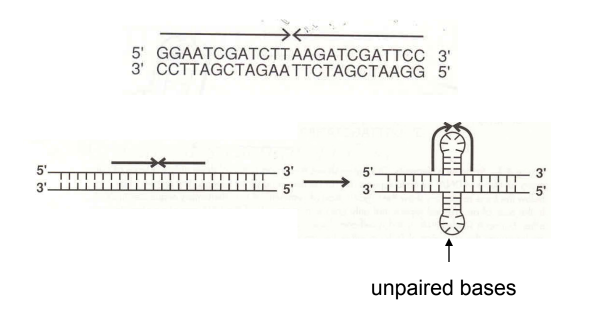


Figure 3.2 Electron micrograph of two forms of DNA. The tangled, twisted molecule is supercoiled DNA, originally called form I DNA. When circular molecules are relaxed (or nicked) (Form II DNA), they lose the twists. A linear molecule (not shown) is called Form III. The plasmid molecules shown are 9000 bp in length. Courtesy of Jack D. Griffith.

Open local structures in negatively supercoiled DNA

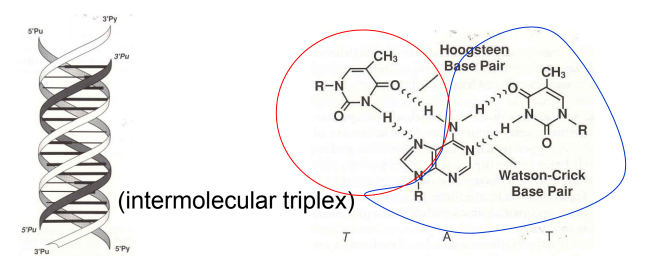
DNA segments of specific sequence can adopt "alternative" local structures

cruciform DNA (inverted repeat)



Open local structures in negatively supercoiled DNA

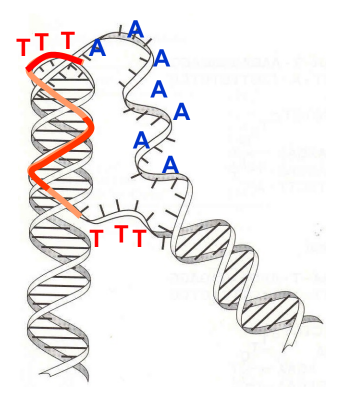
Triplex DNA (homopurine-homopyrimidine stretch with mirror symmetry)

