# DNA REPLICATION, DNA MUTABILITY & DNA REPAIR

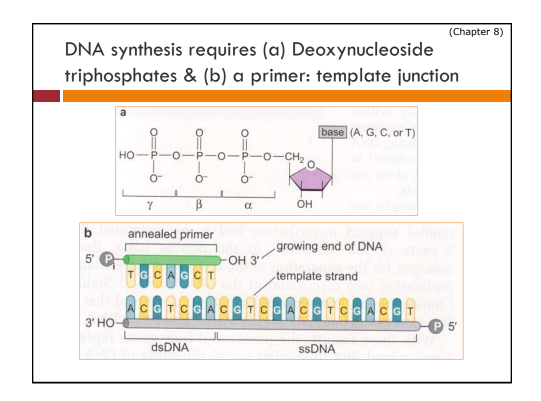
Course: Molecular Biology (BIOL 333)

Instructor: Dr Mahmoud Srour

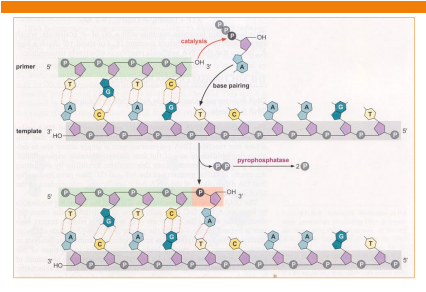
Textbook:

Watson J, et al. (2014). Molecular Biology of the Gene, 7<sup>th</sup> ed.

Chapters 9 and 10



# DNA is synthesized by extending the 3' end of the primer



What is the driving force for DNA synthesis?

- $\square$  XTP + (XMP)n  $\rightarrow$  (XMP)n+1 +  $P \sim P$ 
  - ( $\Delta G$  -3.5kCal/mole)
- $\square$  XTP + (XMP)n  $\rightarrow$  (XMP)n+1 +  $2P_i$

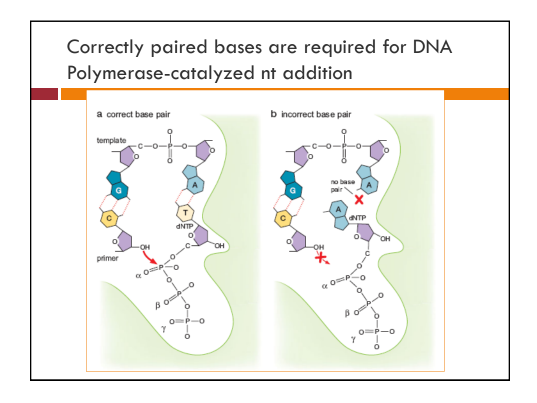
 $(\Delta G - 7kCal/mole)$ 

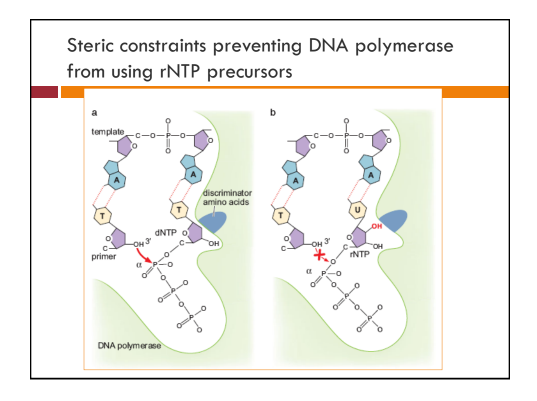
#### The Mechanism of DNA Polymerase

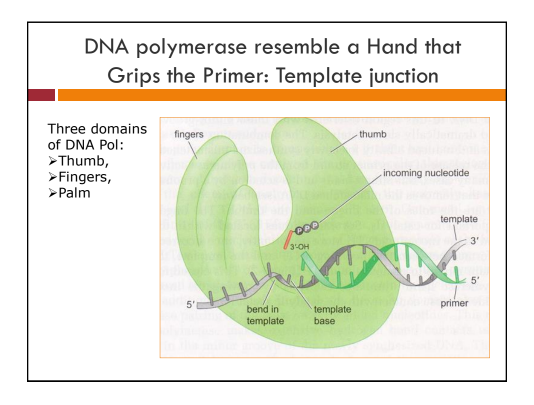
- Unlike most enzymes, DNA Pol uses a single active site that catalyze one reaction, i.e, the addition of any of four dNTPs, HOW?
- DNA Pol explores the nearly identical geometry of A:T and G:C base pairs, or it monitors the ability of incoming nt to form an A:T or G:C base pair, rather than detecting the exact that enters the active site
- □ Only when the correct base pair is formed are the 3'-OH or primer and 5"-P of incoming dNTP are in optimum position for catalysis to occur.

