



# Recombinant DNA technology and DNA cloning

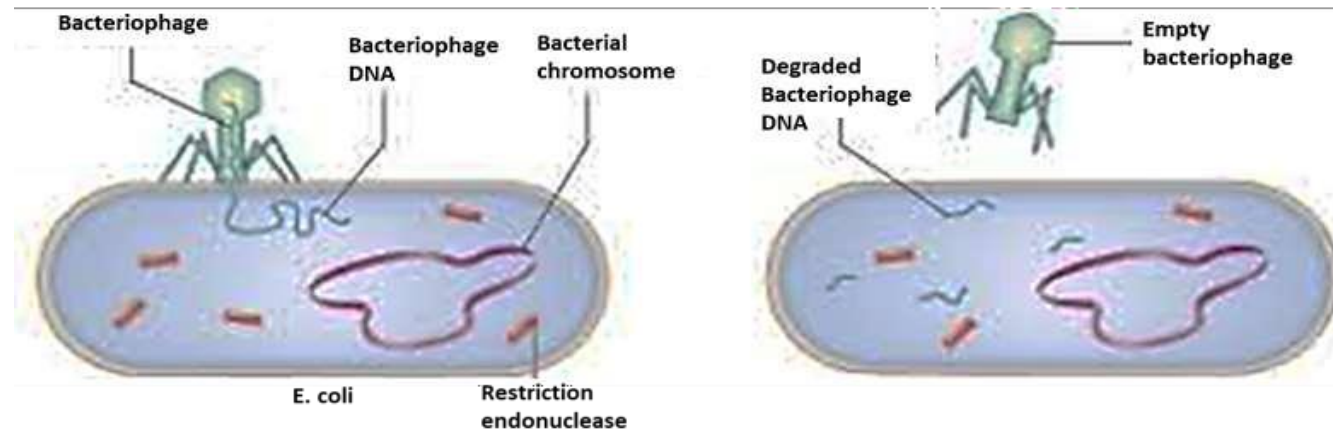
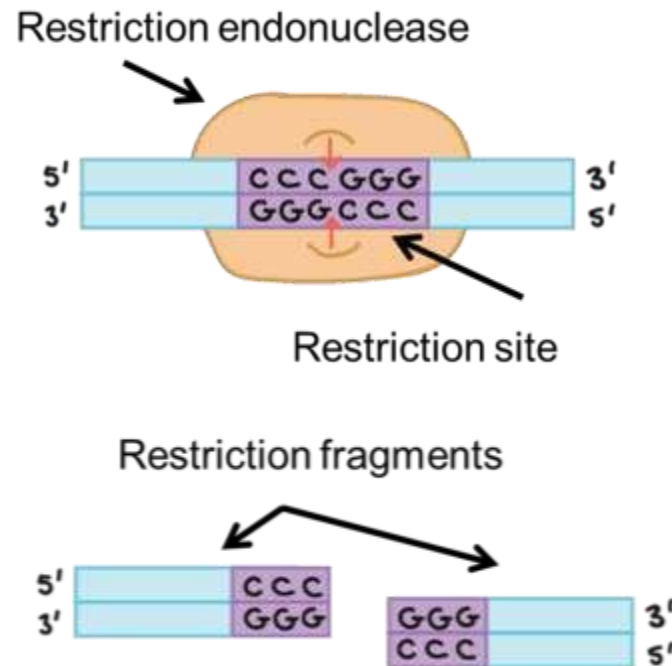
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School of Medicine  
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# What is DNA cloning?

- DNA cloning is a technique that allows for:
  - amplifying a DNA segment into many, many copies in a biological system.
  - expressing a gene inside a biological system such as bacteria, human cells grown in labs, animals, or even the human body as a whole.
- It usually involves:
  - The formation of a recombinant DNA composed of **a vector** (a carrier of the gene or the DNA segment of interest; usually a bacterial plasmid) and **a gene that encodes a protein or a non-coding RNA** using restriction endonucleases.
  - Insertion into the cell(s).

# Restriction endonucleases

- Endonucleases are enzymes that degrade DNA within the molecule.
- Restriction endonucleases: Bacterial enzymes that recognize and cut (break) the **phosphodiester bond** between nucleotides at specific sequences (4- to 8-bp **restriction sites**) generating **restriction fragments**.



# Palindromic sequences



- The sequences recognized by restriction endonucleases—their sites of action—read the same from left to right as they do from right to left (on the complementary strand).

**EcoRI**

5' GAATTC 3'  
3' CTTAAG 5'

**HindIII**

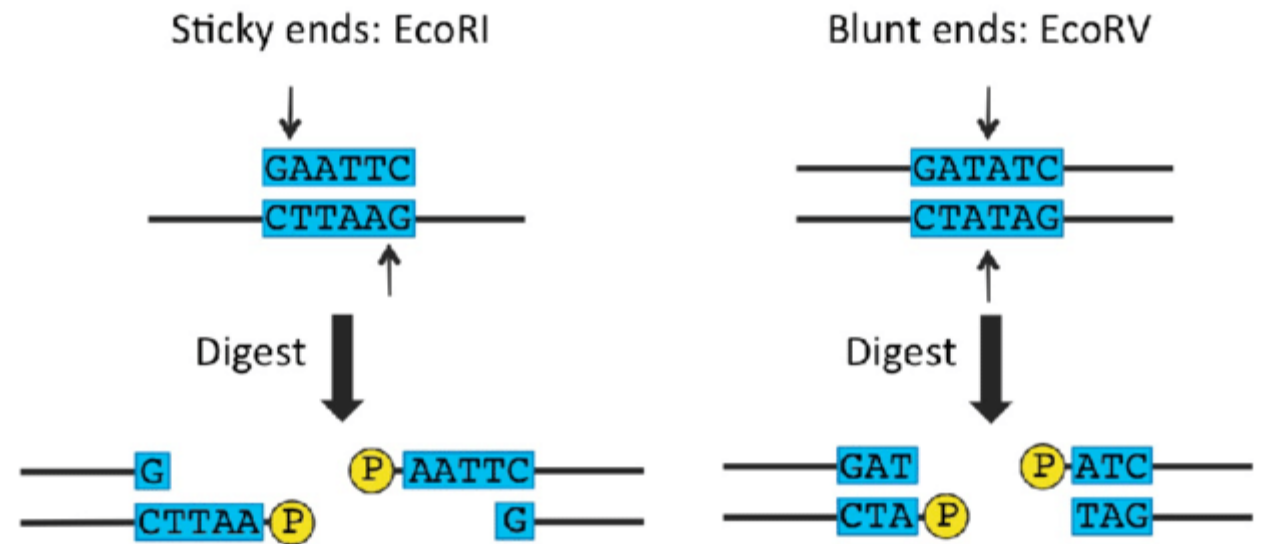
5' AAGCTT 3'  
3' TTCGAA 5'

