



THE MOLECULAR BASIS OF INHERITANCE

Chapter 13

What you must know

- The structure of DNA.
- The knowledge about DNA gained from the work of Griffith; Avery, MacLeod, and McCarty; Hershey and Chase; Wilkins and Franklin; and Watson and Crick.
- That replication is semiconservative and occurs 5' to 3'.
- The roles of DNA polymerase, ligase, helicase, and topoisomerase in replication.
- The general differences between bacterial chromosomes and eukaryotic chromosomes.
- How DNA is packaged can affect gene expression.

Problem:

Is the genetic material of organisms made of DNA or proteins?

Frederick Griffith (1928)

Experiment

Living **S cells**
(control)

infected

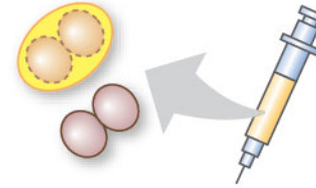
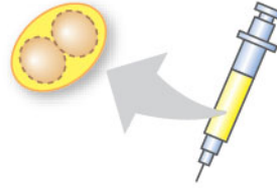
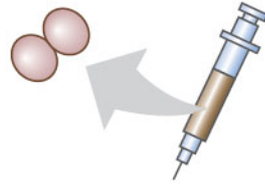
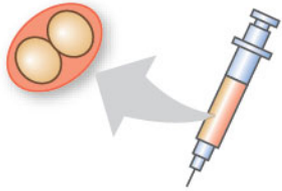
Living R cells
(control)

Denature

Heat-killed S cells
(control)

Denatured infected

Mixture of heat-killed S cells and living R cells



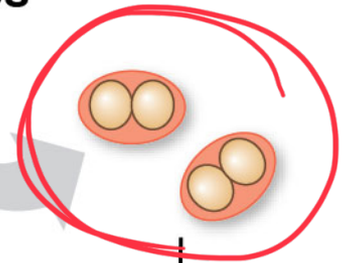
Results

Mouse dies

Mouse healthy

Mouse healthy

Mouse dies



Living S cells

Frederick Griffith (1928)

Conclusion: living R bacteria **transformed** into deadly S bacteria by unknown, heritable substance

Avery, McCarty, MacLeod (1944)

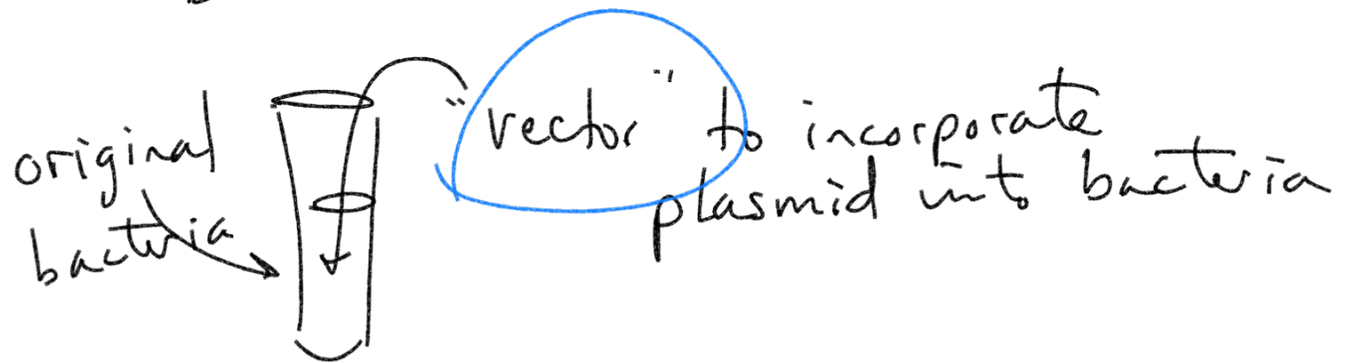
- ▣ Tested DNA, RNA, & proteins in heat-killed pathogenic bacteria
- ▣ Discovered that the transforming agent was **DNA**

T-6B General Biology Week 27 4/11

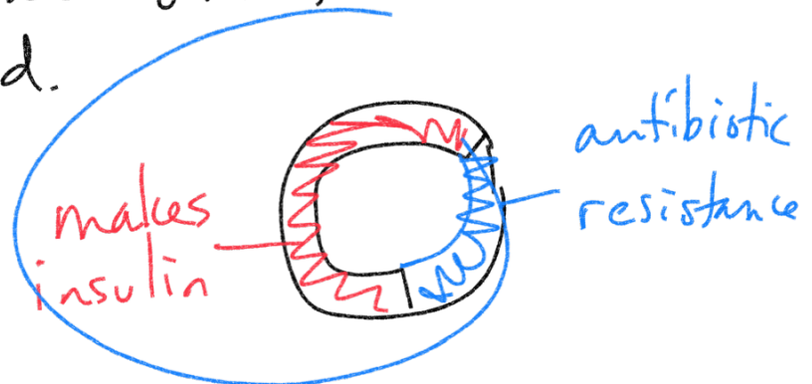
Plasmid → polymer of DNA that we want to introduce to a bacteria

Plasmid for produce insulin

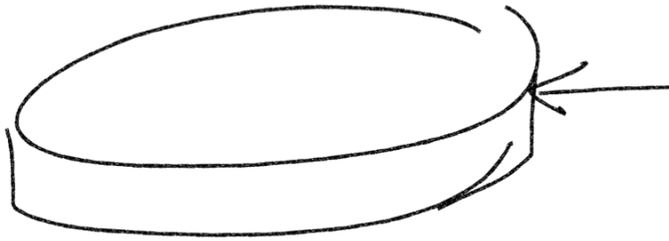
Original Bacteria → no antibiotic resistance



Also — incorporate antibiotic resistance on the insulin plasmid.



original bacteria



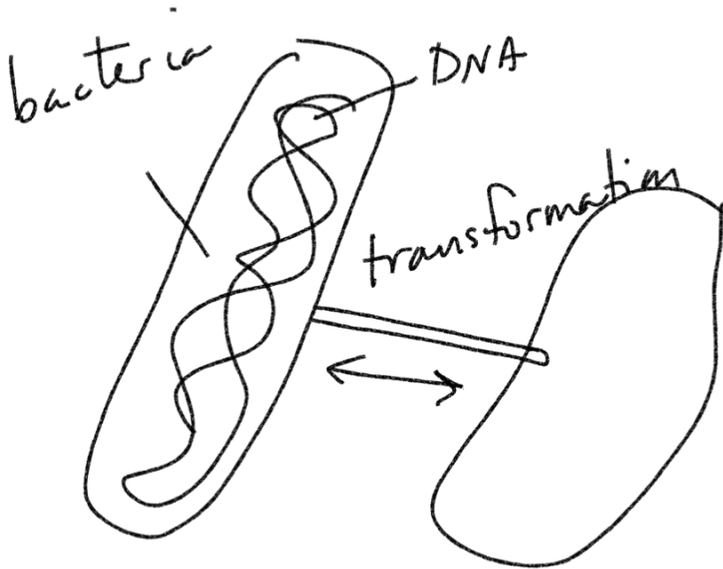
nutrients and antibiotics

original bacteria → Die

original bacteria — Live
w/ plasmid

resistance

make insulin



Hershey and Chase (1952)

- Bacteriophages: virus that infects bacteria; composed of DNA and protein

sulfur

Protein = radiolabel S

DNA = radiolabel P

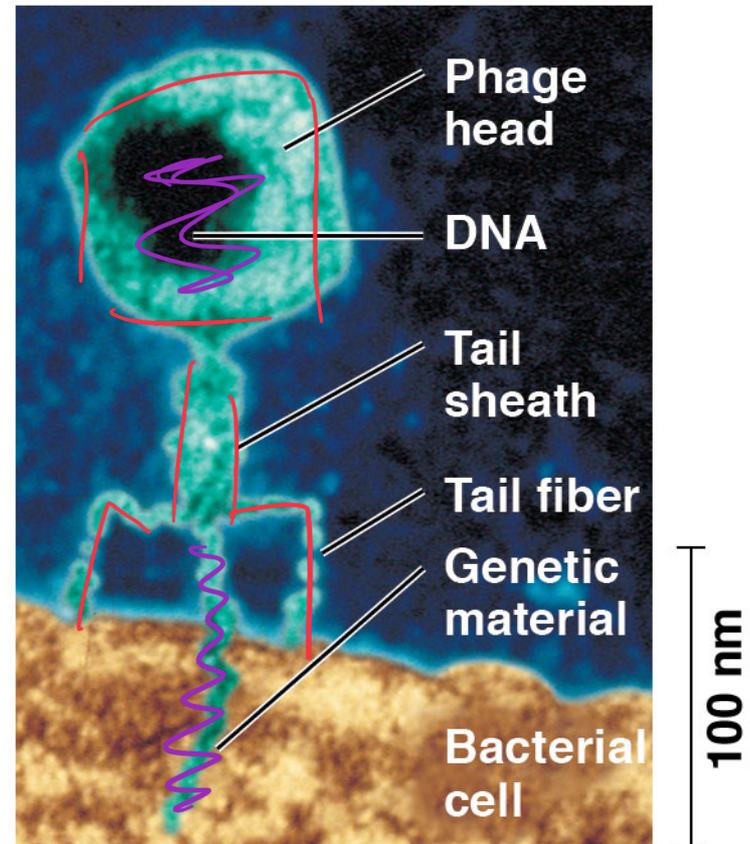
phosphate

"vector"

nucleic acid

CHONPS

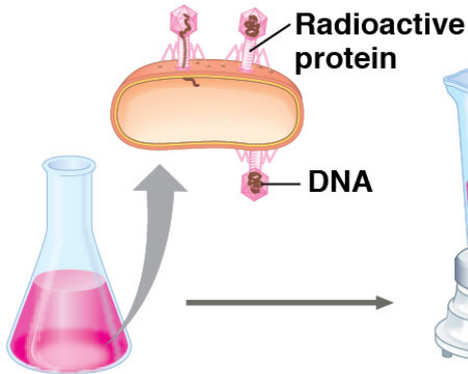
amino acid



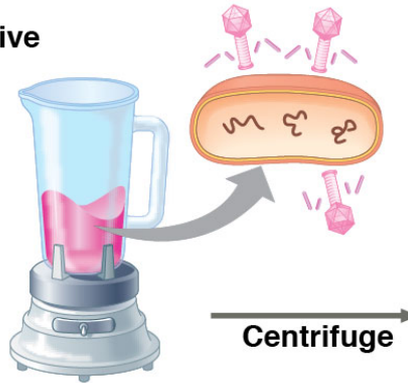
Experiment

Batch 1: Radioactive sulfur (^{35}S) in phage protein

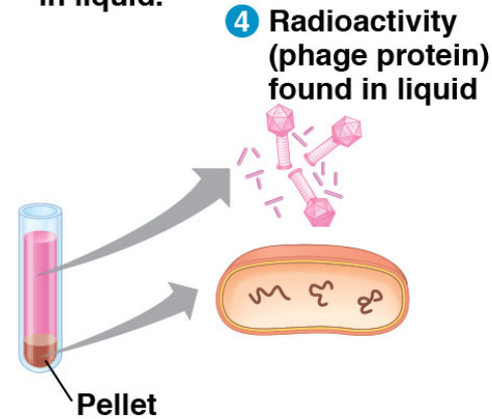
1 Labeled phages infect cells.



