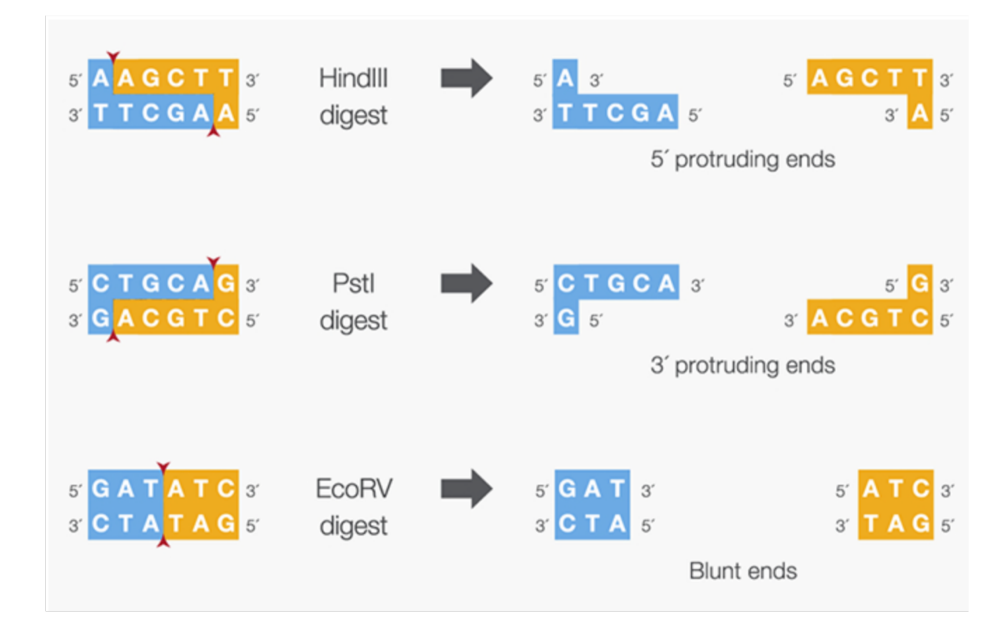
## Endonucelasi di restricion (=enzimi di restrizione) Restriction endonucleases (=restriction enzymes)



## Endonucelasi di restricion (=enzimi di restrizione) Restriction endonucleases (=restriction enzymes)

- Enzymes that cut the double helix of DNA at defined sequences
- 3000 different restriction enzymes exist
- 500 restriction enzymes for the use in the laboratory
- Encoded by prokaryotes (bacteria)

Note: Endonucleases cut DNA at internal position Exonucleases cut DNA starting at termini DNAase: cuts DNA RNase: cuts RNA

## **Discovery of restriction endonucleases**

- Arbor and Dussoix in 1962 discovered that certain bacteria contain Endonucleases which have the ability to cleave DNA.
- In 1970 Smith and colleagues purified and characterized the cleavage site of a Restriction Enzyme.
- Werner Arbor, Hamilton Smith and Daniel Nathans shared the 1978 Nobel prize for Medicine and Physiology for their discovery of Restriction Enzymes.



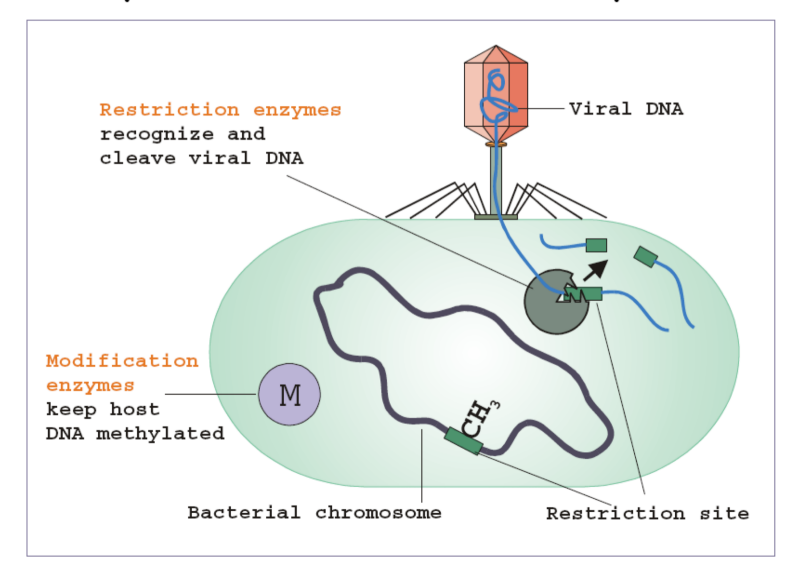
## **Restriction Enzymes**

- Also known as restriction endonucleases
- Scan the DNA sequence
- Find a very specific set of nucleotides
- Make a specific cut (with defined termini or blunt)
- Used to construct recombinant DNA plasmids