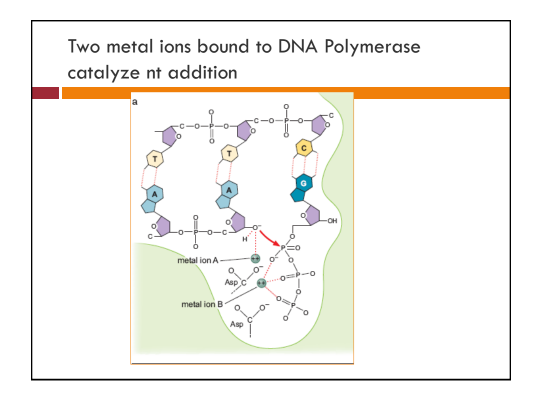
#### The Mechanism of DNA Polymerase (cont'd)

- □ The rate of incorporation of incorrect nt is as much as 10,000-fold slower than incorporation of the correct nt
- DNA pol can distinguish between dNTPs and rNTPs, although rNTPs are present at approx 10-fold higher conc in the cell. This is accomplished by steric exclusion of rNTPs from DNA Pol active site. The nt pocket is too small to allow the presence of 2'-OH of rNTP and this space is occupied by 2 amino acids that make van der Waals contact with sugar ring??

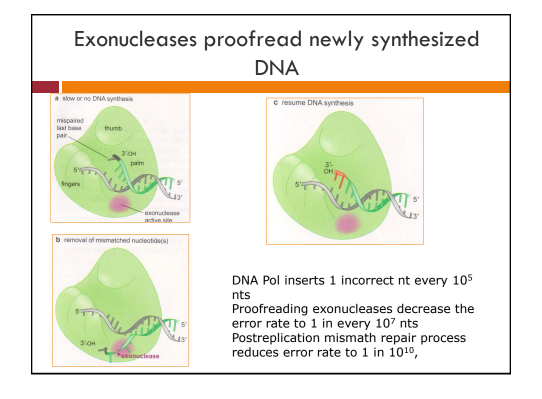




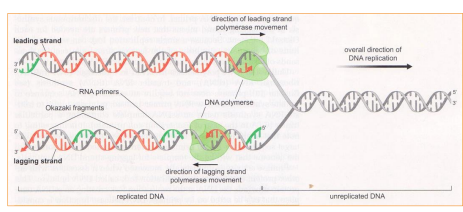




#### 3'-HO 5' **DNA Polymerases are** processive enzymes DNA polymerase binds (slow) "putative" nonprocessive □ Processivity: average processive DNA **DNA** polymerase polymerase William and a second number of nts added Minimum 5 each time the enzyme binds a primer:template junction DNA synthesis (fast) Manual S. □ Catalysis rate: can be up to 1000 nt/second one dNTP many dNTPs added □ Processivity: depending DNA polymerase releases on the specific enzyme, it can range from few nts to >50,000 bases added per binding event

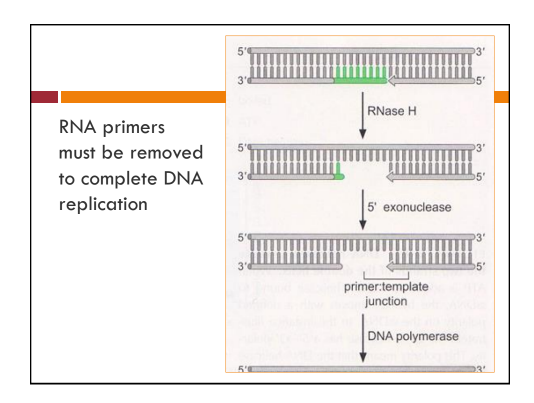


## Replication fork: both strands of DNA are synthesized together at same replication fork



Okazaki fragments length: 1000-2000 bp in bacteria and 100-400 nts in eukaryotes.

RNA primers: 5-10 nts.