**SmaI**

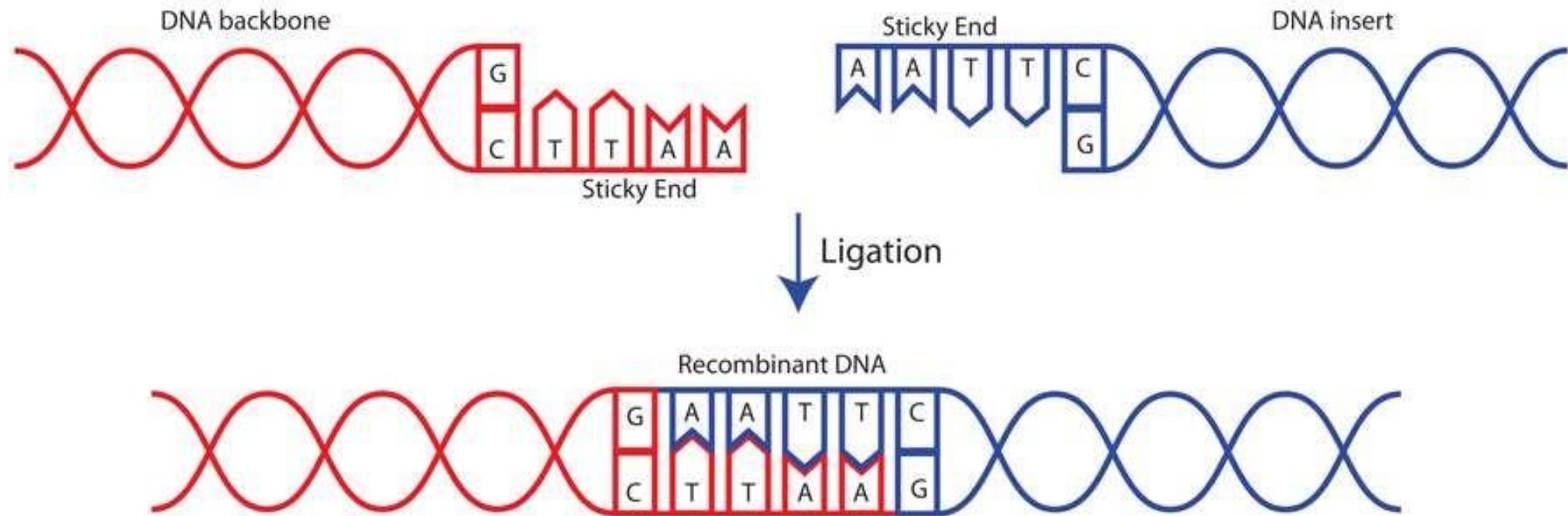
5' CCCGGG 3'  
3' GGGCCC 5'

# Types of cuts by restriction endonucleases

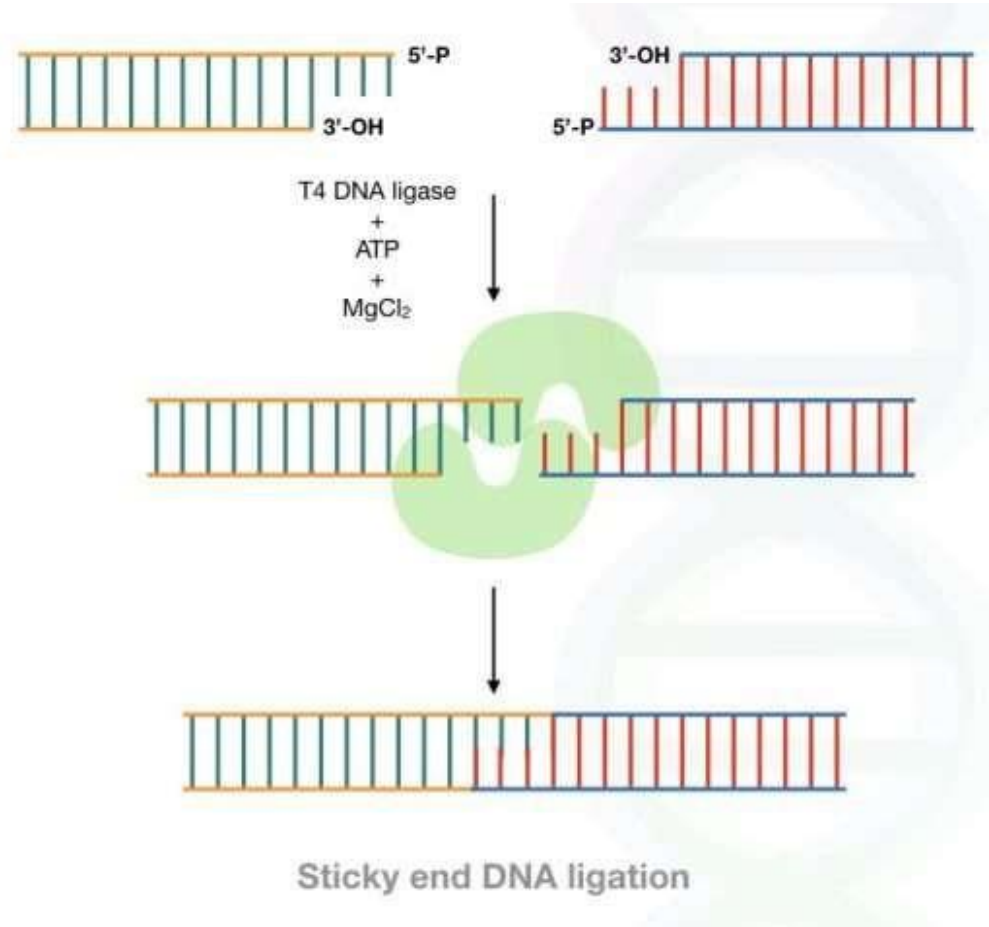
- Staggered (off-center): enzymes cut the two DNA strands at different positions generating sticky or cohesive ends.
- The DNA restriction fragments would have short single-stranded overhangs at each end.
- Blunt: enzymes cut at the same position on both strands giving blunt-ended fragments.