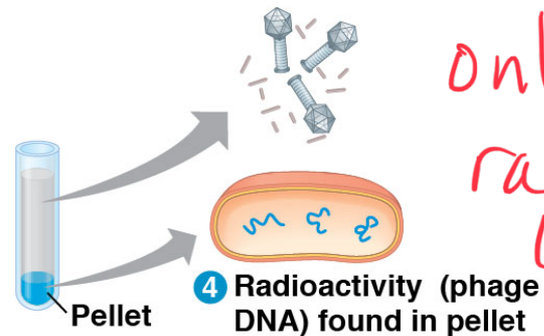
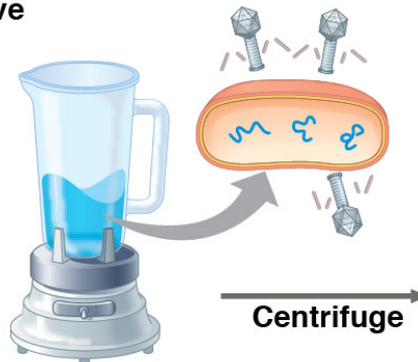
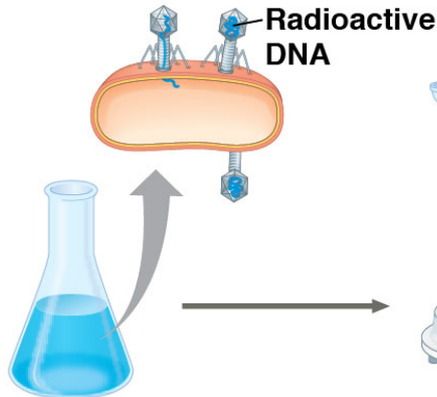
2 Agitation frees outside phage parts from cells.



3 Centrifuged cells form a pellet. Free phages and phage parts remain in liquid.



Batch 2: Radioactive phosphorus (^{32}P) in phage DNA



only find radioactively labeled **(P)**

Conclusion: DNA entered infected bacteria → DNA must be the genetic material!

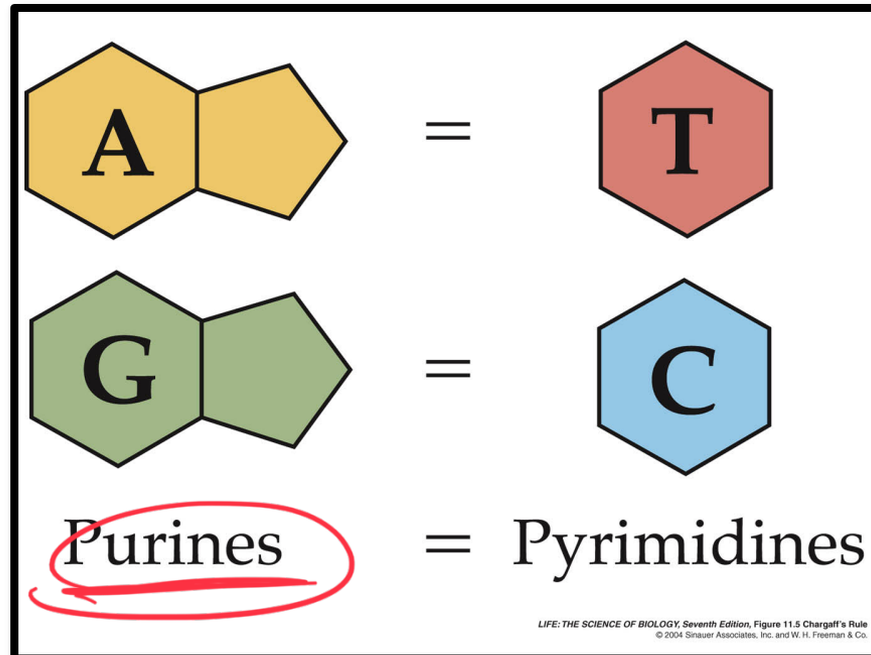
Problem:

What is the structure of DNA?

Edwin Chargaff (1947)

Chargaff's Rules:

- DNA composition varies between species
- Ratios: %A = %T and %G = %C



Pure
As
Gold

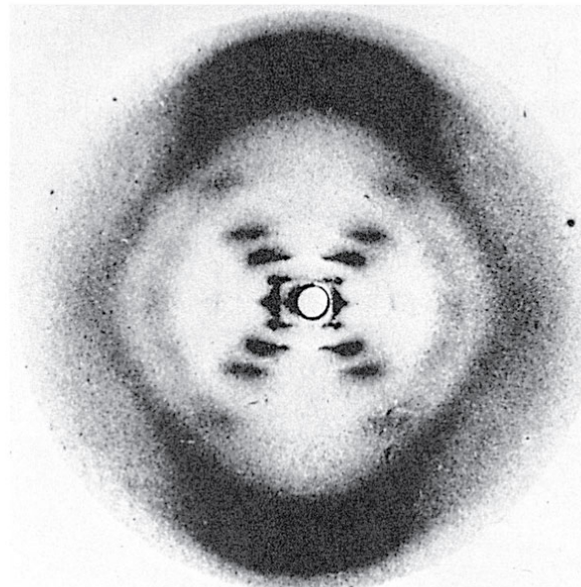
A-T
Pyrimidines
CUT the Py
G-C

Rosalind Franklin (1950's)

- Worked with Maurice Wilkins
- X-ray crystallography = images of DNA
- Provided measurements on chemistry of DNA



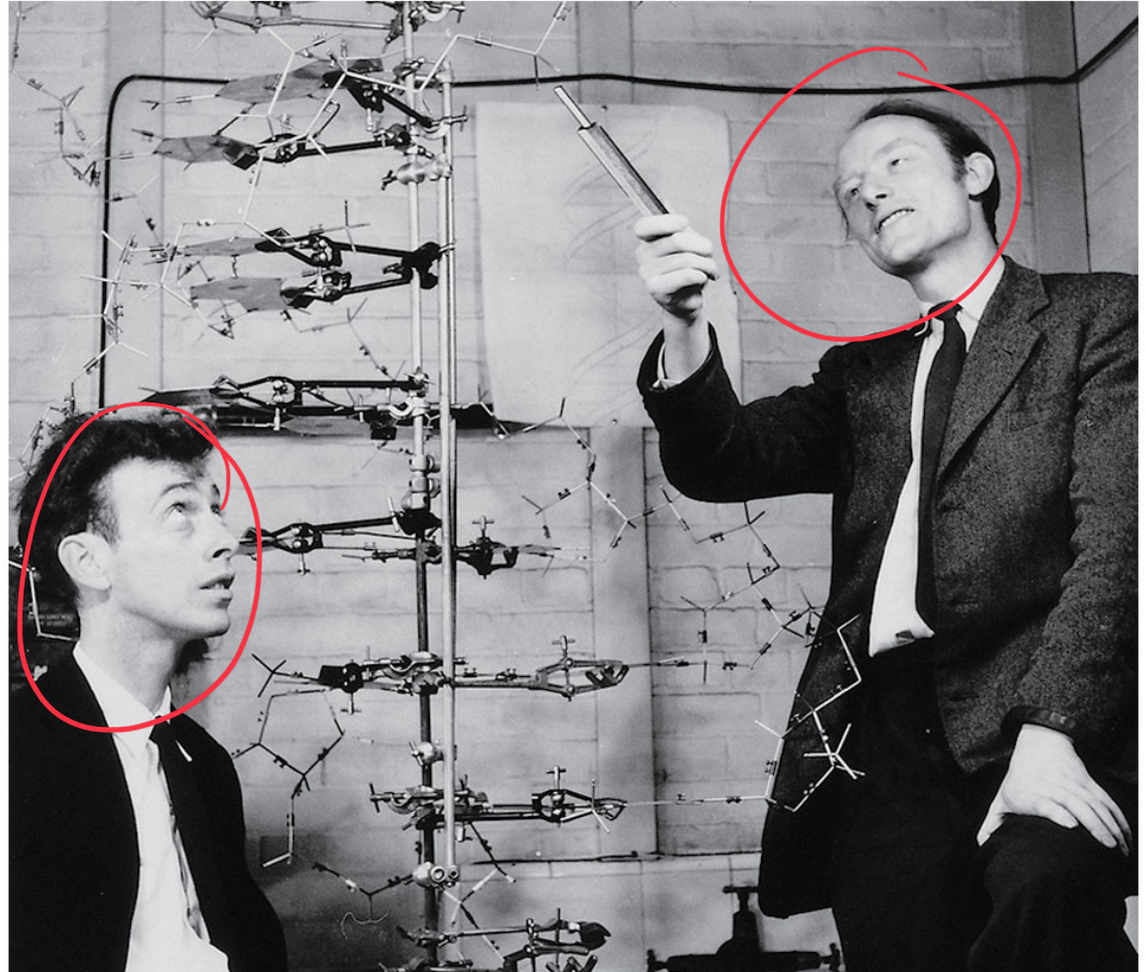
(a) Rosalind Franklin



(b) Franklin's X-ray diffraction photograph of DNA

James Watson & Francis Crick (1953)

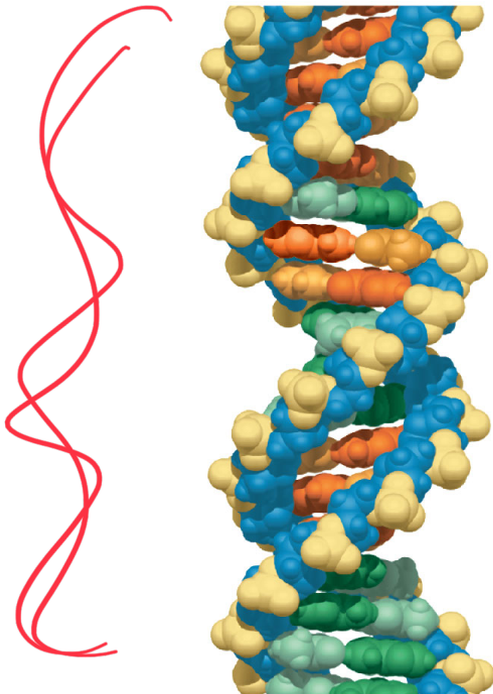
- Discovered the double helix by building models to conform to Franklin's X-ray data and Chargaff's Rules.



DNA = Double Helix

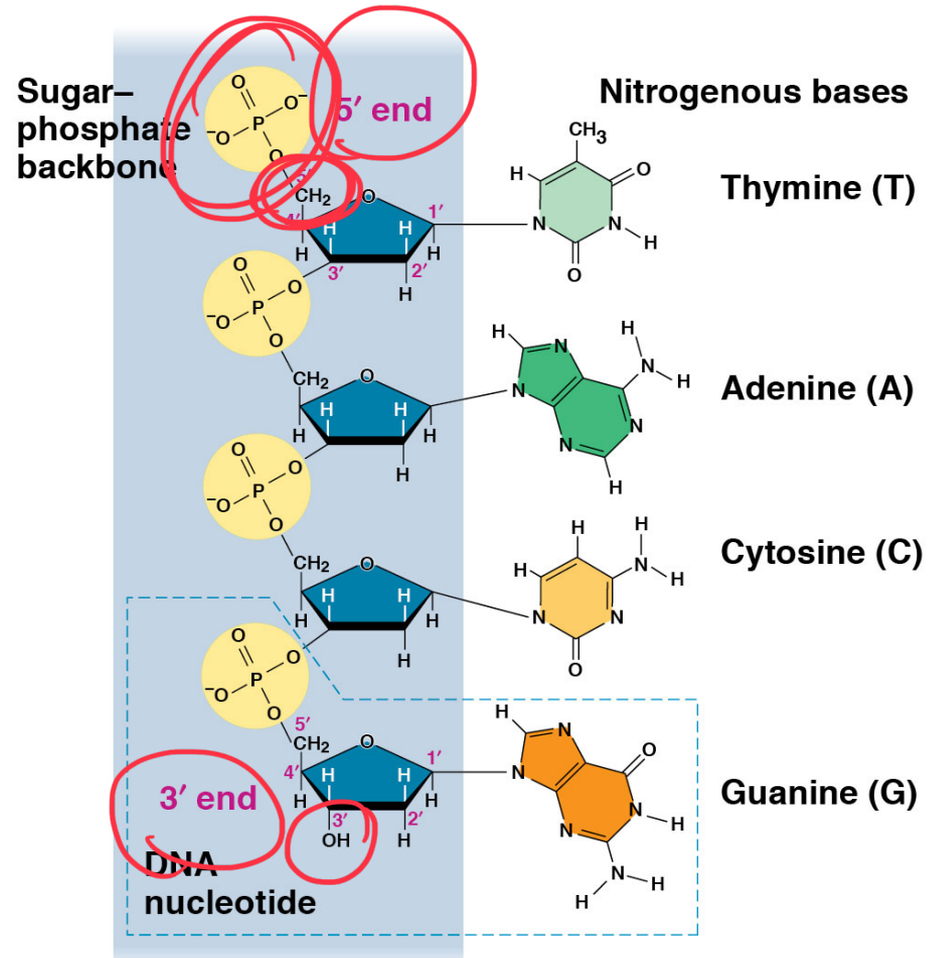
“Backbone” = sugar + phosphate

“Rungs” = nitrogenous bases



(c) Space-filling model

© 2016 Pearson Education, Inc.