# DNA polymerases are specialized for different roles in the cell

Prokaryotic (E. coli)	Number of Subunits	Function
Pol I	111	RNA primer removal, DNA repair
Pol II (Din A)	1	DNA repair
Pol III core	3	Chromosome replication
Pol III holoenzyme	9	Chromosome replication
Pol IV (Din B)	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	DNA repair, translesion synthesis (TLS)
Pol V (UmuC, UmuD' <sub>2</sub> C)	3	TLS

Only Pol I & III have proofreading activity and function in DNA replication

# DNA polymerases are specialized for different roles in the cell

Eukaryotic	Number of Subunits	Function
Pol α	na sagan maa 4	Primer synthesis during DNA replication
Polβ	1	Base excision repair
Polγ	3	Mitochondrial DNA replication and repair
Pol δ	2–3	Lagging-strand DNA synthesis; nucleotide and base excision repair
Pol ε	4	Leading-strand DNA synthesis; nucleotide and base excision repair
Polθ	ucture of a sliding 1784 clamp.	DNA repair of cross-links
Pol ζ	canciated with DN 1 The open	TLS
Pol λ	(1) 1 (1) (1) (1) (1) (1) (1) (1) (1) (1	Meiosis-associated DNA repair
Pol µ	1	Somatic hypermutation
Polκ	1	TLS
Pol η	Slong rive clang 1 out rate to	Relatively accurate TLS past cis-syn cyclobutane dimers
Polı	1	TLS, somatic hypermutation
Rev1	1	TLS

### Class activity??

- □ Both the leading and lagging strands requires primase to initiate DNA synthesis, but the frequency of primase function on the two strands is dramatically different. WHY??
- □ Why the primase adds an RNA primer, not a DNA primer??

### Class discussion

□ Life and biodiversity depend on a happy balance between mutation and its repair!

#### Mutability & DNA Repair:

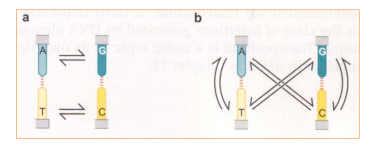
Sources of mutations

(Chapter 9)

- Important sources
  - Inaccuracy in DNA replication
  - Chemical damage to DNA
  - Insertions generated by mobile DNA elements or transposons
- □ Consequences of DNA sequence change
  - Permanent change to DNA (mutation)
  - □ Chemical alteration to DNA and prevent its use as a template
- Cell response to mutations: it must scan the genome to detect errors & mend the lesions and if possible restores original sequence

### Replication errors & their repair

- Mutations include almost every conceivable change in DNA sequence
- □ Switches of one base to another
  - □ Transitions: pyrimidine-to-pyrimidine or purine-to-purine
  - □ Transversions: pyrimidine-to-purine or purine-to-pyrimidine



#### Replication errors & their repair

- □ Insertions & deletions: either small or drastic changes
- $\ \square$  Mutations can arise spontaneously at any given site and ranges from  $10^{-8}$  to  $10^{-11}$  per round of DNA replication
- Some sites are hot spots: simple sequence repeats of dior tri- or tetra-nucleotide sequences (DNA microsatellites)
- Slippage occurs during replication of DNA microsatellites (repetitive DNA sequences), resulting in increase or decrease of repeats >> generates polymorphisms
- □ Examples: the "CA" repeat sites are highly polymorphic in the population

#### Expansion of Triple repeats causes Disease

- □ Huntington Disease
  - Caused by expansion of the "CAG" repeat
  - CAG is the codon for Gln, & its expansion results in extended stretch of Gln in the mutant protein
  - Normal individuals carry 9-35 CAG repeats (average 18-19); affected individuals have >40 repeats; borderline repeats 36-39 repeats have low penetrance
  - The poly-Gln interferes with normal interaction of protein with transcription factor Sp1 & the corresponding Gln-rich patch in the TAFII130 (a subunit of the transcription factor TFIID) > impairs transcription in neurons and brain

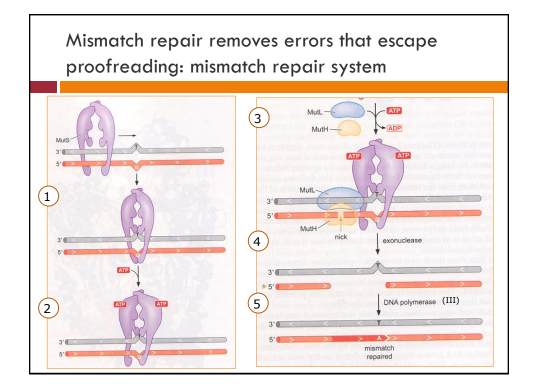
#### Expansion of Triple repeats causes Disease