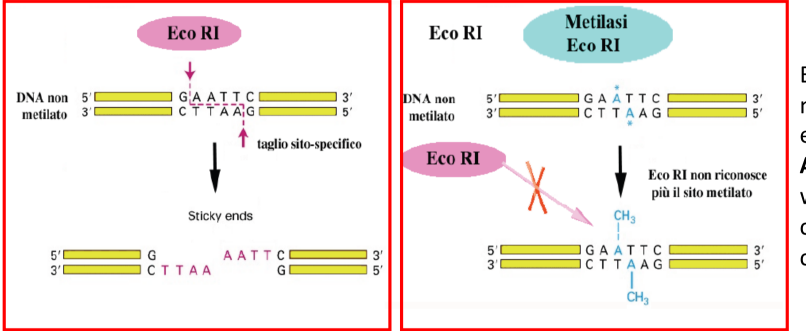
# The origin of restriction endonucleases

Bacterial defense against viral infection by restriction-modification complexes



# The origin of restriction endonucleases

#### Bacteria contain coupled DNA metylation – restriction endonucelase systems



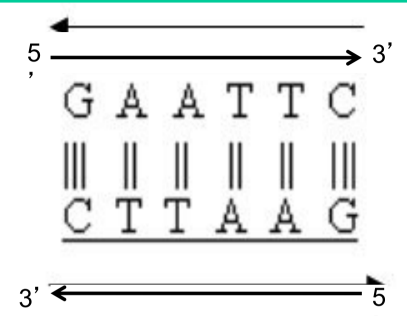
Bacterial DNA methylases can exclusively methylate **A or C**, when present in correct sequence context.

Fage DNA or plasmid DNA with unmethylated or methylated by other DNA methylation system enters bacteria. EcoRI expressed by recipient bacteria cuts at all GAATTC sites in the foreign DNA EcoRI methylase methylates host DNA at defined sequence: GAATTC EcoRI restriction endonuclease expressed by bacteria can cut only un-methylated DNA. DNA integroty is maintained

# **Types of Restriction Enzymes**

	Cleavage site	Location of methylase	Examples
Туре I	Random Around 1000bp away from recognition site	Endonuclease and methylase located on a <b>single</b> <b>protein</b> molecule	EcoK I EcoA I CfrA I
Type II	Specific Within the recognition site	Endonuclease and methylase are <b>separate entities</b>	EcoR I BamH I Hind III
Type III	Random 24-26 bp away from recognition site	Endonuclease and methylase located on a <b>single</b> <b>protein molecule</b>	EcoP I Hinf III EcoP15 I

# Restriction enzymes typically recognize palindromic sequences on double stranded DNA



Genetic palindromes are similar to verbal palindromes. A palindromic sequence in DNA is one in which the 5' to 3' base pair sequence is identical on both strands.

#### (typically 4 or 6 nucleotide palindromes)

4 nucleotide palindromes: 1 cut every 256 nucleotides (4<sup>4</sup>). 6 nuceotide palindromes: 1 cut every 4.096 nucleotides (4<sup>6</sup>).

Enzyme	Target sequence	Cleavage
EcoRI	5' GAATTC 3' 3' CTTAAG 5'	5' G AATTC 3' 3' CTTAA G 5'
EcoRV	5' GATATC 3' 3' CTATAG 5'	5' GAT ATC 3' 3' CTA TAG 5'
HaellI	5' GGCC 3' 3' CCGG 5'	5' GG CC 3' 3' CC GG 5'
HindIII	5' AAGCTT 3' 3' TTCGAA 5'	5' A AGCTT 3' 3' TTCGA A 5'
PpuMI	5' RGGWCCY 3' 3' YCCWGGR 5'	5' RG GWCCY 3' 3' YCCWG GR 5'