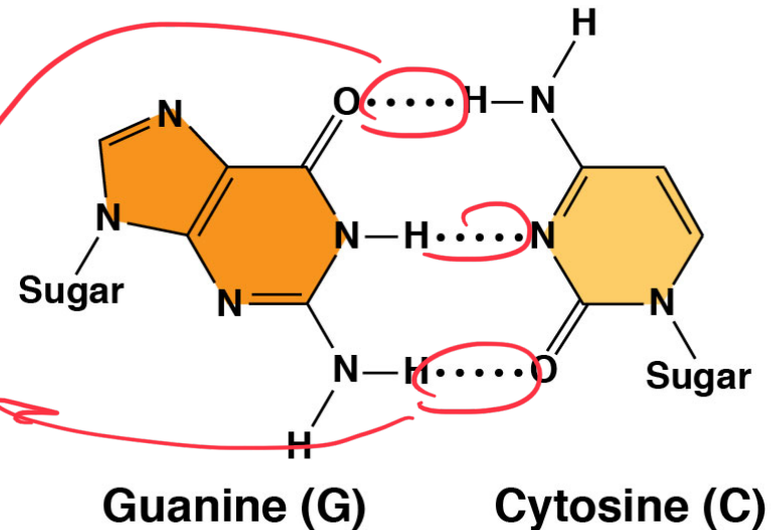
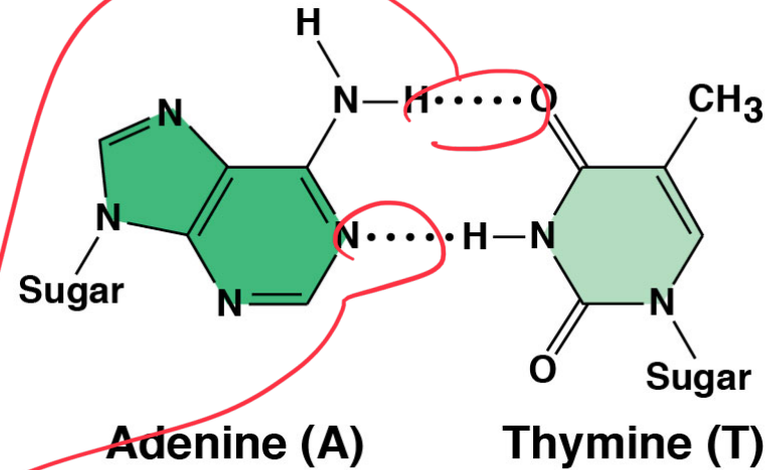
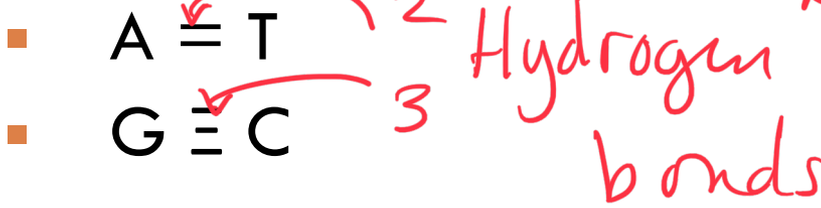
© 2016 Pearson Education, Inc.

Nitrogenous Bases

- Adenine (A)
 - Guanine (G)
 - Thymine (T)
 - Cytosine (C)
- purine
- pyrimidine

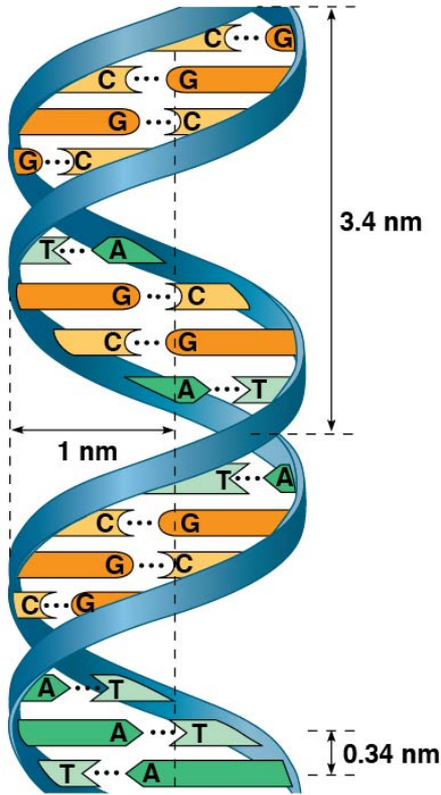
Pairing:

- Purine + Pyrimidine



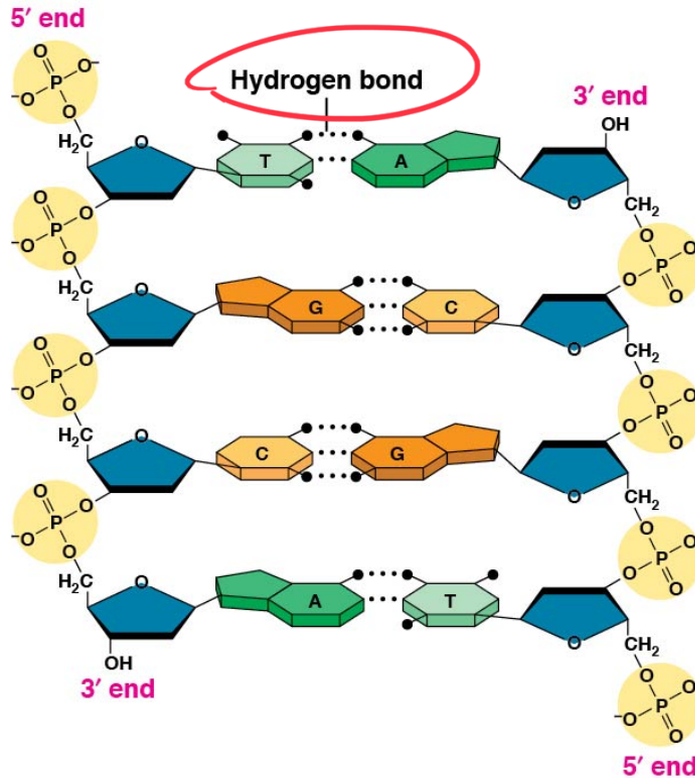
Hydrogen Bonds

Hydrogen bonds bond bases together A=T

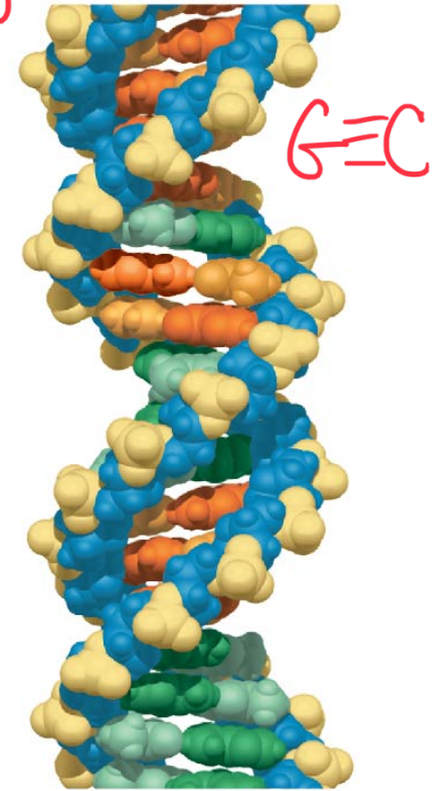


(a) Key features of DNA structure

© 2016 Pearson Education, Inc.



(b) Partial chemical structure

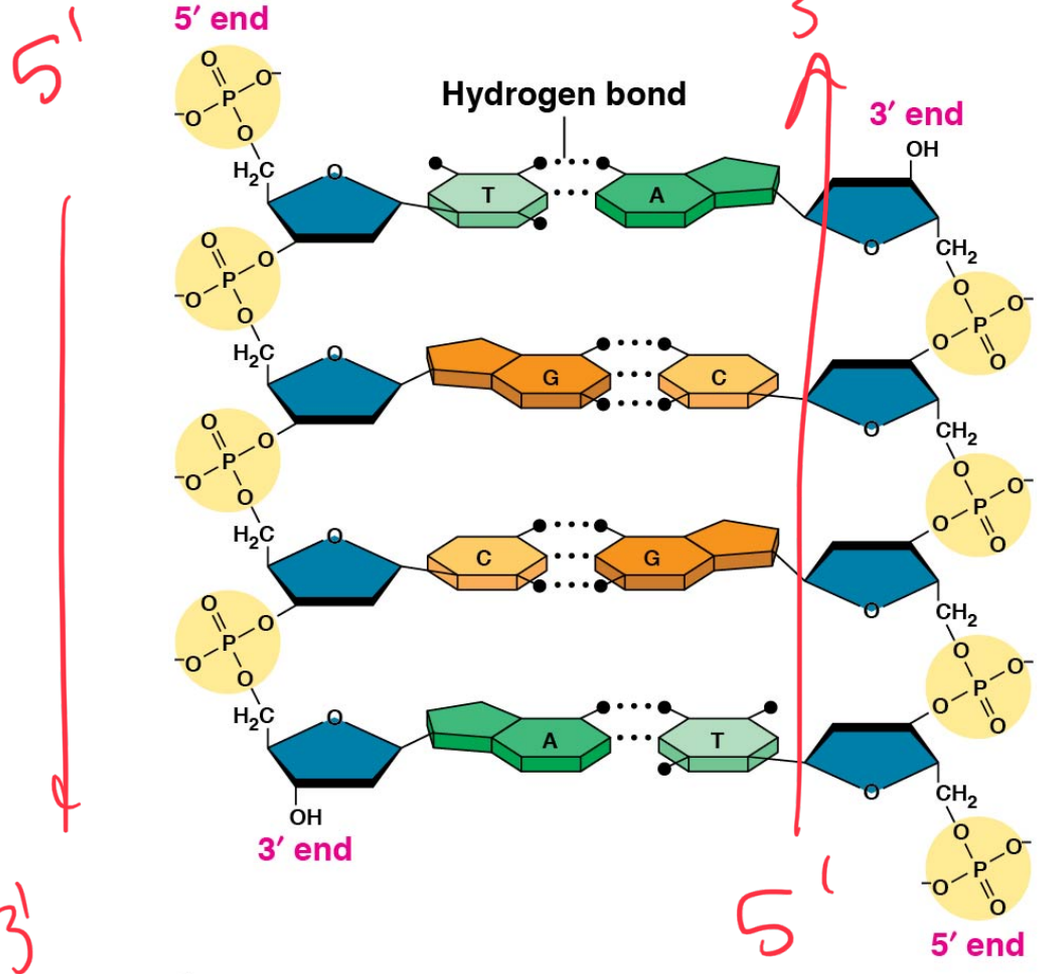


(c) Space-filling model

Hydrogen bonds between base pairs of the two strands hold the molecule together like a zipper.