- □ Other diseases caused by triplet expansion includes
- □ adult muscular (myotonic) dystrophy
  - Caused by expansion of CTG repeat in 3' UTR of a protein kinase gene or DMPK
  - □ Normal range of repeats is from 5 to 30
- □ fragile X syndrome
  - Caused by expansion of CGG repeat in 5' UTR of FMR1 gene)
  - □ Normal range of repeats is up to 60

### 

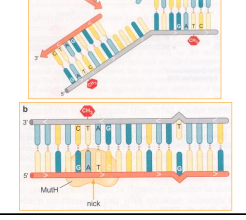
# Mismatch repair removes errors that escape proofreading

- Mismatch repair system: responsible for the fidelity of DNA replication
- □ Functions:
  - It must scan the genome for mismatches, mismatches are transient
  - □ It must correct the mismatch accurately
  - In E. coli mismatches are detected by the MR protein: MutS (has ATPase activity)
  - MutS embraces the mismatch, induces a kink in DNA, recruits MutL. The MutL activates MutH (endonuclease)
  - □ The UvrD Helicase unwinds the DNA starting at incision site



How does the E. coli mismatch repair system recognizes which of the 2 mismatched nts to repair?

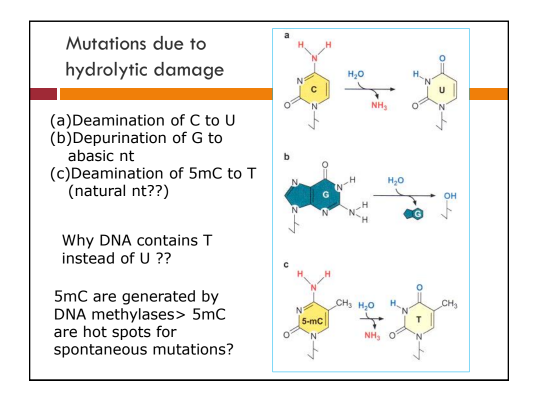
- E. coli Dam methylase, methylates A residues on both strands of the sequence 5'-GATC-3'
- MutH makes incision in unmethylated daughter strand

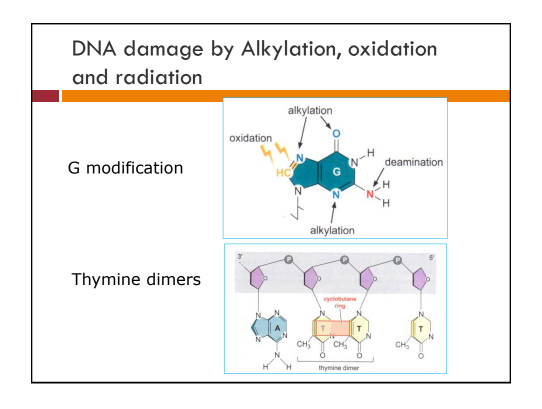


#### DNA damage:

by Spontaneous Hydrolysis, Deamination & depurination

- □ Other Causes of damage:
  - Environmental factors like radiation
  - Chemical mutagens
- □ Examples:
  - Spontaneous deamination of C >> U (U pairs with A rather than G)
  - Spontaneous deamination of A or G >> hypoxanthine (pairs with C) Or xanthine (continues to pair with C but with 2 Hbonds), respectively
  - Depurination by spontaneous hydrolysis of N-glycosyl linkage>> abasic nt