Single Letter Code List
$\mathbf{B} = \mathbf{C} \text{ or } \mathbf{G} \text{ or } \mathbf{T}$
$\mathbf{D} = \mathbf{A} \text{ or } \mathbf{G} \text{ or } \mathbf{T}$
H = A  or  C  or  T
$\mathbf{\kappa} = \mathbf{G} \text{ or } \mathbf{T}$
M = A or C
$\mathbf{N} = \mathbf{A} \text{ or } \mathbf{C} \text{ or } \mathbf{G} \text{ or } \mathbf{T}$
$\mathbf{R} = \mathbf{A} \text{ or } \mathbf{G}$
s = C or G
$\mathbf{v} = \mathbf{A} \text{ or } \mathbf{C} \text{ or } \mathbf{G}$
$\mathbf{w} = \mathbf{A} \text{ or } \mathbf{T}$
$\mathbf{Y} = \mathbf{C} \text{ or } \mathbf{T}$

number

Small

#### Sistemi di restrizione-modificazione

#### Tipo I

•Un singolo enzima contiene attività di restrizione e di metilazione su subunità diverse

•Il taglio viene effettuato in modo non specifico lontano dalla sequenza di riconoscimento (da 100 fino a 1000 bp a valle)

• Mg<sup>2+</sup>, ATP e S-adenosilmetionina come cofattori

#### **Tipo II**

•Due enzimi distinti per il taglio e la metilazione.

- •Non richiedono cofattori se non Mg2+
- •Riconoscono la stessa sequenza **palindromica** e agiscono al suo interno

#### Tipo III

- •Caratteristiche analoghe a quelli di tipo I
- •Riconosce e modifica una sequenza **palindromica** ma taglia a 25-27 bp di distanza

#### Tipo IIs

- Due enzimi separati che riconoscono una sequenza non palindromica
- Tagliano su di un solo lato delle sequenza bersaglio entro 20 bp

1. Le prime tre lettere, scritte in corsivo, sono prese da genere e specie del batterio di origine.

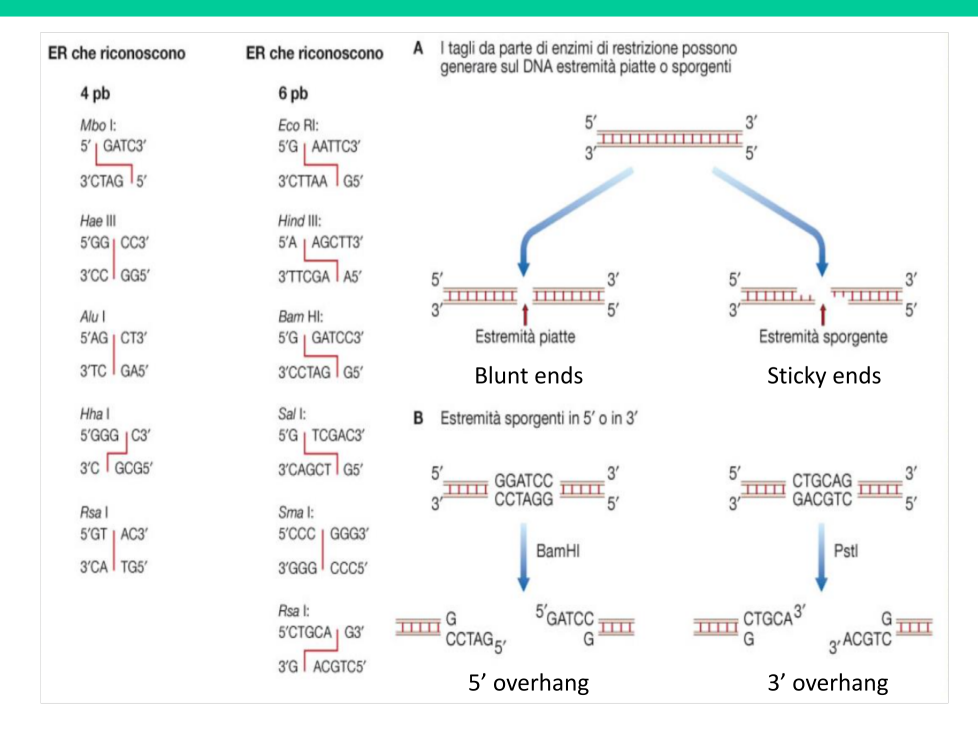
2. Sierotipi differenti dello stesso organismo possono essere identificati da una quarta lettera minuscola (Es. *Hind, Hinf*).