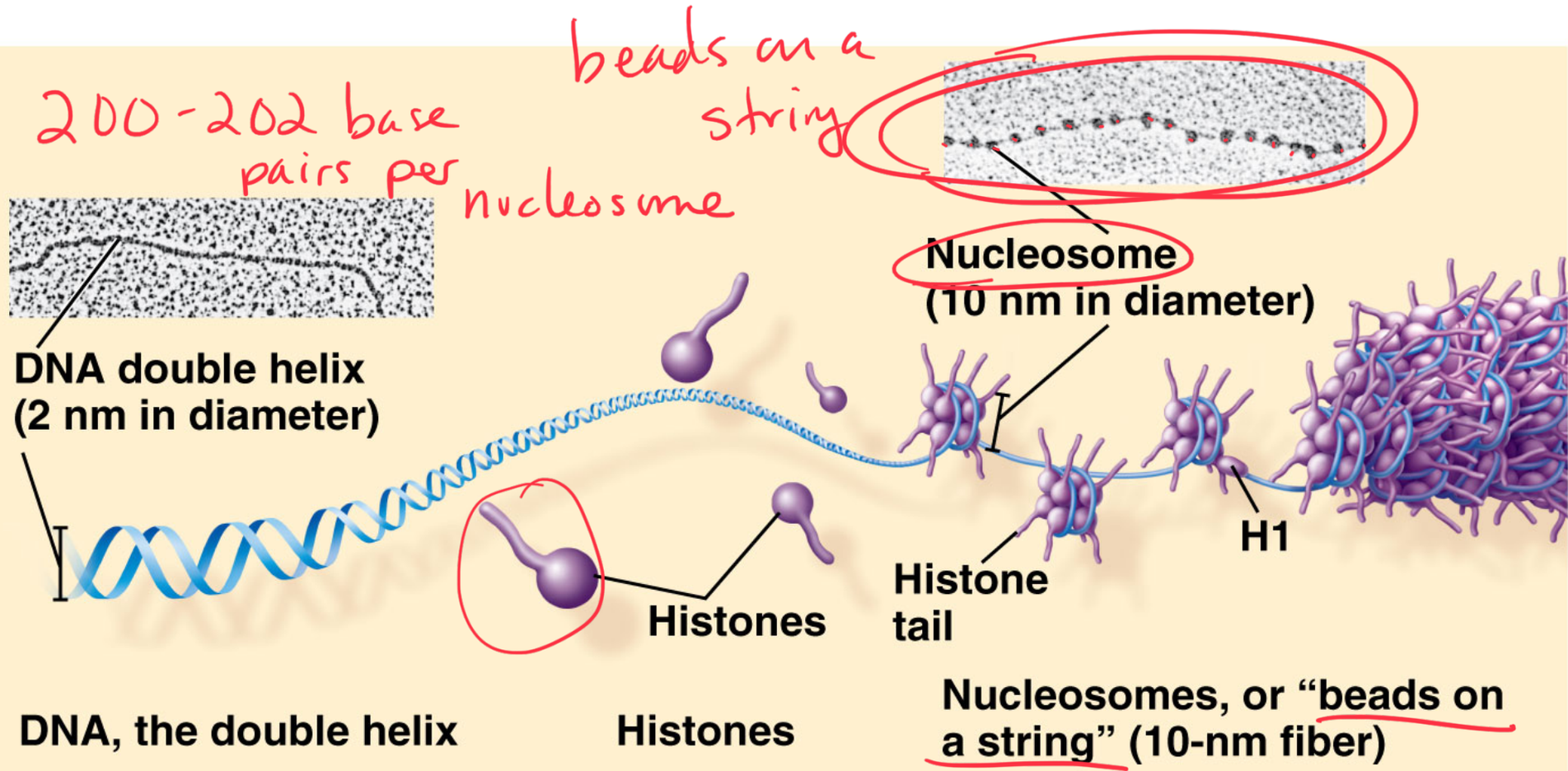
DNA strands are **Antiparallel**

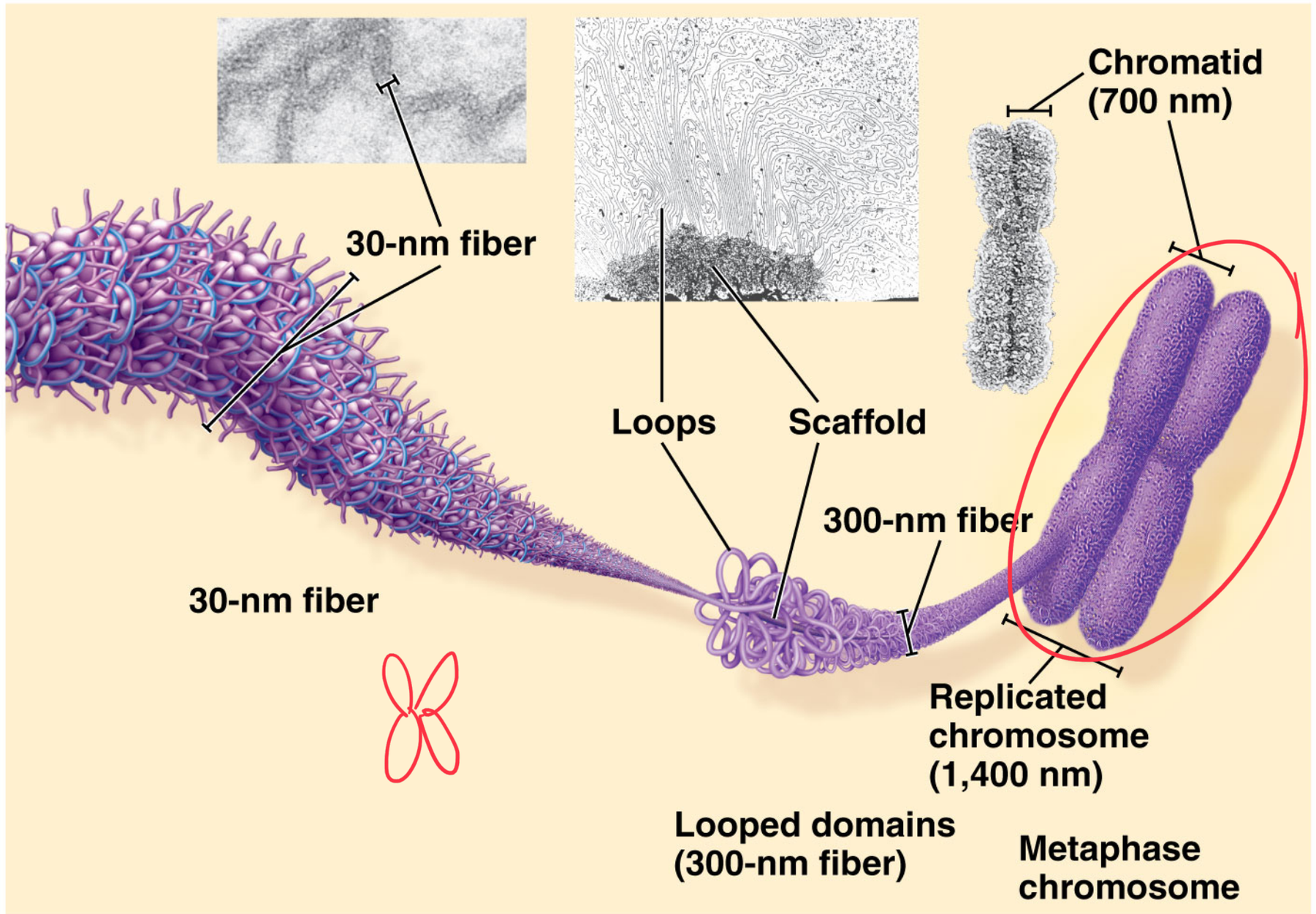
One strand ($5' \rightarrow 3'$),
other strand runs in
opposite, upside-down
direction ($3' \rightarrow 5'$)



(b) Partial chemical structure

How DNA is packaged *through nucleosome of histones*





DNA Comparison

Prokaryotic DNA

- Double-stranded
- Circular *plasmids*
- One chromosome
- In cytoplasm
- Supercoiled DNA (nucleoid)
- No histones

Eukaryotic DNA

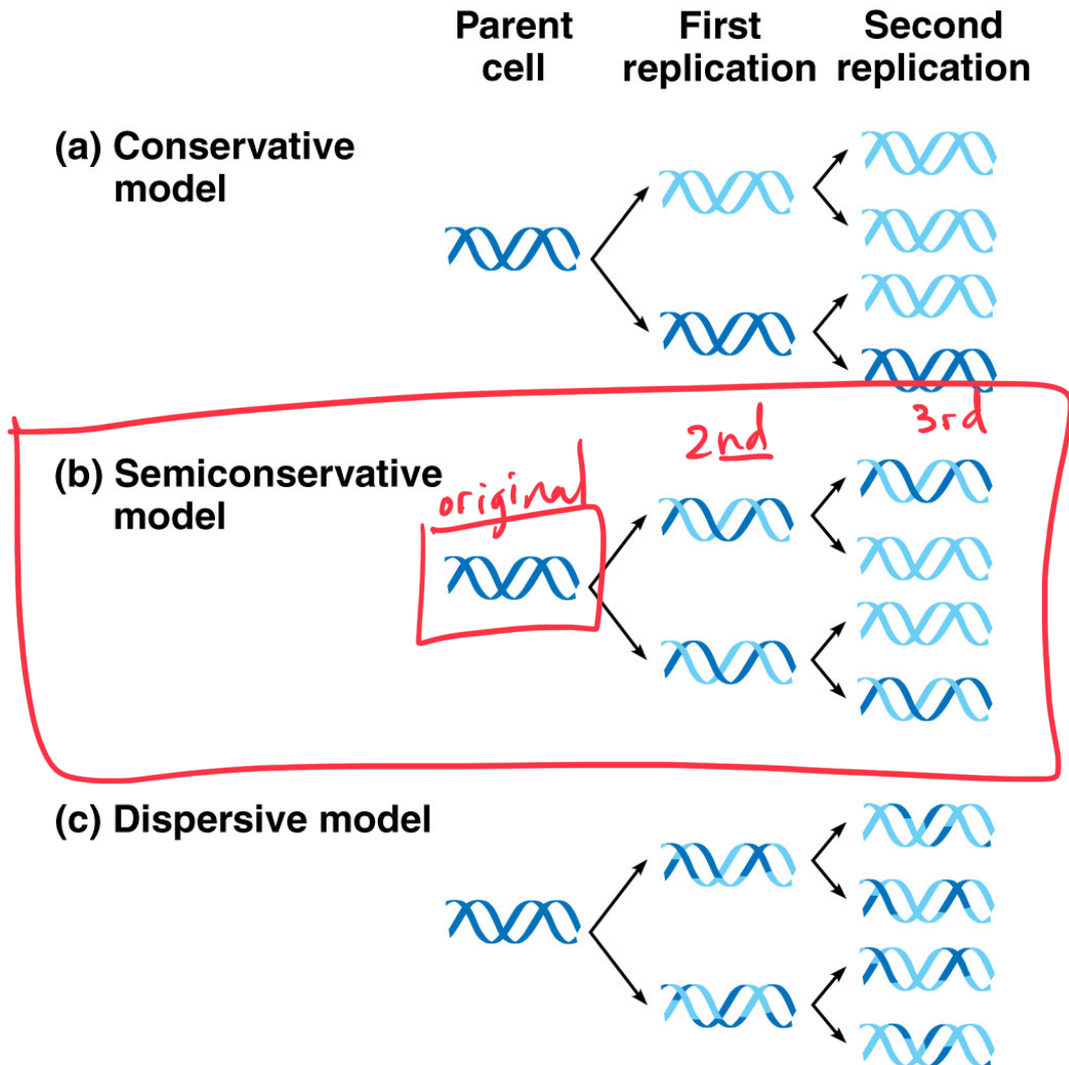
- Double-stranded
- Linear
- Usually 1+ chromosomes
- In nucleus
- Chromatin = DNA wrapped around histones (proteins)

Problem:

How does DNA replicate?