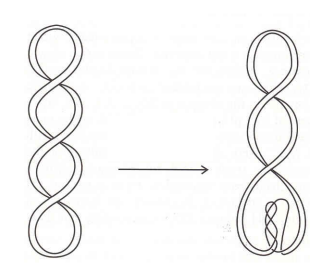


Otevřené lokální struktury v negativně nadšroubovicové (sc) DNA

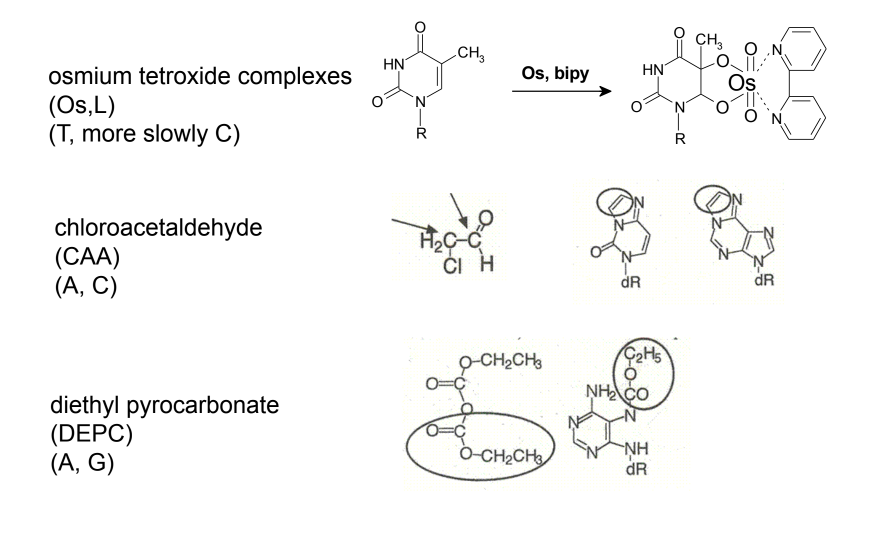
Intramolecular triplex

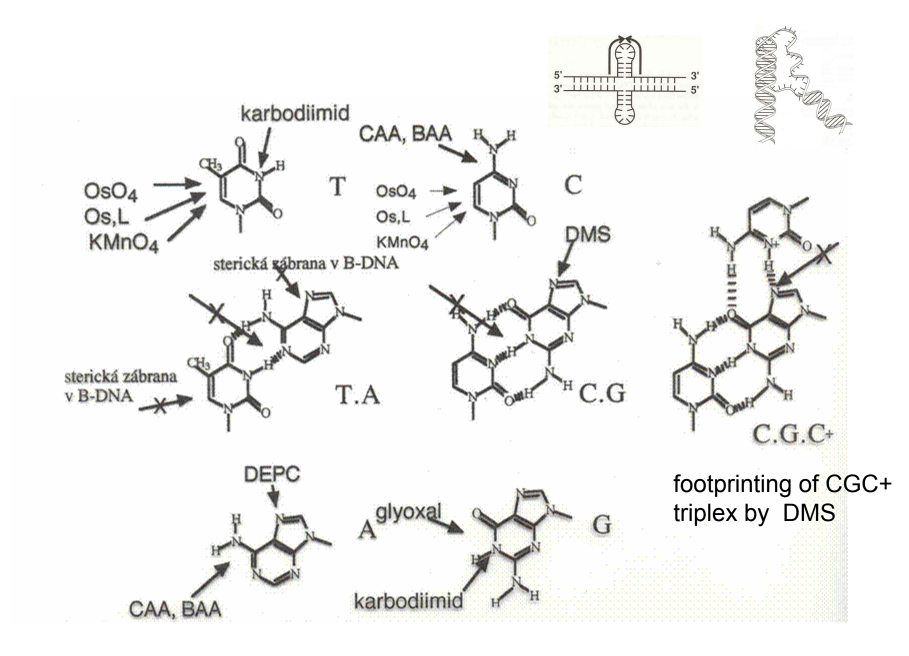
(homoPu•homoPy segment within negatively supercoiled DNA)





Chemicals selectively reacting with unpaired bases:

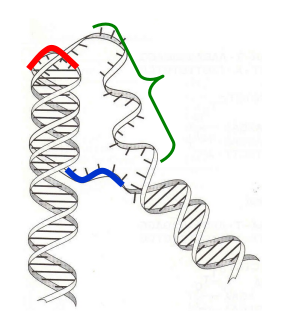




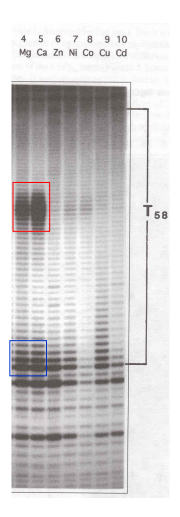
Using the Maxam-Gilbert technique, it is possible to determine with a high preciseness which nucleotides are forming the local structure

- ➤ modification of supercoiled DNA
- ➤ restriction cleavage, radiactive labeling
- ➤ hot piperidine
- ➤ sequencing PAGE

the structure can be deduced from the modification pattern







DNA damage and repair

Why is it important to study "DNA damage"?