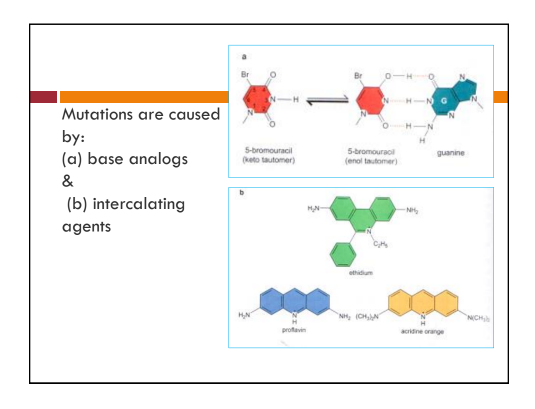


#### DNA damage by Alkylation, oxidation and radiation

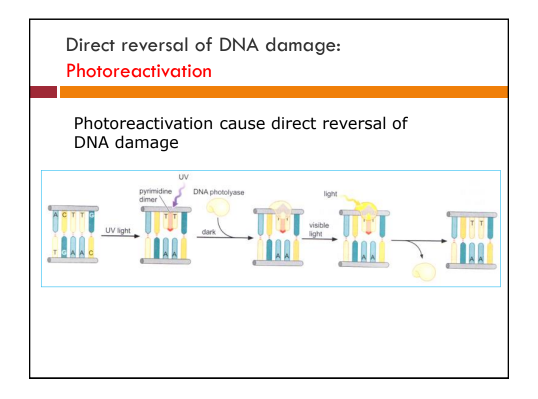
- Alkylation: methyl or ethyl groups are transferred to reactive sites on bases & phosphates in DNA backbone
- $\Box$  Example: Nitrosamines, which methylates G to O<sup>6</sup>-methylguanine which mispairs with T (G:C > A:T)
- Oxidation: via ROS (O2-, H2O2, OH.) generated from ionizing radiation & chemicals
- $\hfill\Box$  Oxidation of G > oxoG which pairs with A (G:C to A:T) & C
- □ G:C to A:T transversion is very common in human cancers

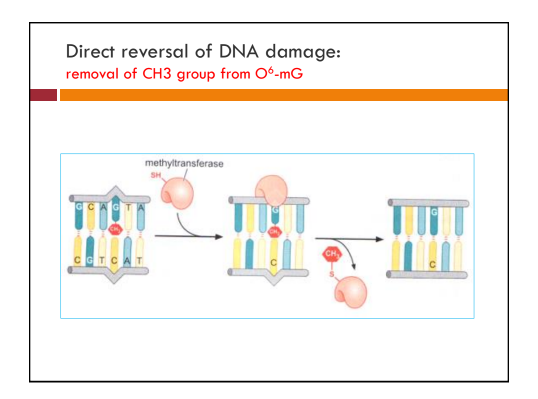
#### DNA damage by Alkylation, oxidation and radiation

- □ Radiation with UV light: forms T-dimers
- □ Gamma radiation & X-rays (ionizing radiation): cause double-strand breaks and can generate free radicals
- Many anticancer drugs like Bleomycin cause breaks in DNA (clastogenic agents)



DNA repair systems				
Туре	Damage	Enzyme		
Mismatch repair	Replication errors	MutS in E. coli, MSH in humans		
Photoreactivation	Pyrimidine dimers	DNA photolyase		
Base excision repair	Damaged bases	DNA glycosylase		
Nucleotide excision repair	Pyrimidine dimers, bulky adducts on bases	UvrA/B/C/D in E. coli XPC/A/D, ERCCI-XPF, XPG in humans		
Double-strand break repair	Double-strand breaks	RecA & RecBCD in E. coli		
Translesion DNA synthesis	Pyrimidine dimers or apurinic site	Y-family DNA polymerases, such as UmuC in E. coli		

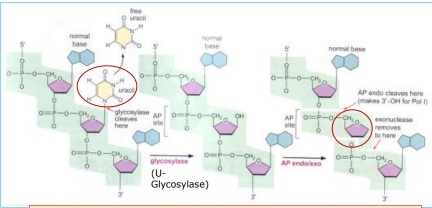




#### Repair of DNA damage:

Base excision repair pathway

Glycosylase: U-glycosylase; AP: apurinic or apyrimidinic



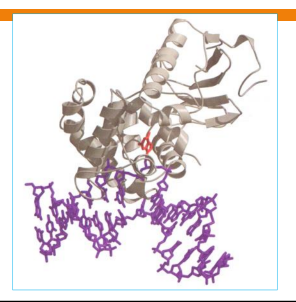
After the damaged nt is removed a repair DNA Pol and DNA ligase restore the intact strand

## Structure of a DNA Glycosylase complex

How the glycosylase recognize the damaged base buried in the helix?