# Zoom into the sticky ends



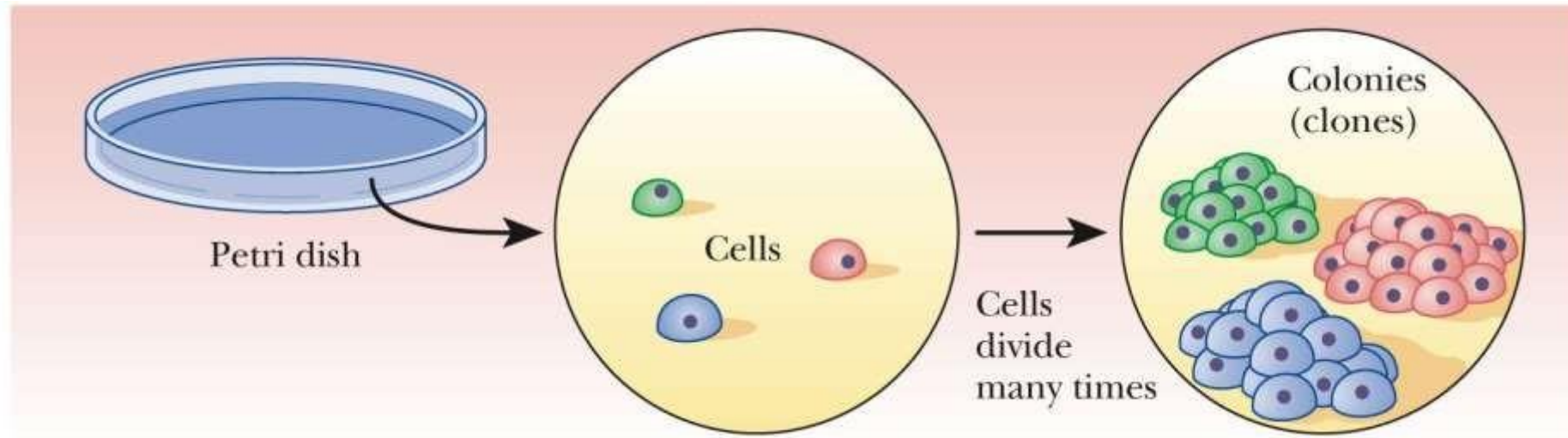
# DNA ligase



It covalently joins DNA ends (example, restriction fragments) by catalyzing the ATP-dependent formation of phosphodiester bonds between the 3'-hydroxyl group of one strand and the 5'-phosphate end of another strand.

# Cloning

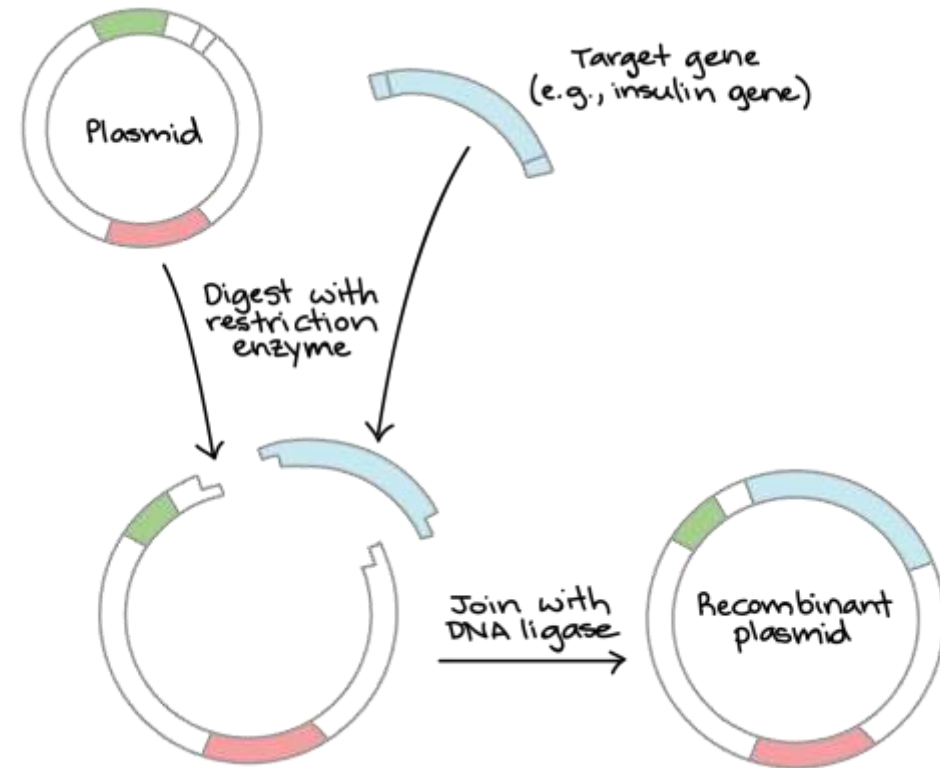
- Cloning means that you make several copies of one thing.
- A clone is a genetically identical population, whether of organisms, cells, viruses, or DNA molecules.
- Every member of the population is derived from a single cell, virus, or DNA molecule.





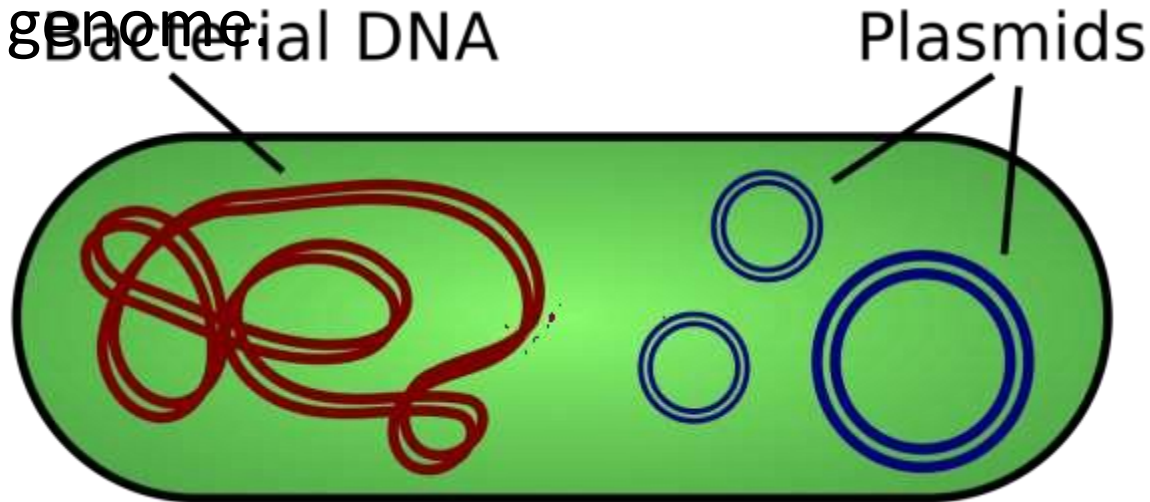
# How do we clone a DNA molecule?

- a DNA fragment of interest is inserted into a DNA carrier (called a **vector**) that can be replicated.
- The resulting DNA molecule is what is known as a **recombinant DNA molecule**.
- The procedure is known as **recombinant DNA technology**, which is part of genetic engineering.