DNA: the genetic material ensuring

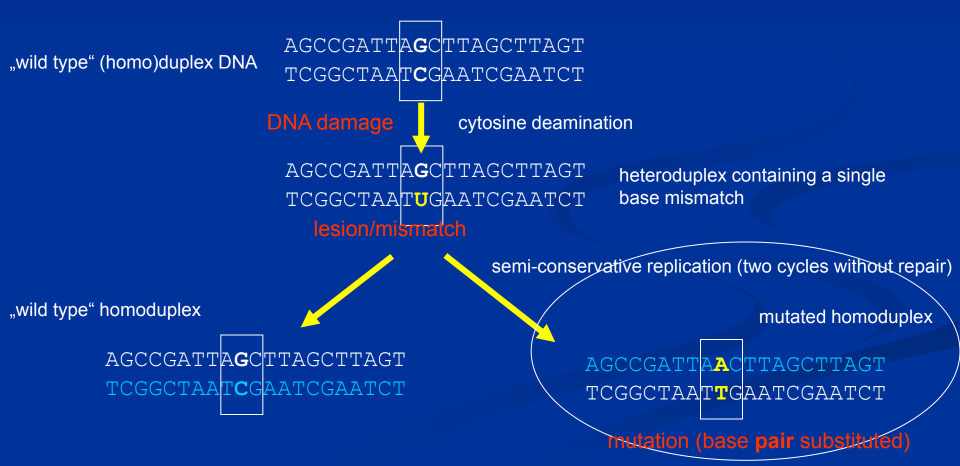
Damage to DNA may

- preservation of the genetic information
- its transfer to progeny
- its transcription and translation into proteins

- lead to change of the genetic information (mutation)
- affect gene expression
- have severe health impacts

DNA damage, mutation, lesion, mismatch...?

 mutation may arise from (among others) DNA damage which is not repaired prior to DNA replication, e.g..

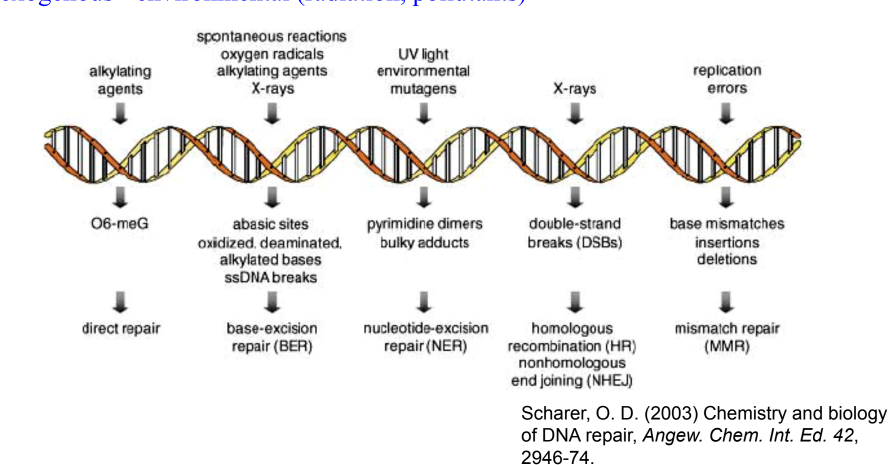


DNA damage, mutation, lesion, mismatch...?

- **mutations arise from unrepaired DNA damage** (or from replication errors)
- damaged DNA is not <u>mutated</u> yet! (damage is usually repaired in time i.e. before replication – lesions and/or mismatches are recognized by the reparation systems)
- DNA with <u>mutated</u> nucleotide sequence does not behave as <u>damaged</u>! All base pairs in such DNA are "OK" (no business for the DNA repair machinery) but the genetic information is (hereditably) altered.

DNA in the cells is permanently exposed to various chemical or physical agents

endogenous - products and intermediates of metabolism
exogenous - environmental (radiation, pollutants)



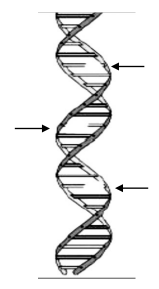
Most frequent products of DNA damage ("lesions")