3. Può seguire una lettera maiuscola o un numero, che identifica un ceppo particolare di quel batterio.

4. Un numero romano indica l'ordine di scoperta, qualora dallo stesso batterio vengano isolati enzimi diversi.

Enzima	Organismo di provenienza
Smal	Serratia marcescens, 1º enzima
Haelli	Haemophilus aegyptius, 3° enzima
Hindl	Haemophilus influenzae, ceppo d, 2º enzima
HindIII	Haemophilus influenzae, ceppo d, 3º enzima
BamHt	Bacillus amyloliquefaciens, ceppo H, 1° enzima

#### **Types of DNA cuts by restriction endonucelases**



### **Isoschizomers and Neochischizomers**

• Restriction enzymes that have the same recognition sequence as well as the same cleavage site are **Isoschizomers**.

CGTACG	CGTACG
GCATGC	GCATGC
Sphl	Bbul

 Restriction enzymes that have the same recognition sequence but cleave the DNA at a different site within that sequence are **Neochizomers.** Eg:Smal and Xmal

CCCGGG	CCCGGG
GGGCCC	GGGCCC
Xma I	Sma I

#### Isocaudomers

Isocaudomers are pairs of restriction enzymes that have slightly different recognition sequences, but upon cleavage of DNA, generate identical overhanging termini sequences. These sequences can be ligated to one another, but then form an asymmetrical sequence that cannot be cleaved by a restriction enzyme.

Mbo I NGATCN NCTAGN BamH I GGATCC CCTAGG

#### **Example of laboratory plasmid sequence**

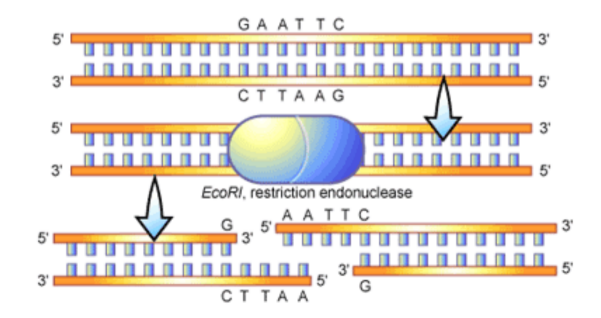


# **Mechanism of Action of restriction endonucleases**

Restriction Endonuclease scan the length of the DNA, binds to the DNA molecule when it recognizes

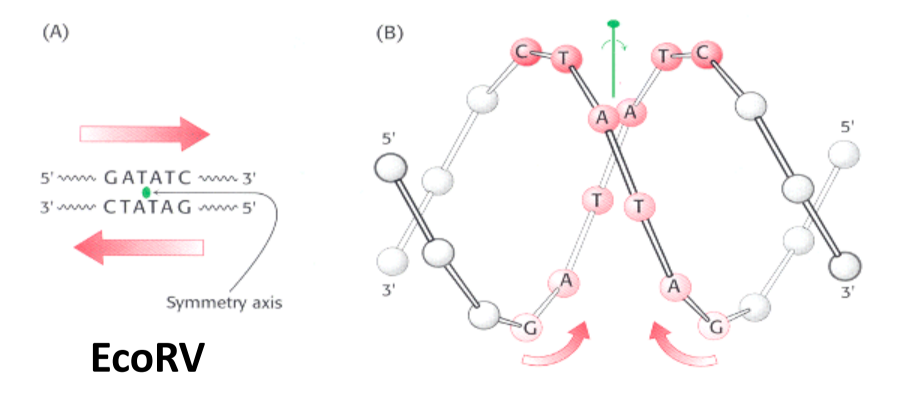
specific sequence and makes one cut in each of the sugar phosphate backbones of the double helix – by

hydrolyzing the phoshphodiester bond. Specifically, the bond between the 3' O atom and the P atom is broken.