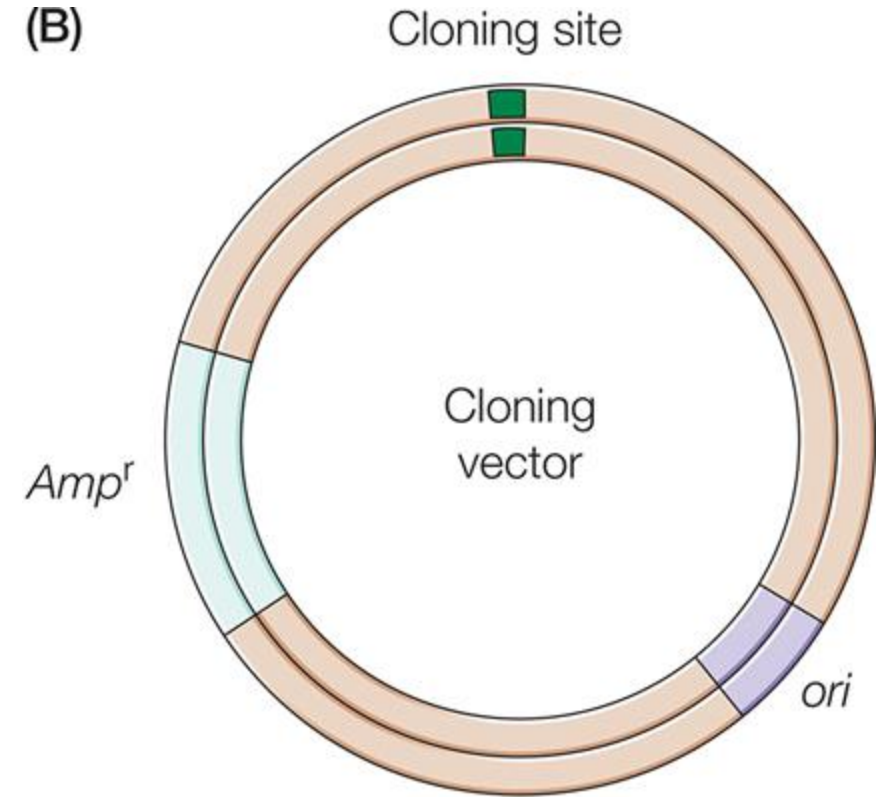
# Using plasmids as vectors

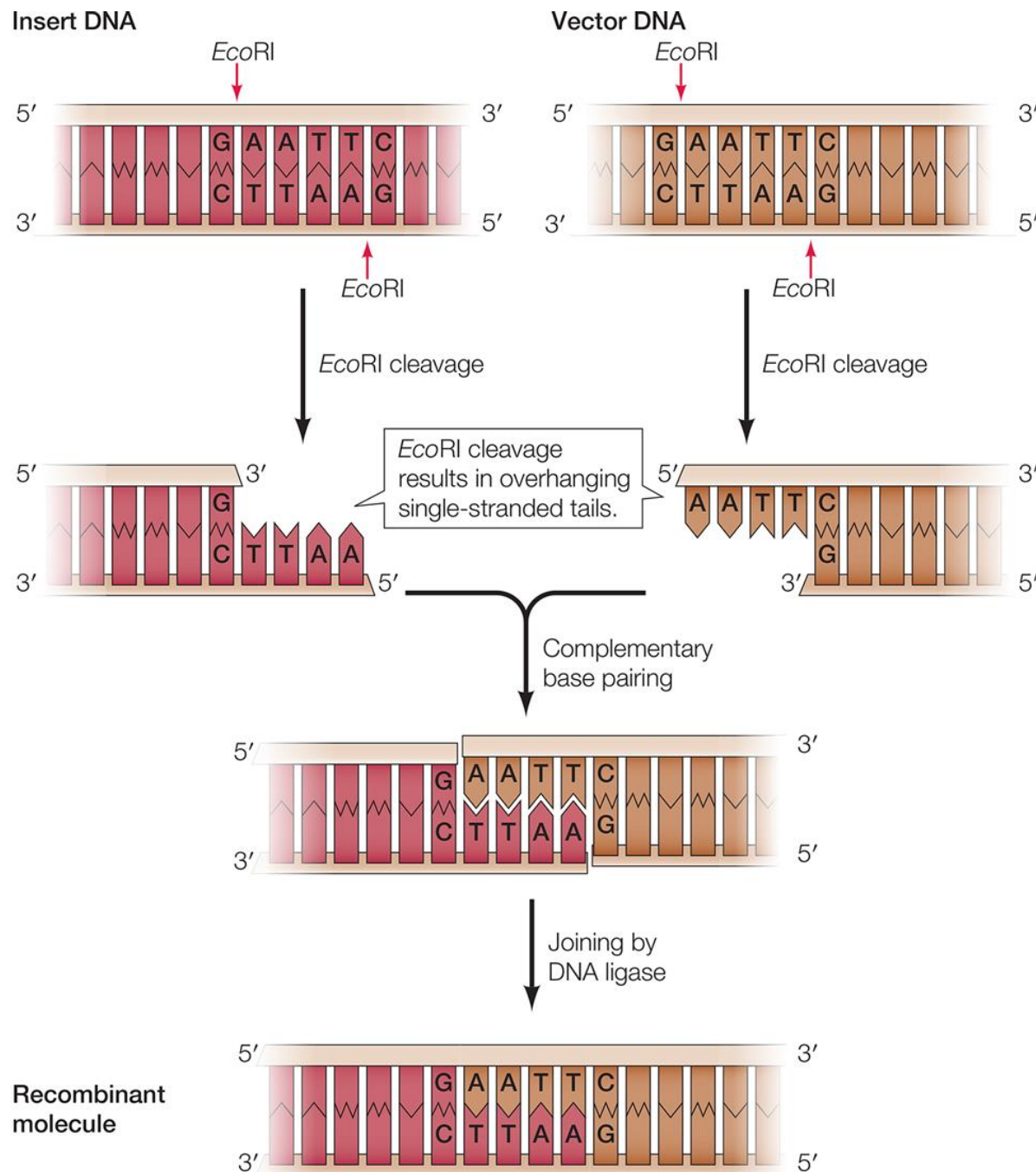
- Bacterial plasmids are considered excellent vectors that are used for cloning (**cloning vectors**) or expression (**expression vectors**).
- These are natural bacterial circular DNA that is not part of the main circular DNA chromosome of the bacterium.
- A plasmid exists as a closed circle and replicates **independently** of the main bacterial genome.



# Features of plasmid cloning vectors

- Plasmid cloning vectors must have the following three components:
  - Their own origin of replication (OriC) that allows them to replicate independently of the bacterial chromosome.
  - Aselectable gene such as an antibiotic resistance gene that allows for selecting for/against the cells that have them.
  - A restriction site that allows for insertion of the DNA segment of interest into the plasmid.