interruptions of DNA sugar-phosphate backbone

interruption of the N-glykosidic linkage







single-strand break

double-strand break

abasic sites

reactive oxygen species
action of nucleases
consequence of base damage

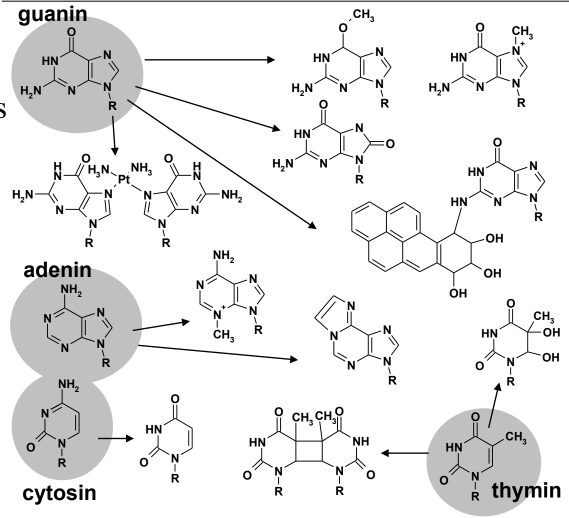
spontaneous hydrolysis
 (depurination)
 consequence of base
 damage

Most frequent products of DNA damage ("lesions")

base damage: chemical modifications

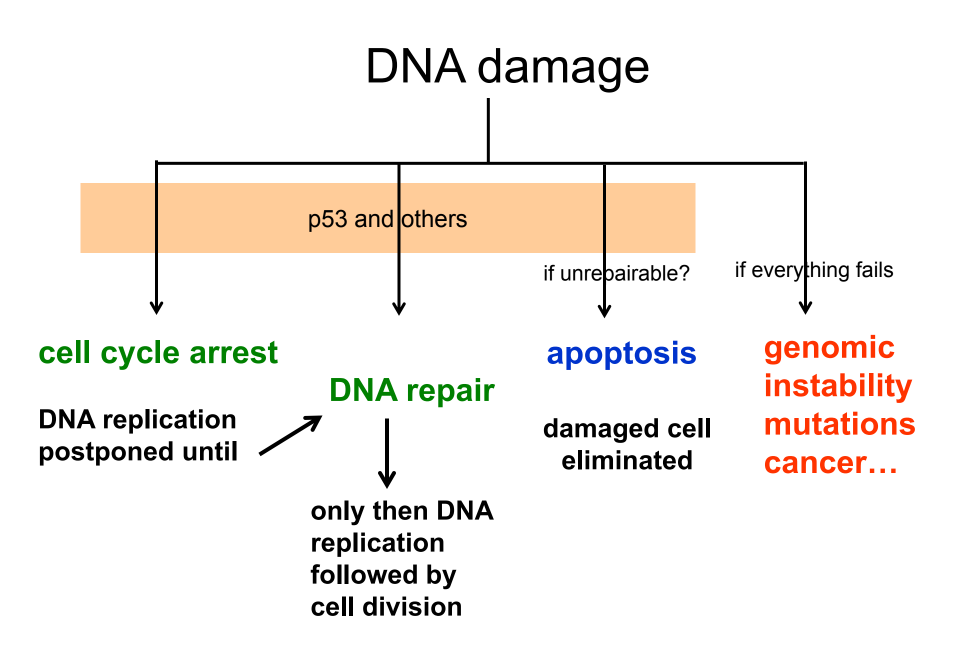
>alkylation
>oxidative damage
>deamination
>damage by UV radiation
(sunlight)
>metabilically activated
carcinogens

≻anticancer drugs



Importance of DNA repair

- estimated number of DNA-damage events in a single human cell: 10⁴-10⁶ per day!!
- only a small number of base pairs alterations in the genome are in principle sufficient for the induction of cancer
- DNA-repair systems must effectively counteract this threat
- in an adult human (10¹² cells) about 10¹⁶–10¹⁸
 repair events per day

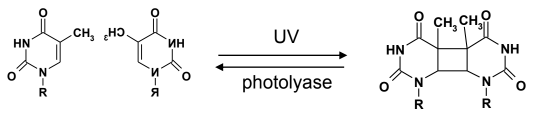


DNA repair pathways

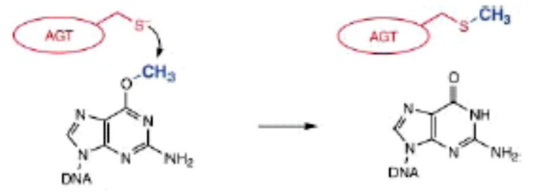
- direct reversal of damage
- base excision repair
- nucleotide excision repair
- mismatch repair
- repair of double strand breaks