Replication: Making DNA from existing DNA

3 alternative models of DNA replication



Meselson & Stahl

Experiment

1 Bacteria cultured in medium with ^{15}N (heavy isotope)



2 Bacteria transferred to medium with ^{14}N (lighter isotope)

Results

3 DNA sample centrifuged after first replication



4 DNA sample centrifuged after second replication



Less dense
More dense

Meselson & Stahl

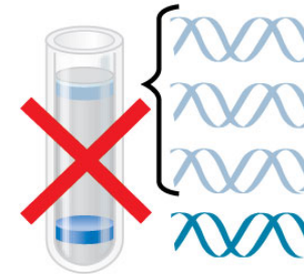
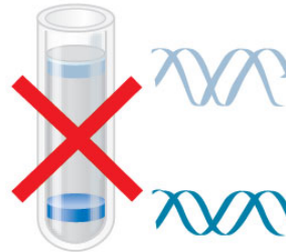
Conclusion

Predictions:

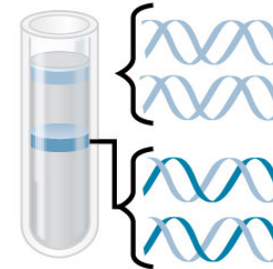
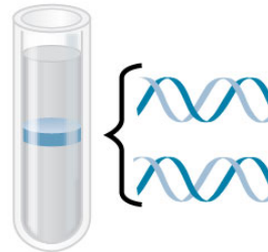
First replication

Second replication

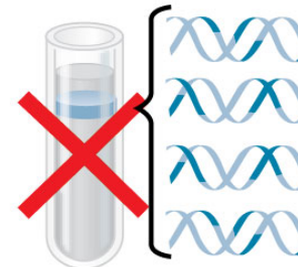
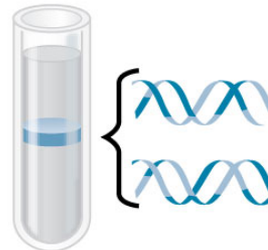
Conservative model



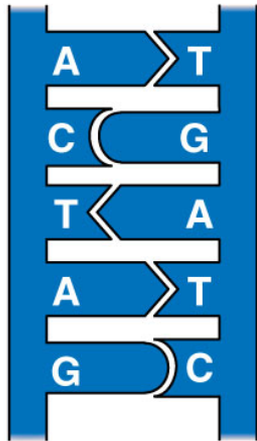
Semiconservative model



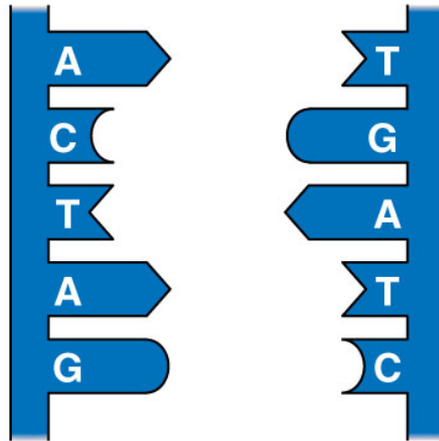
Dispersive model



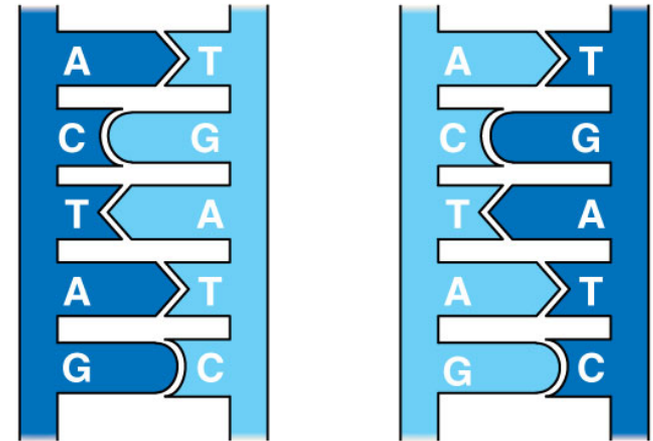
Replication is **semiconservative**



(a) Parent molecule



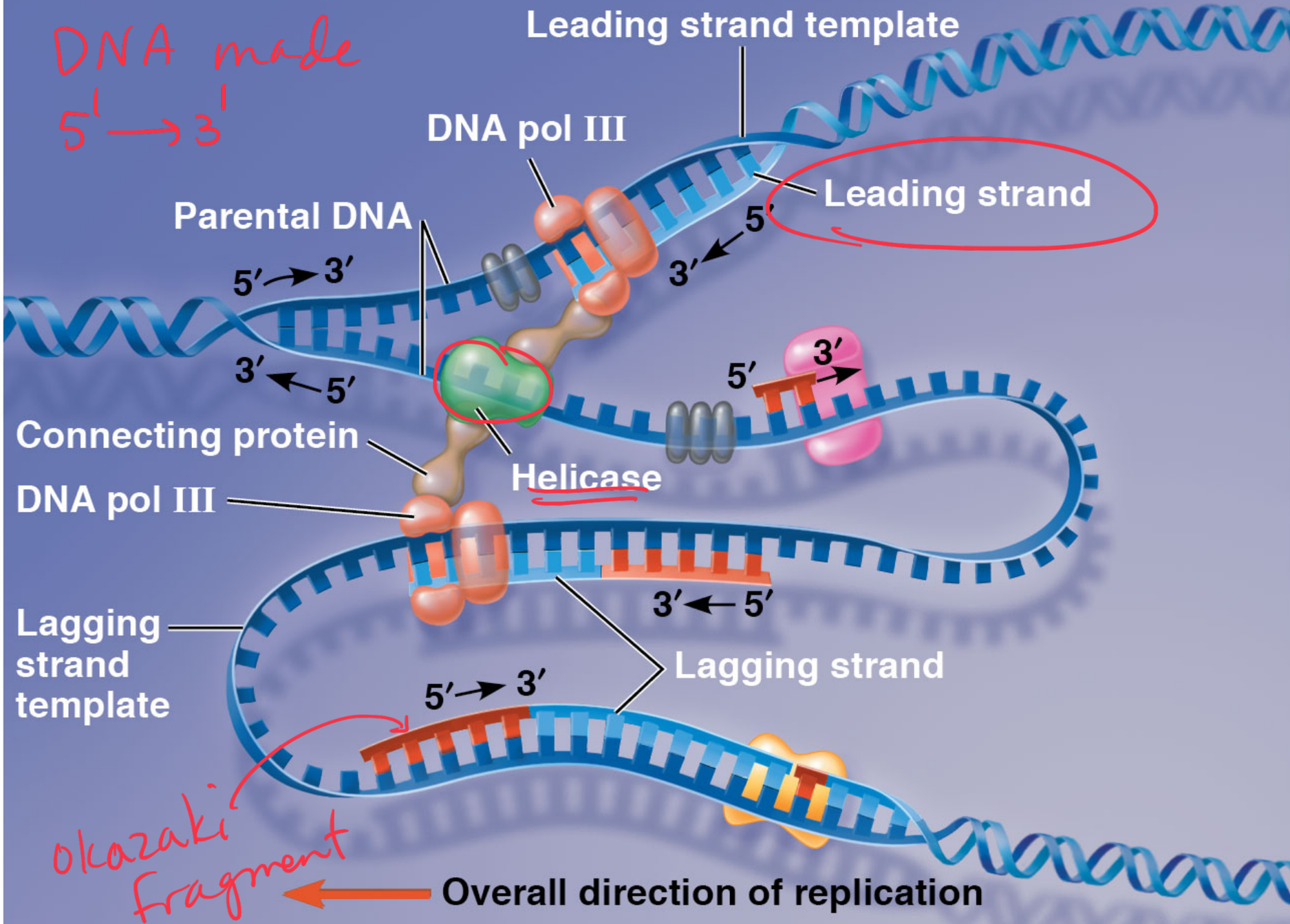
(b) Separation of strands



(c) "Daughter" DNA molecules, each consisting of one parental strand and one new strand

DNA Replication Video

<http://www.youtube.com/watch?v=4jtmOZalvS0&feature=related>

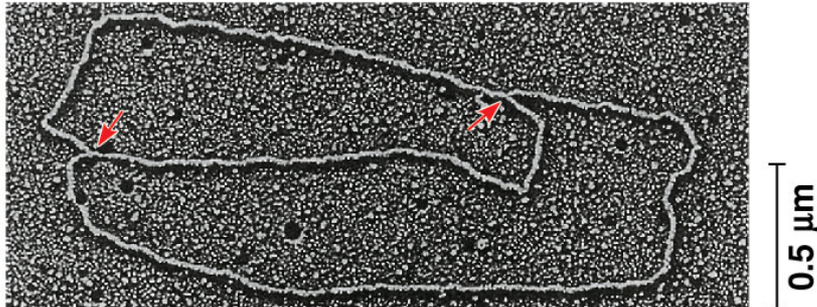
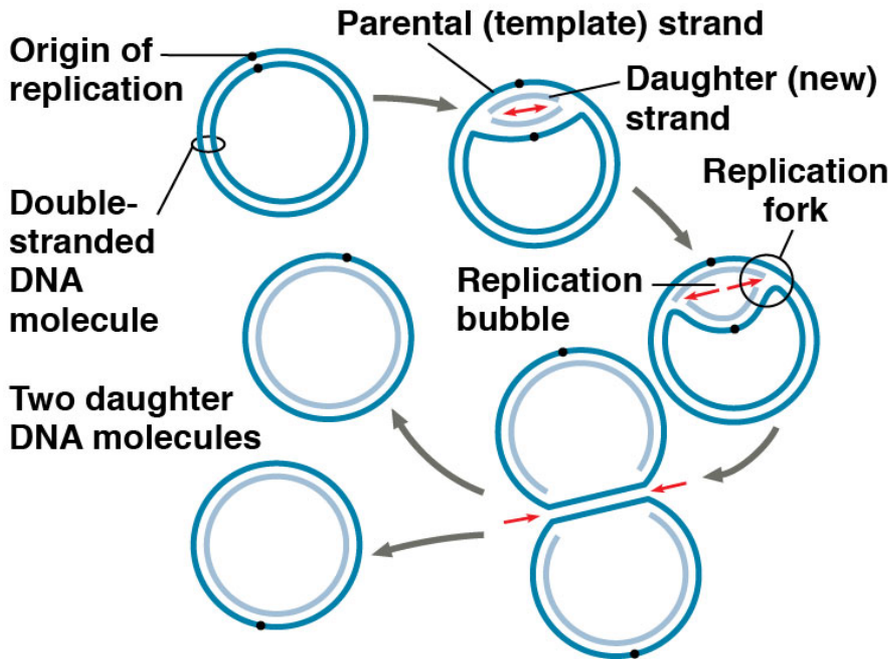


Major Steps of Replication:

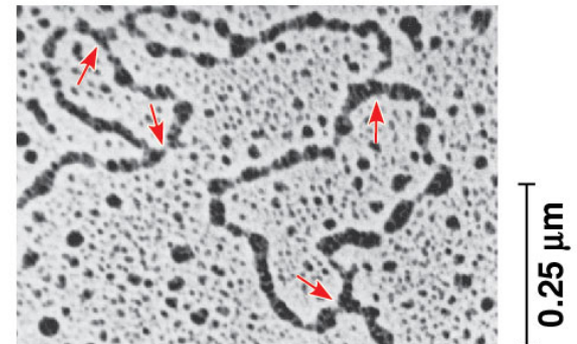
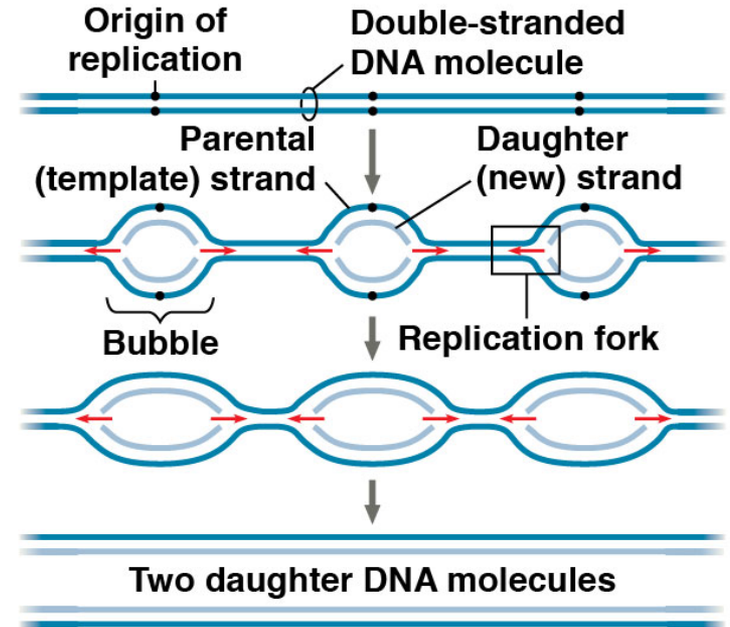
1. **Helicase:** unwinds DNA at *origins of replication*
2. Initiation proteins separate 2 strands → forms *replication bubble*
3. **Topoisomerase:** relieves overwinding strain ahead of replication forks by breaking, swiveling, rejoining DNA strands
4. **Primase:** puts down RNA primer to start replication
5. **DNA polymerase III:** adds complimentary bases to *leading strand*
(new DNA is made 5' → 3')
DNA is made by adding on to primer
6. *Lagging strand* grows in 3' → 5' direction by the addition of *Okazaki fragments*
7. **DNA polymerase I:** replaces RNA primers with DNA
8. **DNA ligase:** seals fragments together
Okazaki fragment *DNA polymerase II troubleshooter*