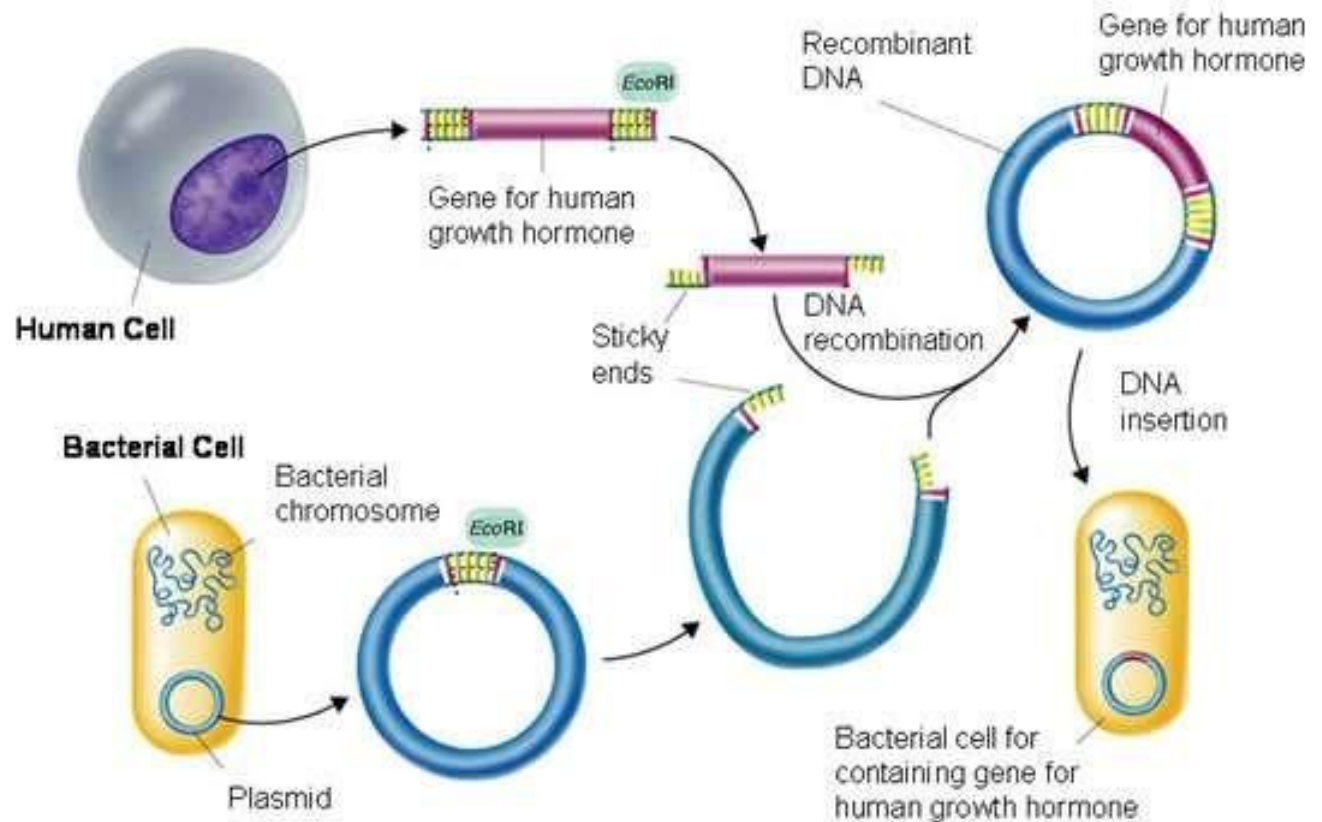


## Creation of a recombinant DNA

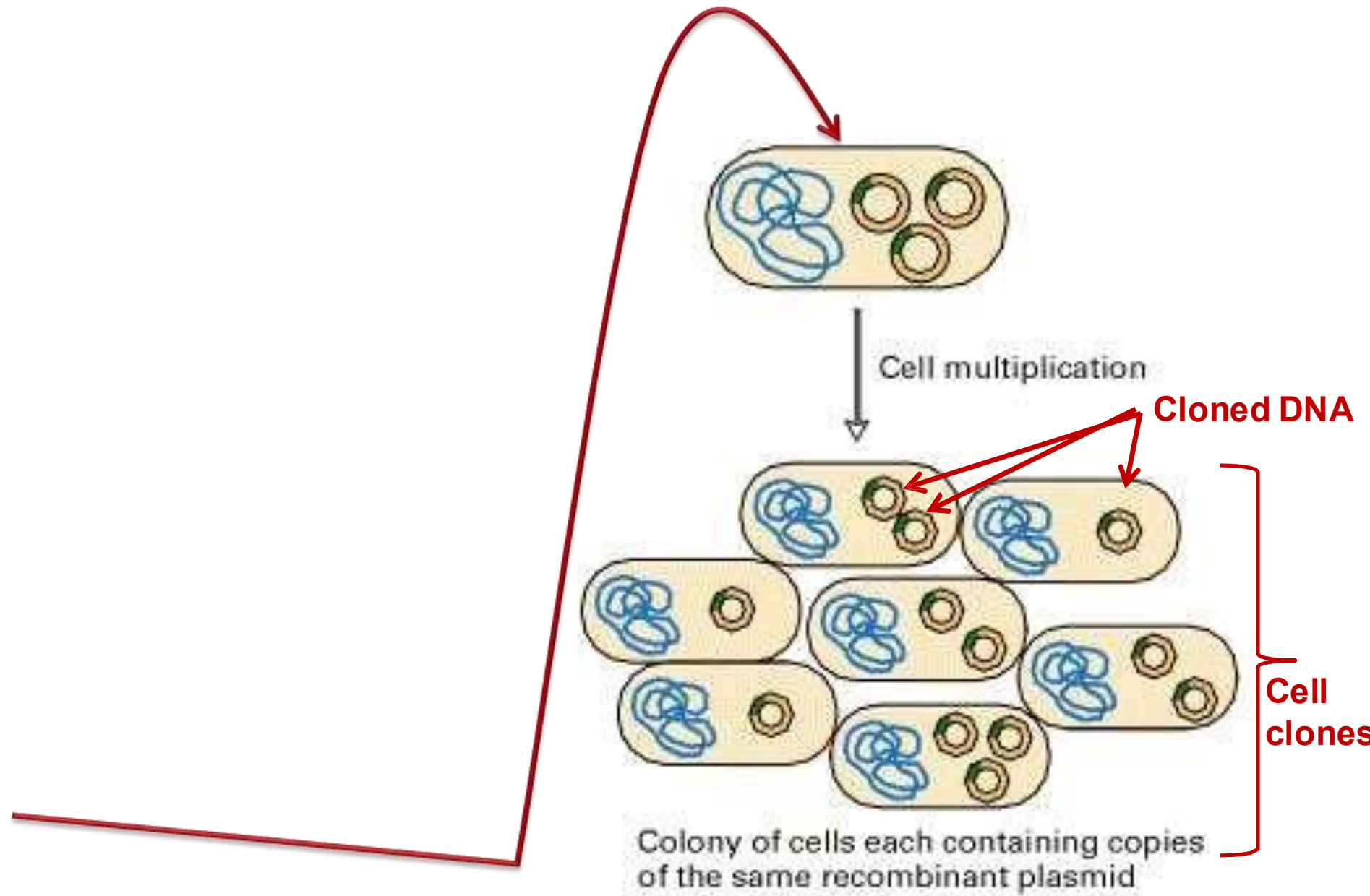
- Insert and vector DNAs are digested with a restriction endonuclease (such as *EcoRI*), which cleaves at staggered sites leaving overhanging single-stranded tails.
- Insert and vector DNAs can then associate by complementary base pairing, and covalent joining of the DNA strands by DNA ligase yields a recombinant molecule.