Direct reversal of DNA damage

• photolyases: repair of cyclobutane dimers



 O6-alkylguanine transferase: reverses O6-alkylguanine to guanine by transferring the alkyl group from DNA to a reactive cysteine group of the protein

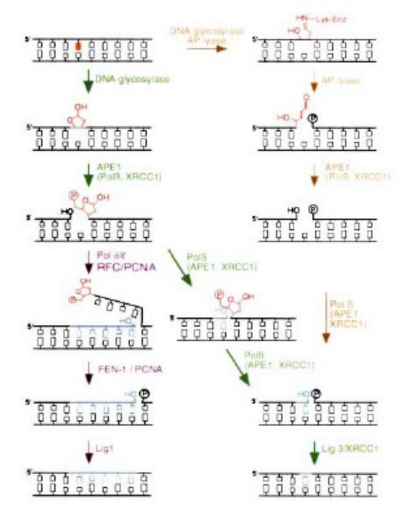


Base excision repair

- repair of damage by deamination (U, I), oxidation (8-oxoG), and alkylation
- initiated by DNA glycosylases, which recognize damaged bases and excise them from DNA by hydrolyzing the Nglycosidic bond
- substrate specificity of the glycosylases: developed to repair expectable "errors"?

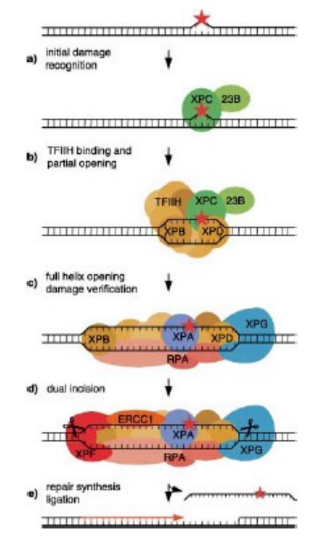
Enzyme	Most important substrate	AP lyase
UNG	U, 5-OH-U in ss/dsDNA	no
SMUG1	U, 5-OH-U in ss/dsDNA	no
• TDG	U:G, T:G, &C	no
MBD4	U:G, T:G	no
OCG1	8-oxoG:C, fapy	yes
 MYH 	A:8-oxoG	no
NTH 1	ox. pyrimidine, fapy	yes
NEI 1	ox. pyrimidine, fapy	yes
AAG (MPG)	3-MeA, 7-MeG, EA, Hx	no





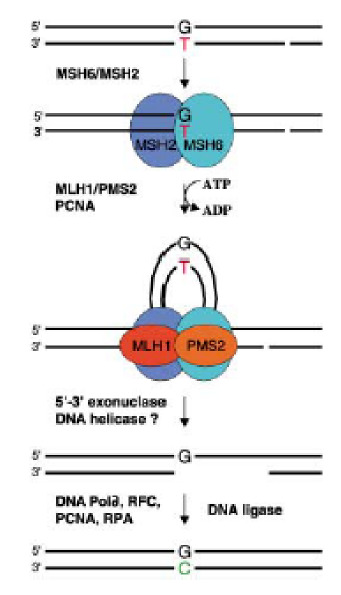
Nucleotide excision repair

- removes bulky base adducts (such as those formed by UV light, various environmental mutagens, and certain chemotherapeutic agents) from DNA
- broad substrate specificity: dealing with unexpected environmental DNA damaging agents
- excision of the damaged oligonucleotide
- then filling the gap & the sealing break