1. **Helicase** unwinds DNA at *origins of replication* and creates *replication forks*

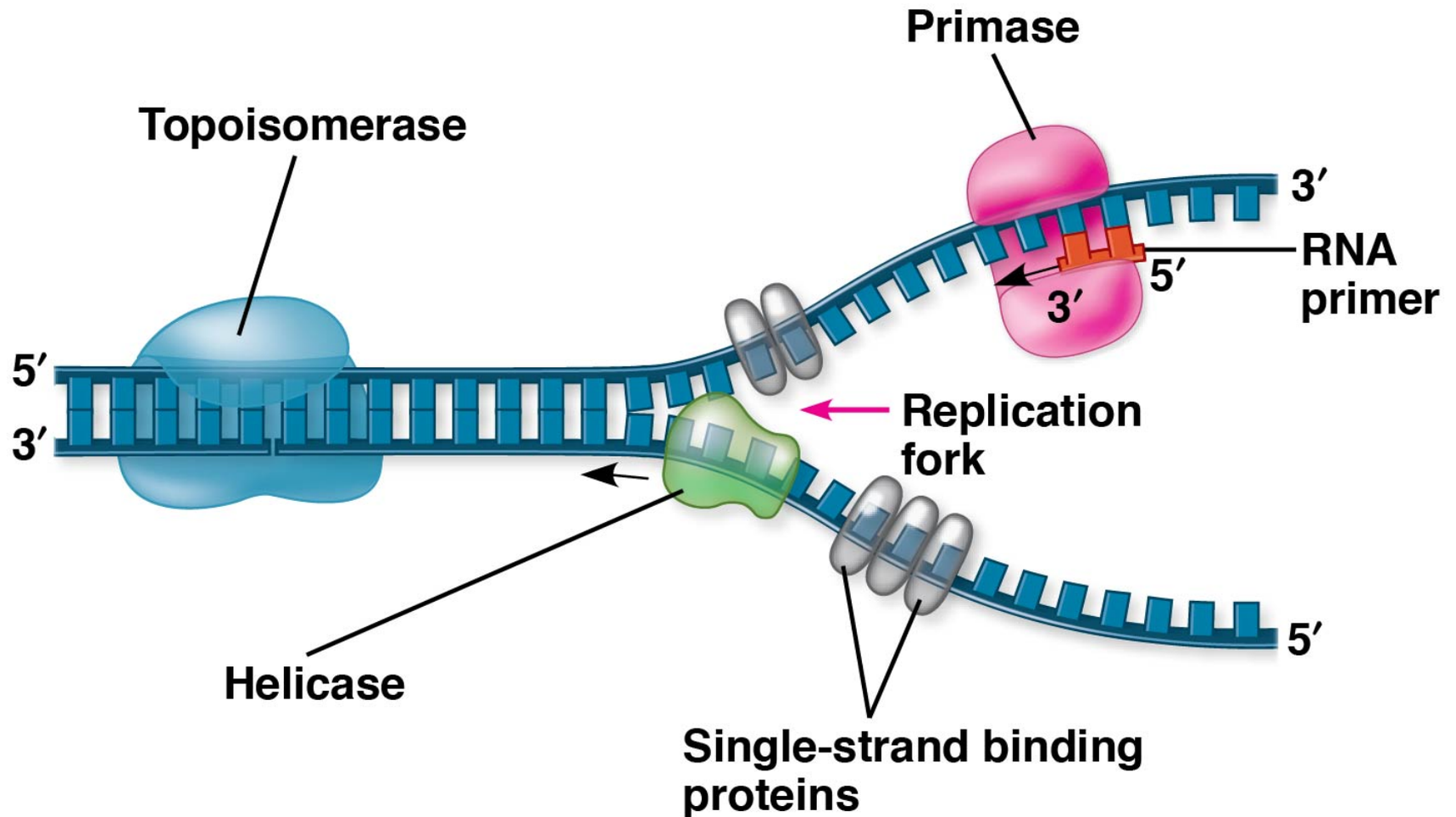
(a) Origin of replication in an *E. coli* cell



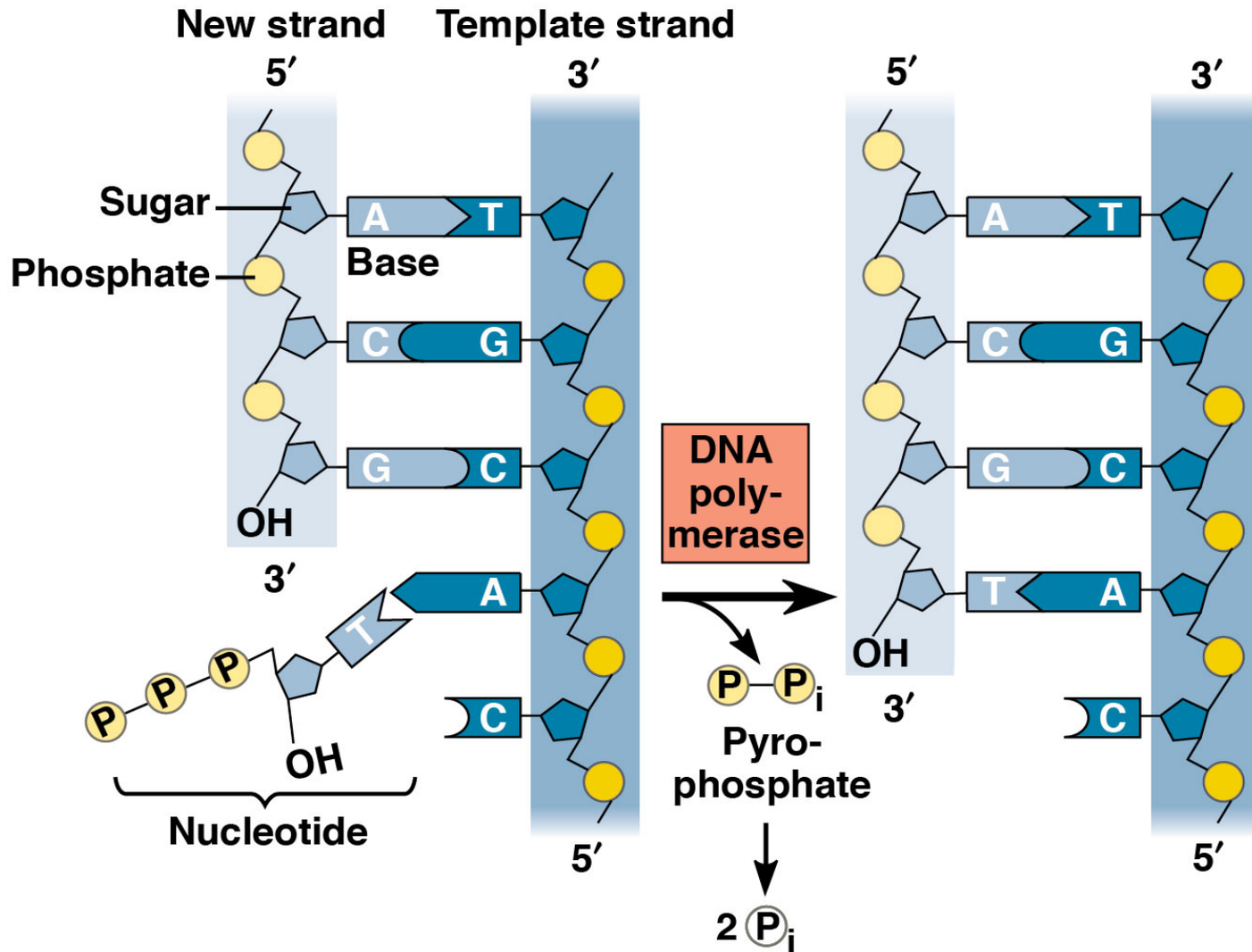
(b) Origins of replication in a eukaryotic cell