# The making of a recombinant DNA

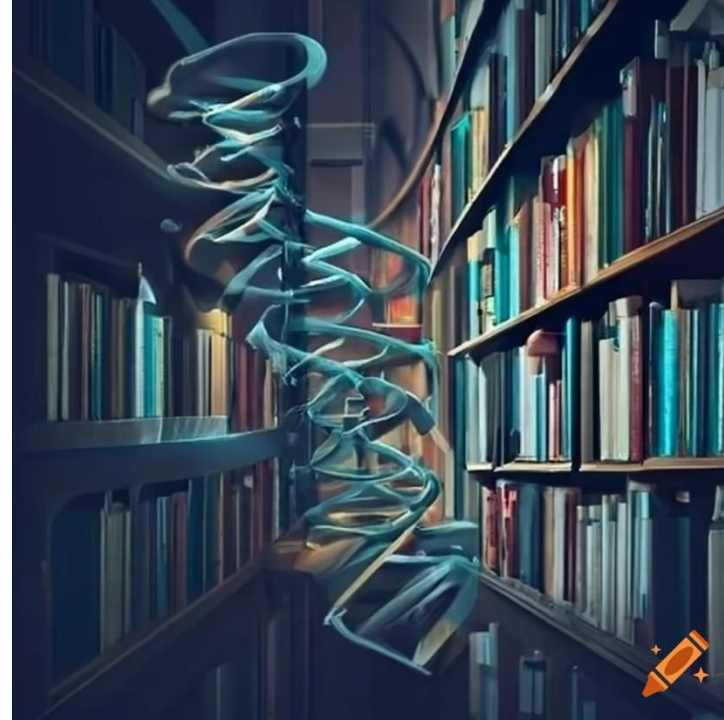
- Both DNA fragments (the DNA to be cloned and a vector) are cut by the same restriction endonuclease that makes DNA fragments with same sticky-ends hybridize (anneal) to each other, when mixed.
- A DNA ligase is added to “close” the plasmid.