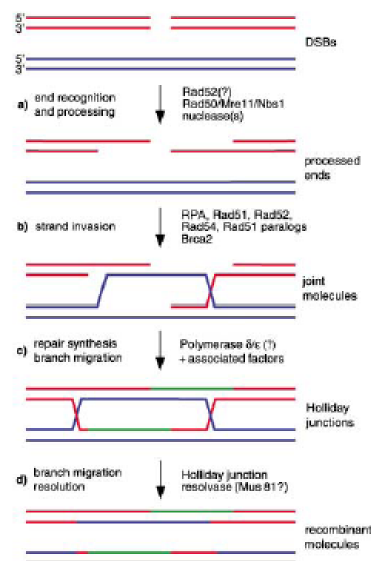
Mismatch repair

- dealing with replication errors
- polymerases introduce about one erroneous nucleotide per 10⁵ nucleotide; their 3'→5'exonuclease activity decreases incidence of the errors to 1:10⁷
- the MMR contributes to replication fidelity by a factor of 10³ by removal of base-base mismatches, insertions and deletions (hence the resulting incidence of mutations due to erroneous replication is only 1:10¹⁰)
- the system must be able discrimitate between parental and daughter DNA strand!
- MutS binds to mismatches and insertion/deletion loops
- "repairosome" formation, removal of a part of the daughter strand by 5'→3'- exonuclease
- new DNA synthesis and ligation



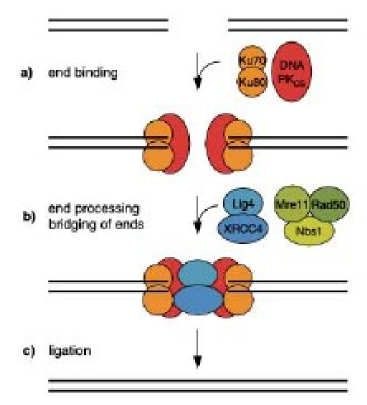
Repair of double strand breaks

- consequences of DSBs can be very severe (chromosome aberrations)
- two repair pathways:
- homologous recombination: an intrinsically accurate repair pathway that uses regions of DNA homology (such as sister chromatids) as coding information.



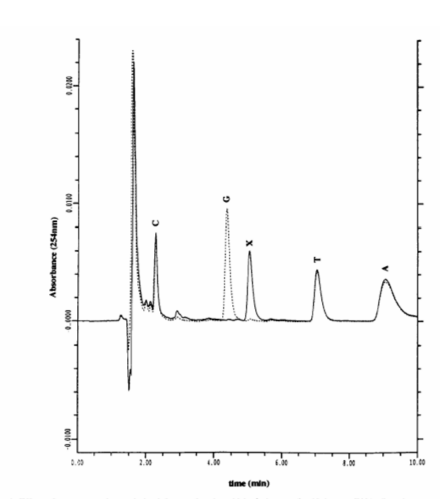
Repair of double strand breaks

- consequences of DSBs can be very severe (chromosome aberrations)
- two repair pathways:
- non-homologous end joining: conceptually simple pathway that involves the religation of broken ends (without using a homologous template
- less accurate: may loss of a few nucleotides at the damaged DNA ends

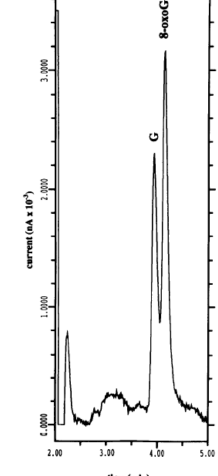


Examples of techniques used to detect DNA damage

 Techniques involving complete DNA hydrolysis followed by determination of damaged entities by chromatography or mass spectrometry





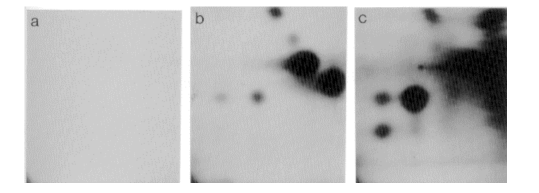


time (min)

Fig. 4. Effect of guanase on bases derived from a formic acid hydrolysate of calf thymus DNA. Samples were a HPLC with UV detection prior to (--) and following (--) guanase treatment as described in Materials and Methods. G, guanine; X, xanthine; T, thymine; A, adenine.

Fig. 1. Reversed-phase HPLC analysis, using electrochemical detection at +600 mV, of a solution containing 500 nM guanine (G) and 40 nM 8-oxoguanine (8-oxoG). Chromatographic conditions were as described in Materials and Methods except the mobile phase was 50 mM sodium acetate, 1 mM EDTA, pH5.1 containing 2% methanol.