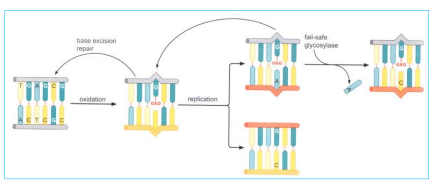
Damaged base is flipped out of the helix (?)

Cells have 8 different glycosylases, & DNA glycosyalses are lesion-specific: e.g. U- and oxoGglycosylases



### oxoG:A repair and the Fail-safe Glycosylase

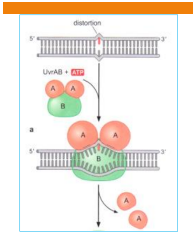
oxoG Glycosylase removes the undamaged incorrect A opposite oxoG



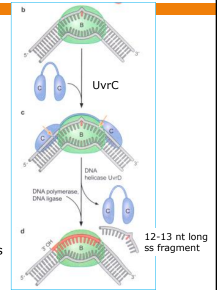
Another Fail-safe glycosyalse removes T opposite to G, that results from deamination of 5mC to T  $\,$ 

#### Repair of DNA damage:

Nucleotide excision repair pathway



Nucleotide excision repair in eukaryotes is similar but more complex, involving 25 or more polypeptides



#### Clinical correlations:

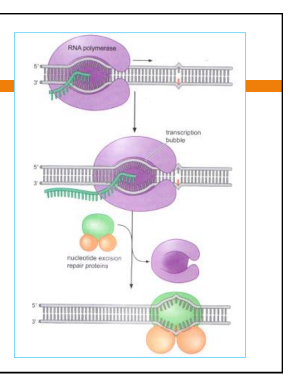
#### UVR proteins & Xeroderma pigmentosum

- Mutants of the uvr genes are sensitive to UV light and lack the capacity to remove T-dimers
- □ In human seven XP genes (XPA-G) are responsible for the Nucleotide excision repair system
- □ Defects in the XP genes give rise to Xeroderma pigmentosum (sun-light induced skin cancer)

## Transcription-coupled repair >

RNA polymerase serves as another damagesensing protein. It recruits the Nucleotide excision repair proteins and backs up or dissociates from DNA

In eukaryotes, the TFIIH is crucial for this process with its subunits XPA & XPD



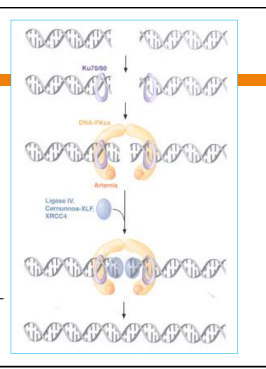
#### Recombination repairs DNA breaks

- □ How do cells repair double-strand breaks (DSB) in DNA in which both strands of the duplex are broken?
- >>By the DSB repair pathway> retrieves sequence information from undamaged sister chromosome
- Alternatively and under some conditions, SBs are repaired by direct joining of broken ends> nonhomologous end joining (NHEJ)
- □ NHEJ protects and process broken ends then join them
- □ NHEJ is mutagenic

## Mammalian pathway for NHEJ

In mammalian cells, 7 components of NHEJ pathway are known: Ku70, Ku80, DNA-PKs, Artemis, Ligase IV, Cernunnos-XLF & XRCC4.

Artemis has a 5' to 3' exonuclease and endonuclease activities that are activated by phosphorylation by DNA-PKcs.



## Translesion DNA synthesis:

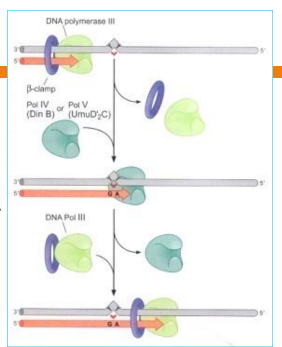
It enables Replication to proceed across DNA damage

TLS is highly error-prone!!

In E.coli, TLS is carried out by UmuC & UmuD'>> a member of Y family of DNA Polymerases

Nucleotide synthesis is not guided by base pairing but may not be random!!

In E. coli Translesion Polymerase is not present under normal conditions



### Translesion DNA synthesis:

- In TLS Nucleotide synthesis is not guided by base pairing but may not be random!! For example, in human a member of Y family of translesion DNA Polymerases correctly inserts two A residues opposite a thymine dimer!
- In E. coli Translesion Polymerase is not present under normal conditions. The genes encoding translesion polymerases are expressed as part of a pathway known as the SOS response