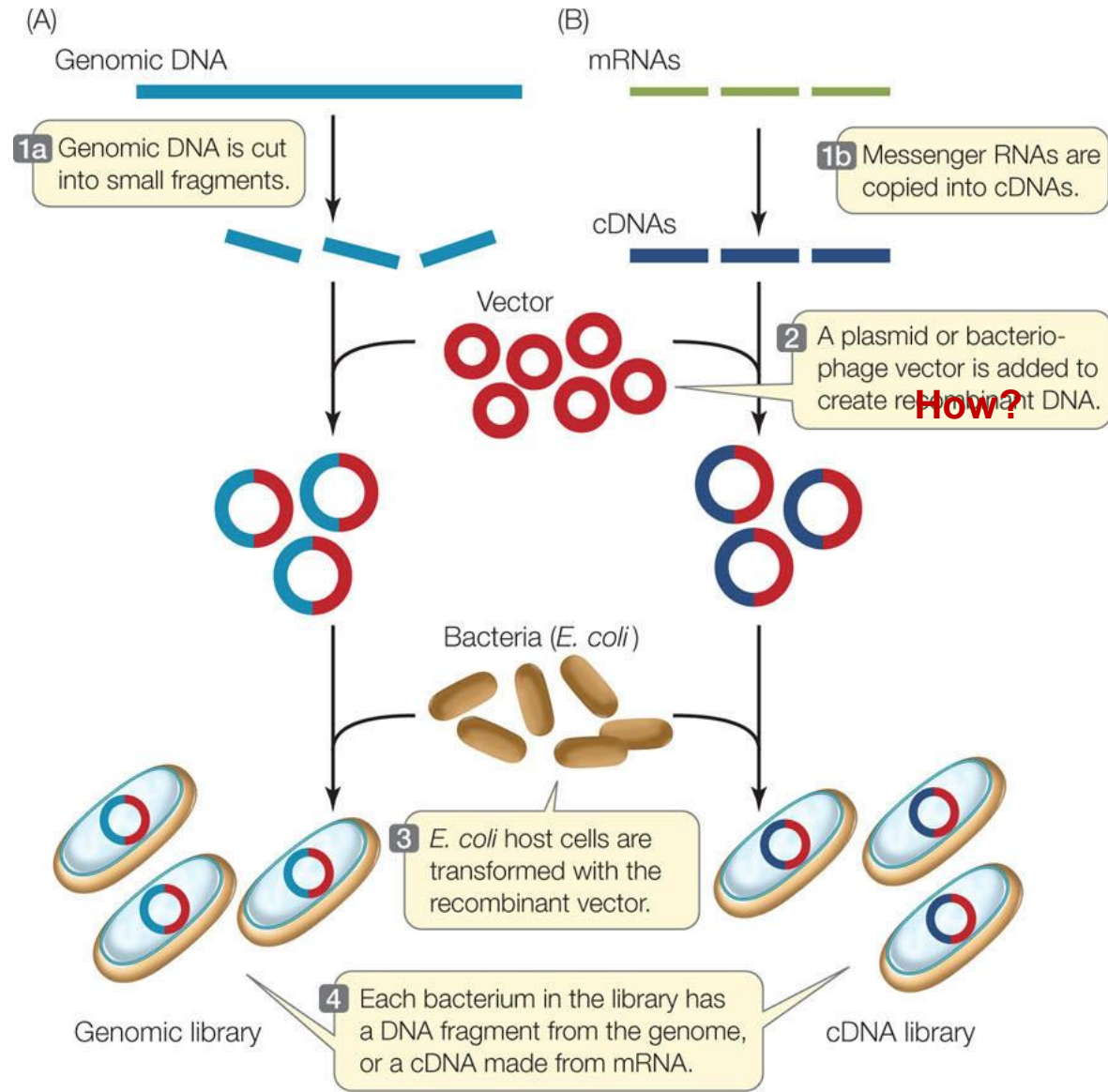




# DNA libraries

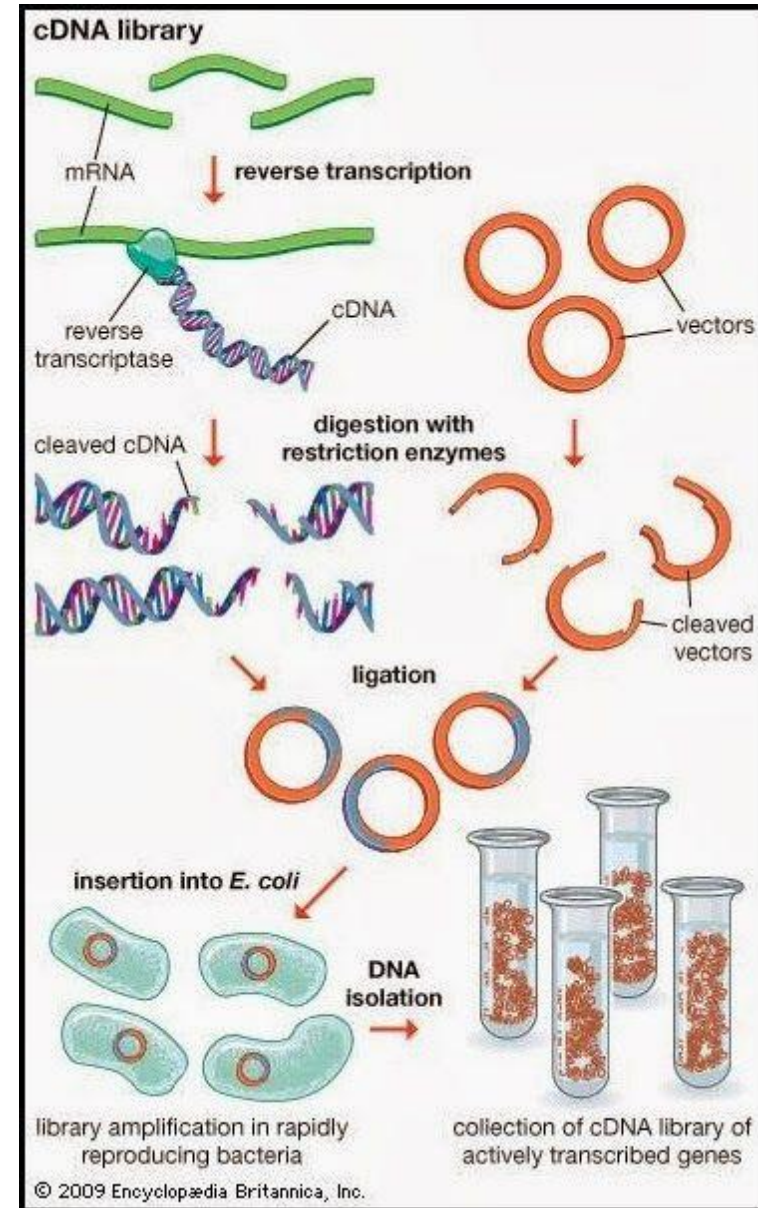
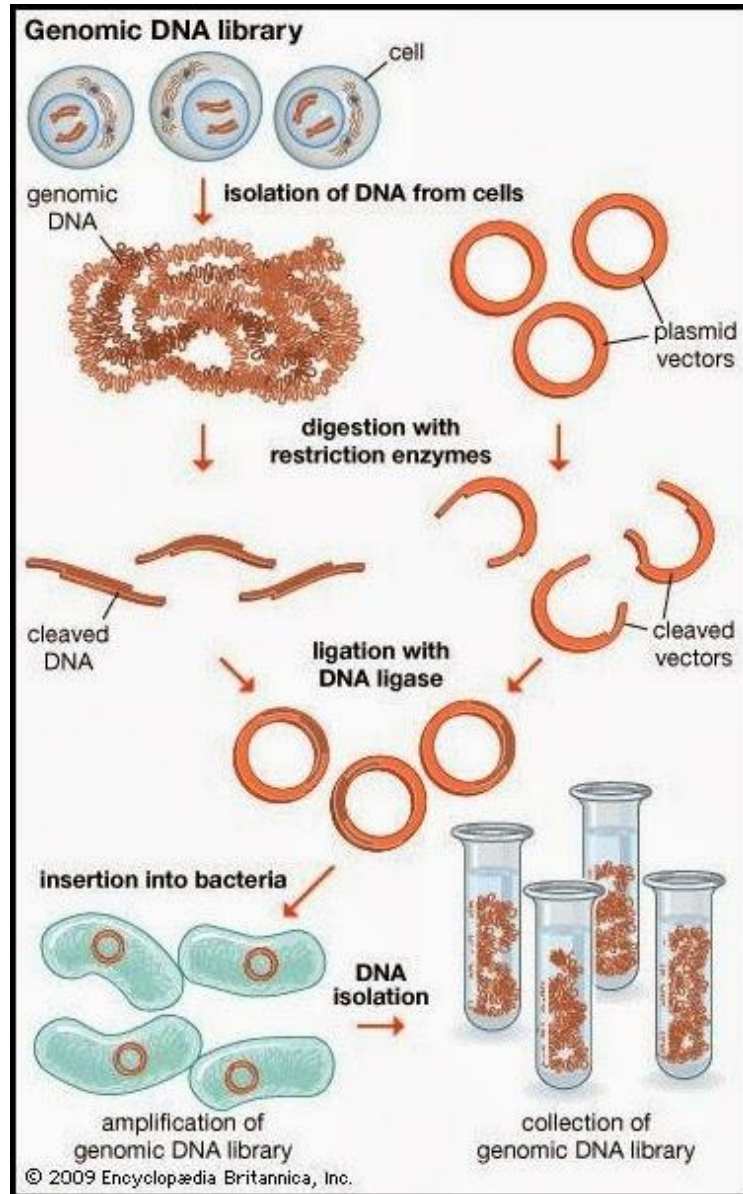