³²P-postlabeling: analysis of base adducts



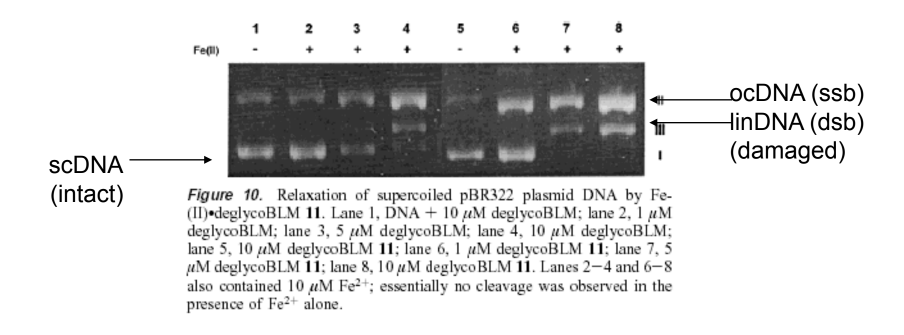
Enzymatic hydrolysis of DNA to 3'-mononucleotides using micrococcal endonuclease and spleen phophodiesterase Np + Np + Np + - - + Xp + Yp + Zp + - -Normal 3 mononucleotides Adducted 3'-mononucleotides Adduct enrichment by selective removal of normal 3'-mononucleotides a) nuclease P1 b) n-butanol extraction Xp + Yp + Zp + - - - - -Radiolabel adducted 3'-mononucleotides with ³²P *pXp + *pYp + *pZp + - - - -Multidimensional thin-layer chromatography of adducts on PEI-cellulose Autoradiography or storage phosphor imaging to determine the distribution of radioactivity on the chromatograms Quantitation of the radioactive areas on the chromatograms

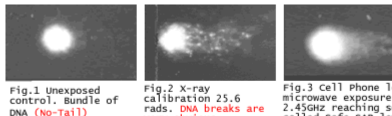
-Np-Np-Np-Np-Xp-Yp-Zpxenobiotic-modified DNA Techniques involving complete DNA hydrolysis followed by determination of damaged entities by chromatography or mass spectrometry

 Monitoring of changes in whole (unhydrolyzed) DNA molecules: electrophoretic and immunochemical techniques

detection of strand breaks:

relaxation (and/or linearization) of plasmid supercoiled DNA





,,comet assay" (dsb)

Fig.1 Unexposed control. Bundle of DNA (No-Tail) very obvious

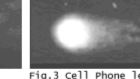
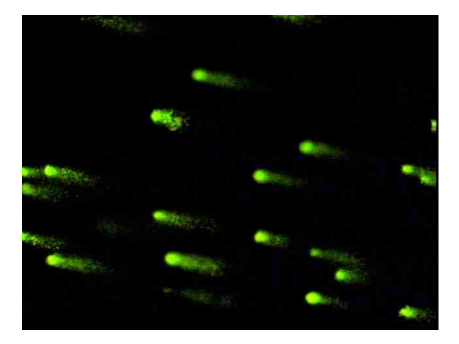


Fig.3 Cell Phone level microwave exposure 2hrs 2.45GHz reaching so called safe SAR levels Comet Tail = DNA Damage

"alkaline elution assay" (ssb + alkali-labile sites)

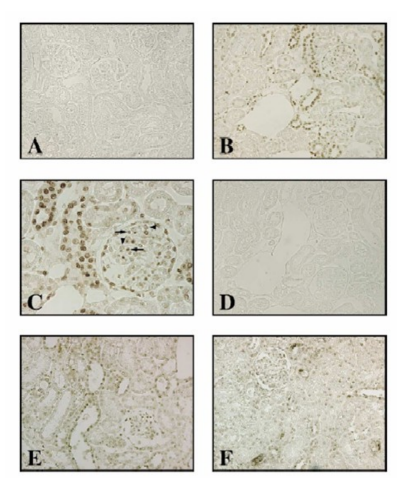


imunochemical techniques

when antibodies against the adducts available

►ELISA

▶In situ techniques



8-oxo guanine detection in situ in kidney tissue