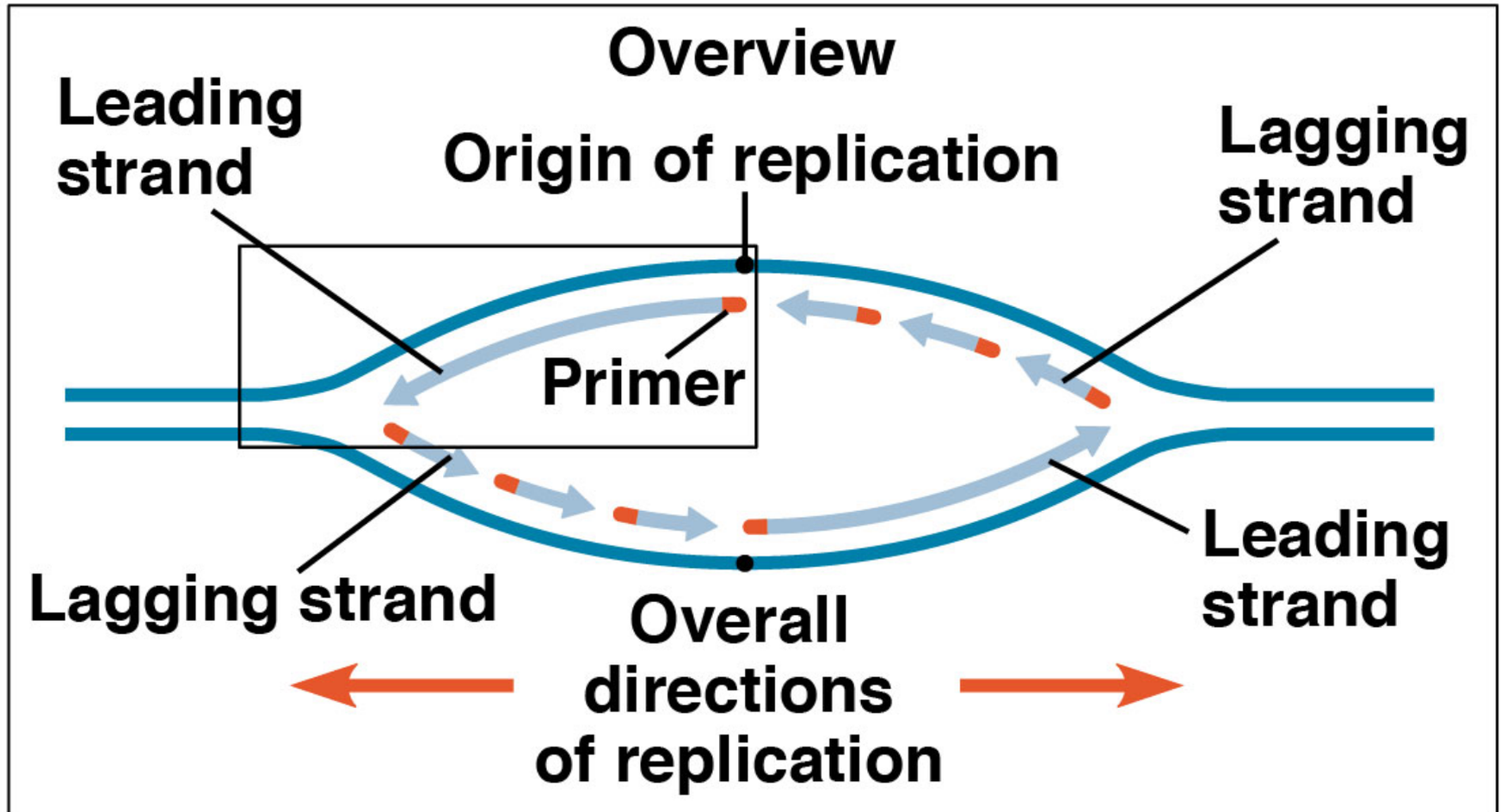
4. Primase adds *RNA primer*



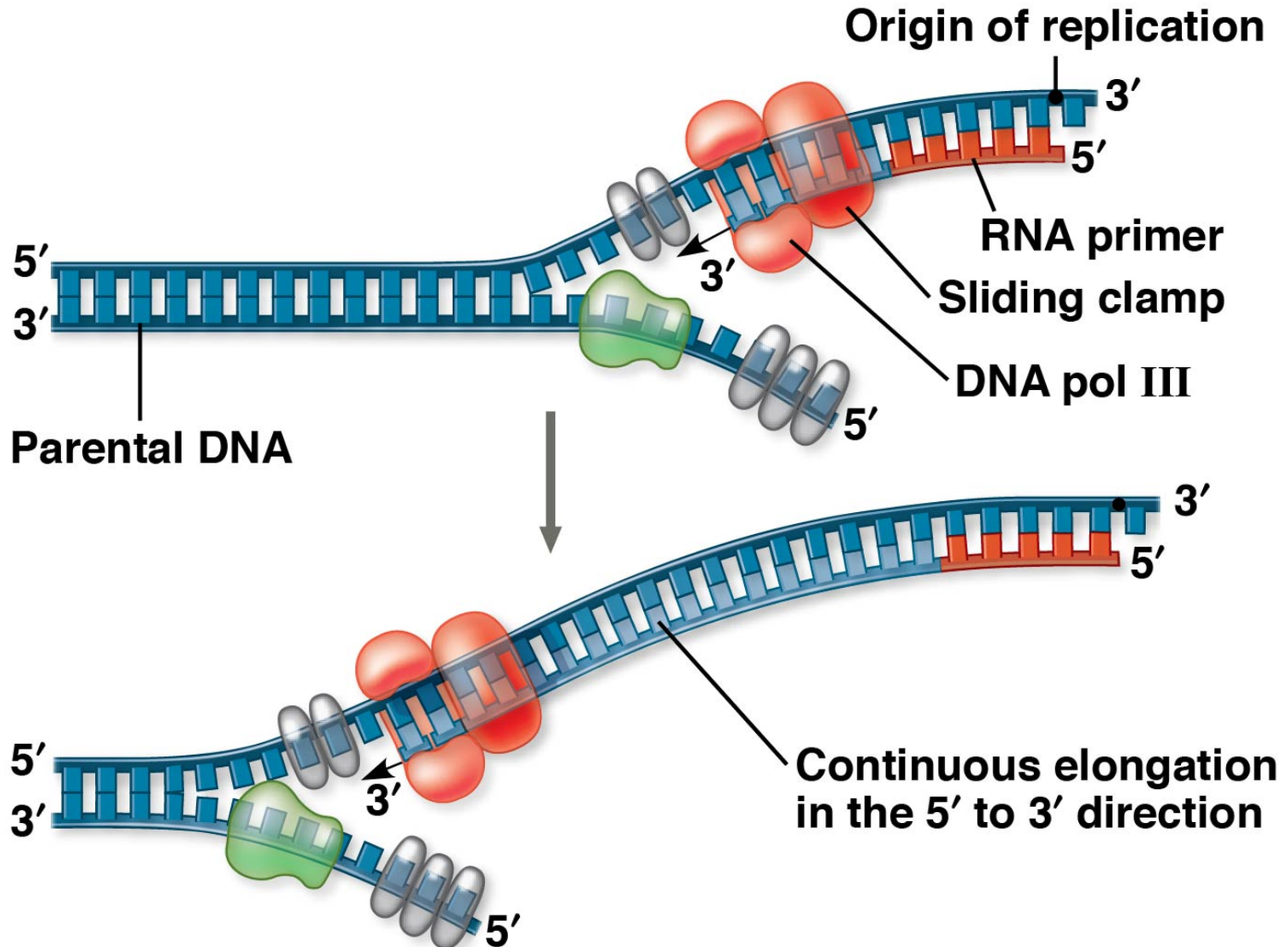
5. DNA polymerase III adds nucleotides in 5'→3' direction on *leading strand*



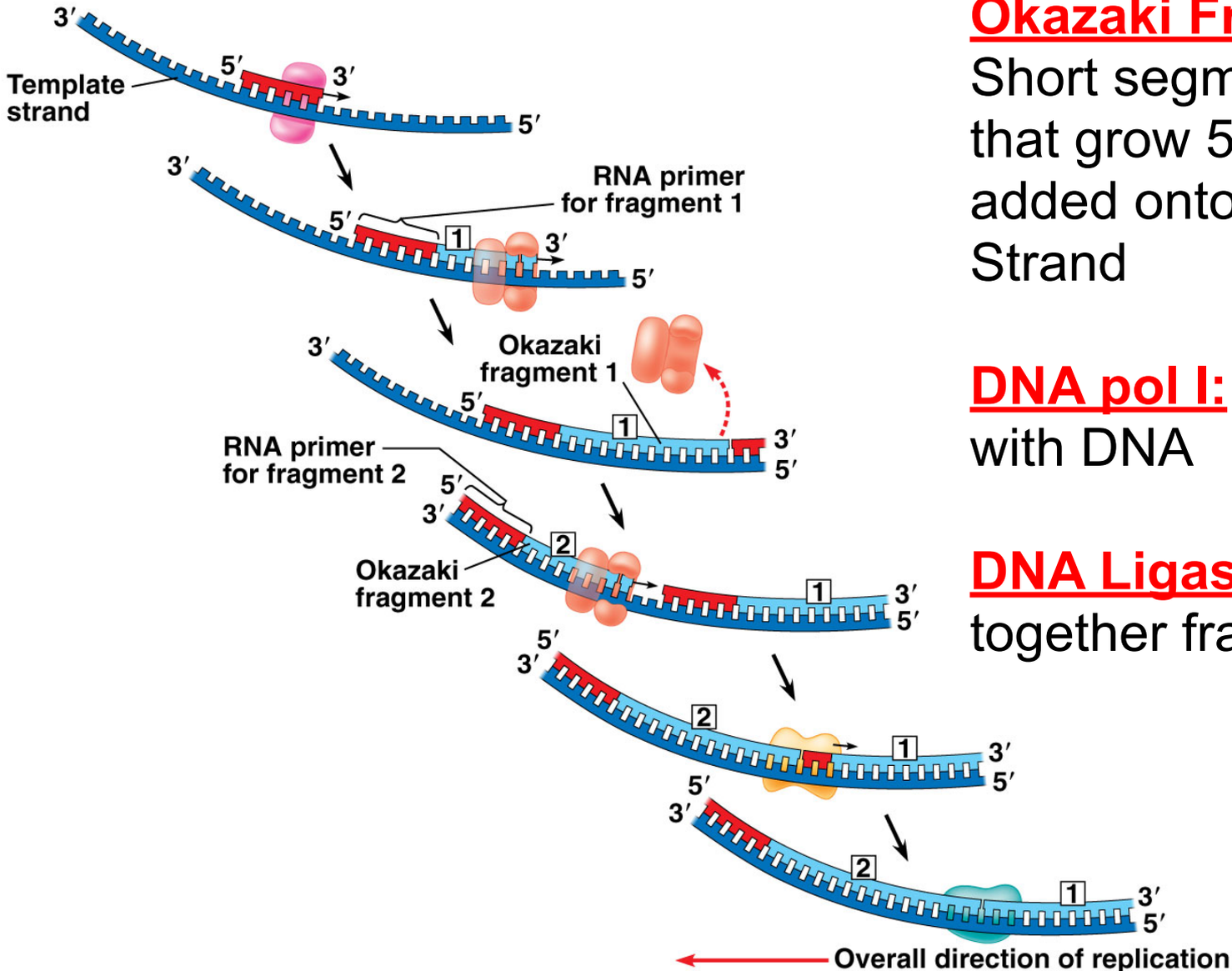
Leading strand vs. Lagging strand



Replication on leading strand



Replication on lagging strand



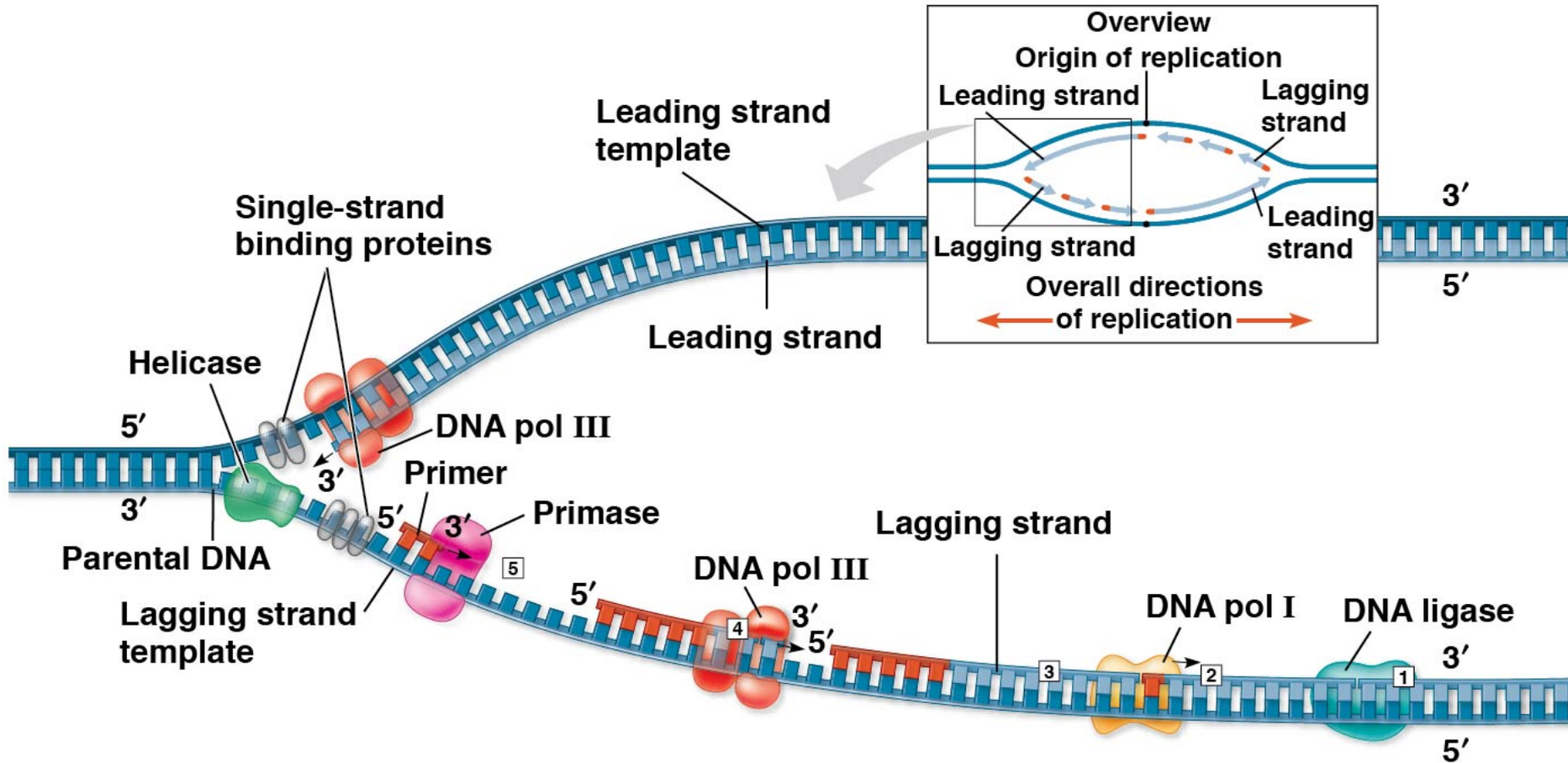
Okazaki Fragments:

Short segments of DNA that grow 5' → 3' that are added onto the Lagging Strand

DNA pol I: replace RNA with DNA

DNA Ligase: seals together fragments

Summary of DNA Replication

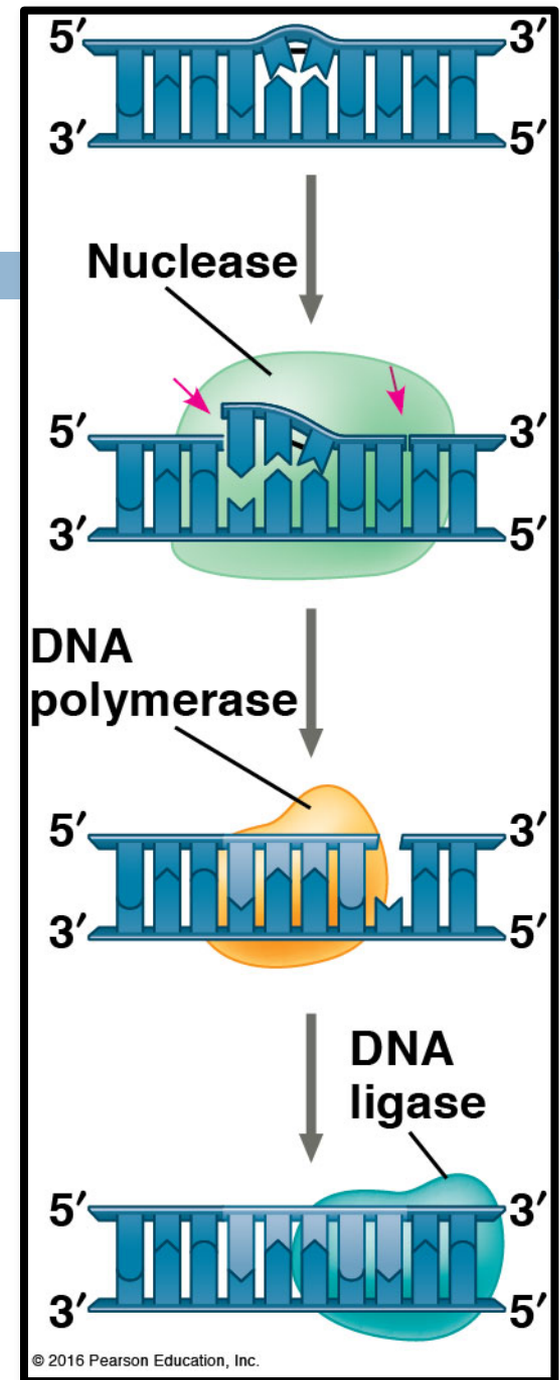


Proofreading and Repair

- DNA polymerases proofread as bases added
- Errors:
 - Pairing errors: 1 in 100,000 nucleotides
 - Complete DNA: 1 in 10 billion nucleotides
- Mismatch repair: special enzymes fix incorrect pairings

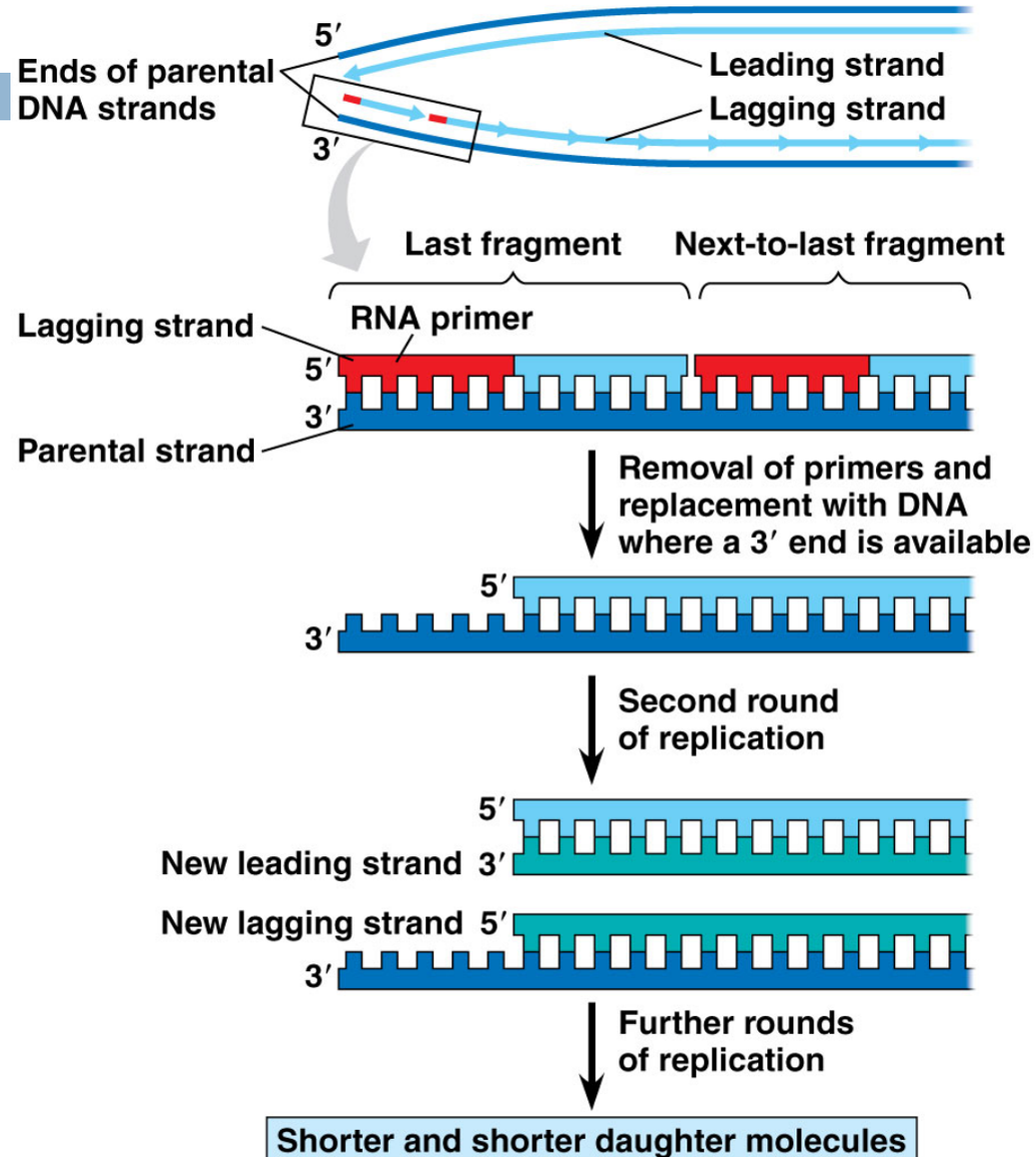
Nucleotide Excision Repair

- **Nucleases** cut damaged DNA
- DNA poly and ligase fill in gaps



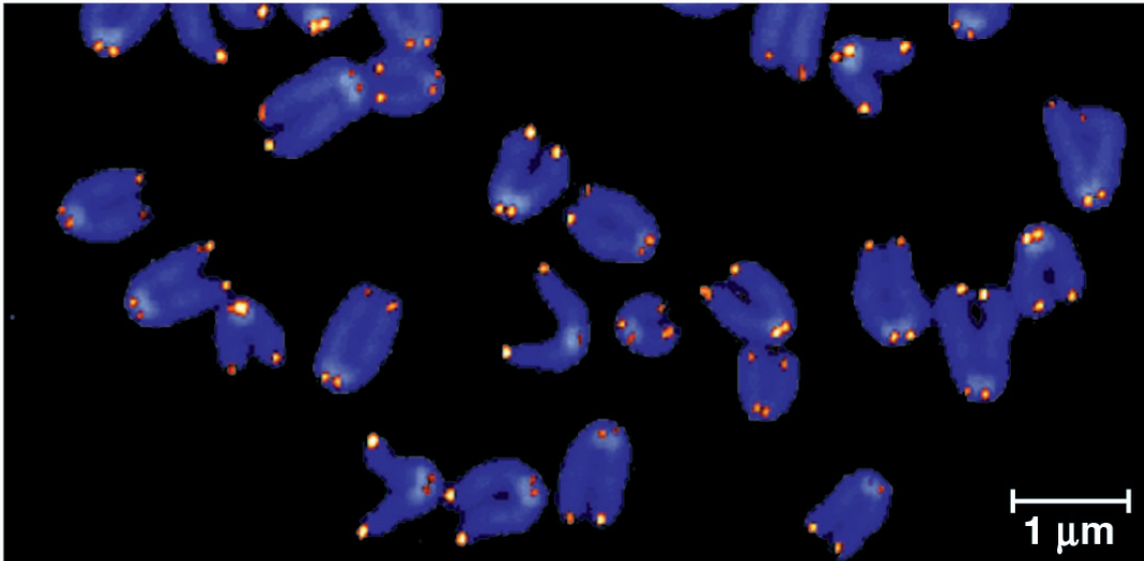
Problem at the 5' End

- DNA poly only adds nucleotides to 3' end
- No way to complete 5' ends of daughter strands
- Over many replications, DNA strands will grow shorter and shorter



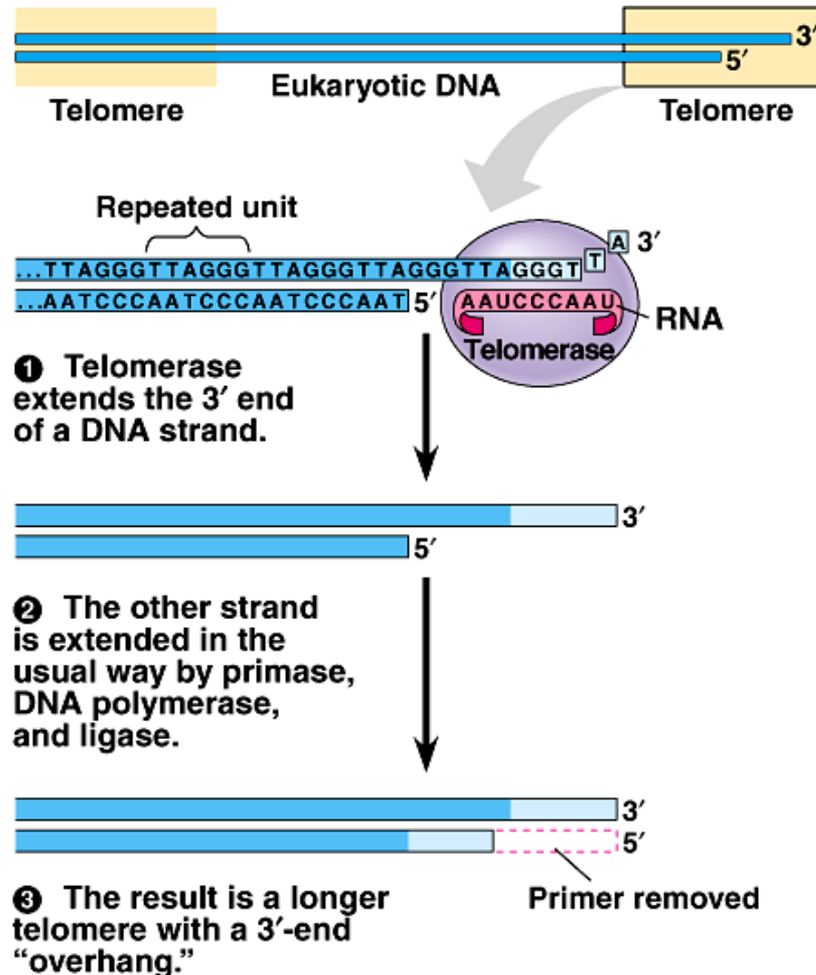
Telomeres: repeated units of short nucleotide sequences (TTAGGG) at ends of DNA

- Telomeres “cap” ends of DNA to postpone erosion of genes at ends (TTAGGG)
- **Telomerase**: enzyme that adds to telomeres
 - Eukaryotic germ cells, cancer cells



Telomeres stained orange at the ends of mouse chromosomes

Telomeres & Telomerase



(b)

BioFlix: DNA Replication

[http://media.pearsoncmg.com/bc/
bc_0media_bio/bioflix/bioflix.htm?8apdnarep](http://media.pearsoncmg.com/bc/bc_0media_bio/bioflix/bioflix.htm?8apdnarep)