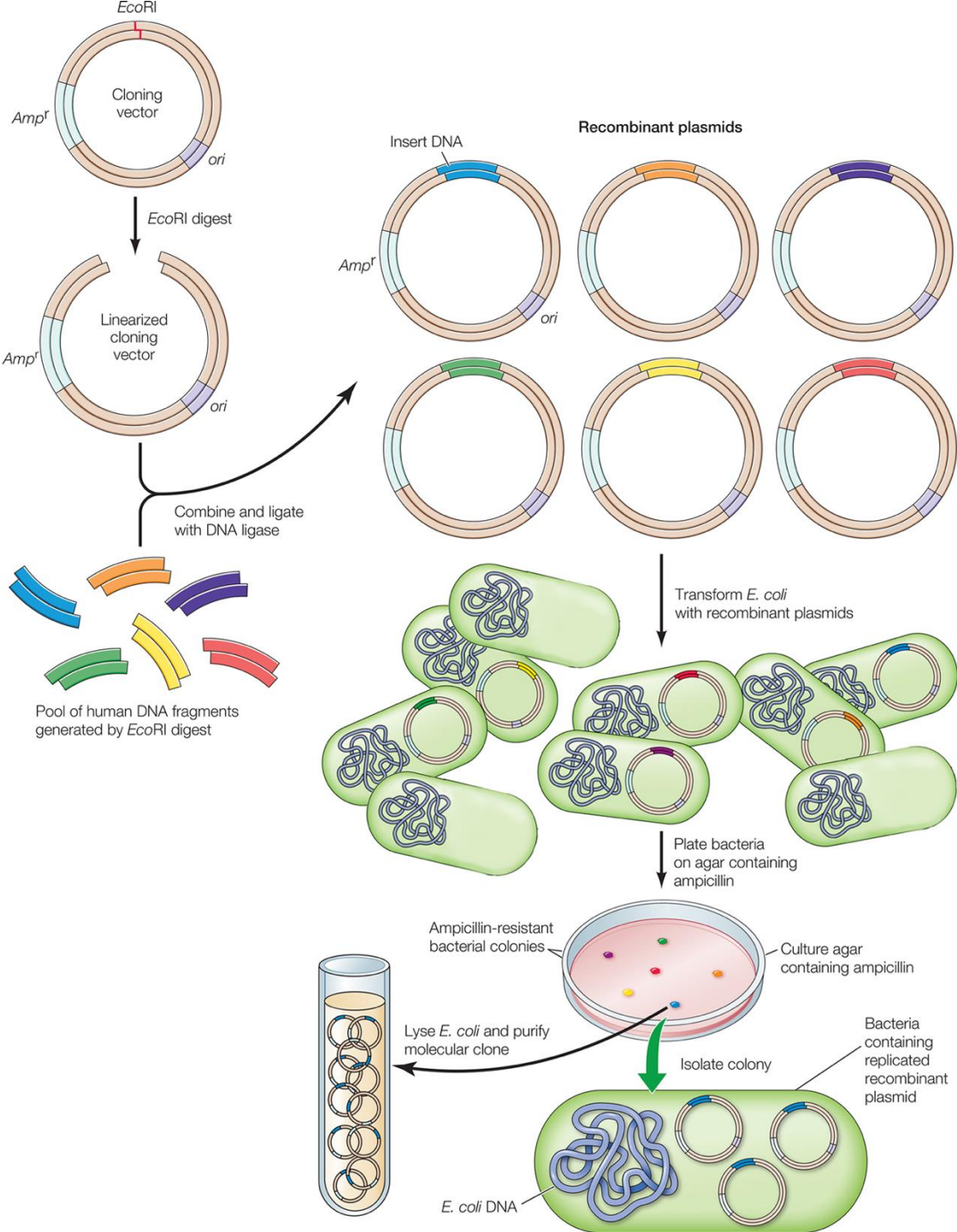
# Genomic vs. cDNA libraries





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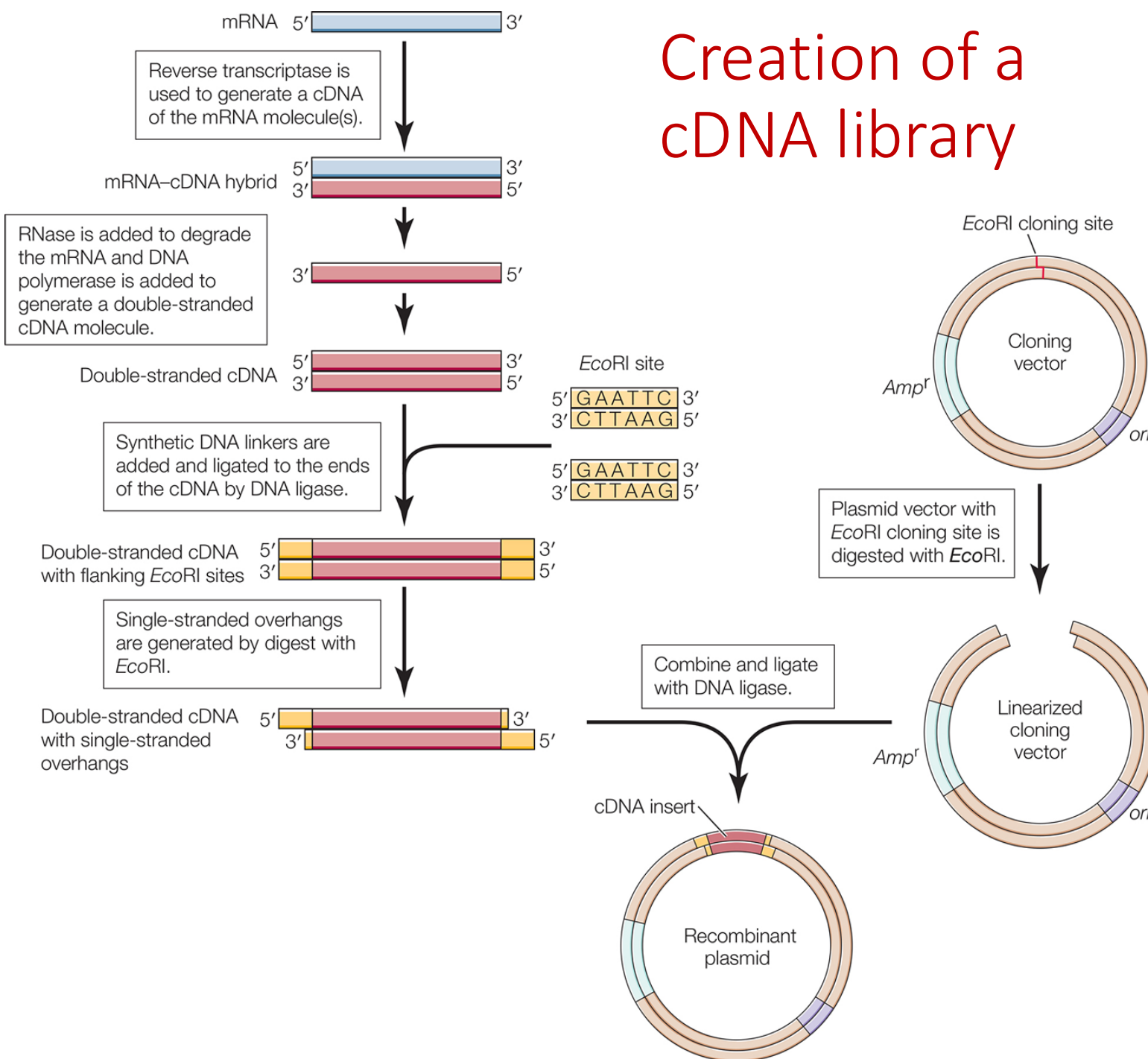


## Creation of a genomic library

- A genome is cleaved by the same restriction endonuclease as that used for the vector.
- Each fragment is ligated into a vector.
- Bacterial cells are transformed with the plasmid vector with each cell having one plasmid DNA.
- Each cell can grow into millions of cells and each cell can make multiple copies of every plasmid ending up with billions of copies of plasmid with each plasmid having a specific DNA fragment.



# Creation of a cDNA library



- Messenger RNAs are isolated and reverse transcribed by reverse transcriptase into a cDNA molecule that is replicated by DNA polymerase to form a double-stranded cDNA.
- Synthetic linkers containing a restriction site are ligated to the ends of the cDNAs and then digested with the restriction endonuclease to form overhangs.
- The cDNAs are then cloned into a plasmid.