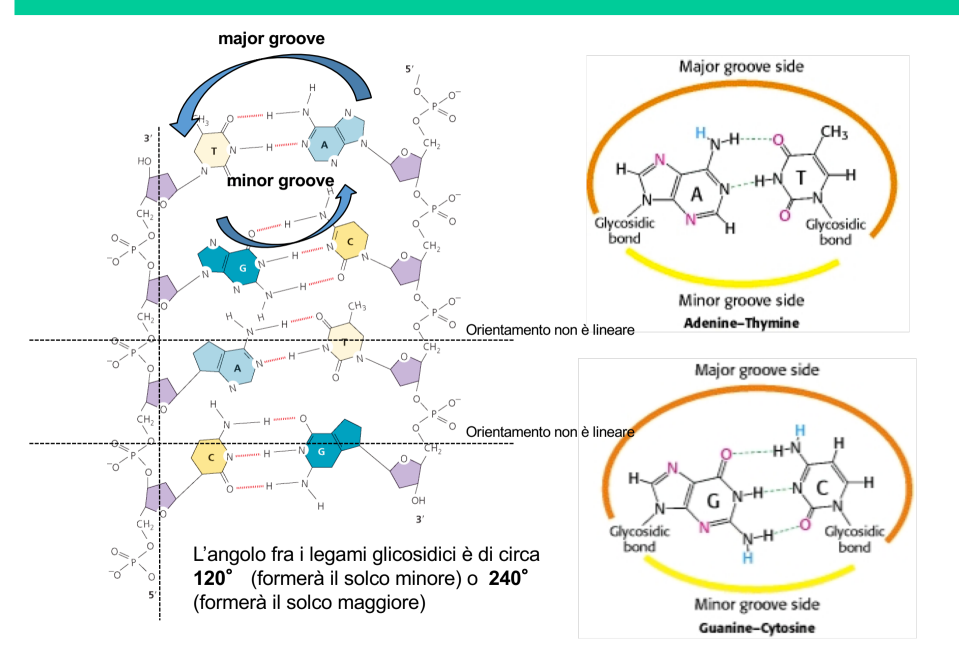
## **Mechanism of restriction endonucleases**

#### Recognition sites of most restriction enzymes have a twofold rotational symmetry



Restriction enzymes have corresponding symmetry to facilitate recognition and usually cleave the DNA on the axis of symmetry

# La doppia elica presenta un solco minore ed un solco maggiore (minor and major groove)



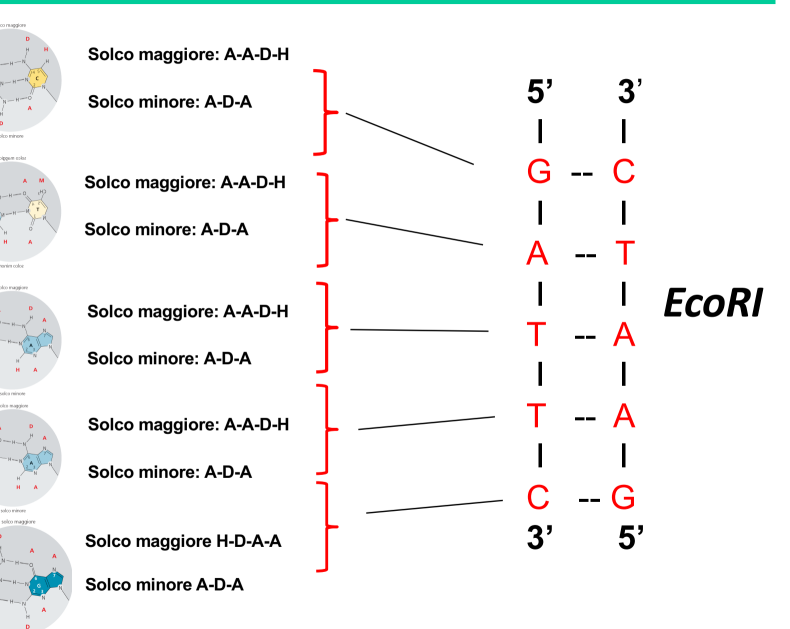
# Restriction enzymes are DNA binding proteins that recoginice specific seqenced by interaction with chemical groups in major and minor groove

A: accettore legame idrogeno D: donatore legame idrogeno M: gruppo metilico H: idrogeno non polare

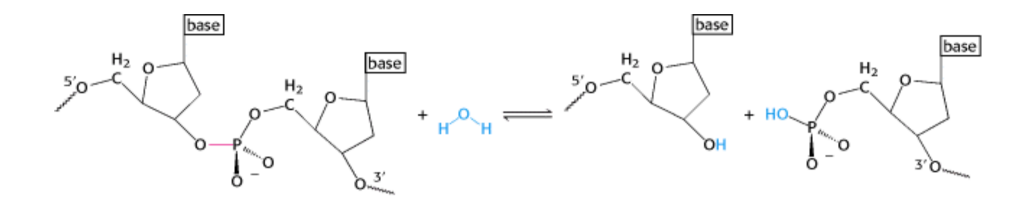
Queste codice formato da gruppi chimici posti al interno del solco maggiore/minore identificano in modo specifico le coppie di basi.

Le proteine possono riconoscere specifiche sequenze di DNA senza che sia necessario aprire o rompere la doppia helica !!! Esempio: →Fattori di trascrizione →Elicasi, etc...

→Endonucleasi



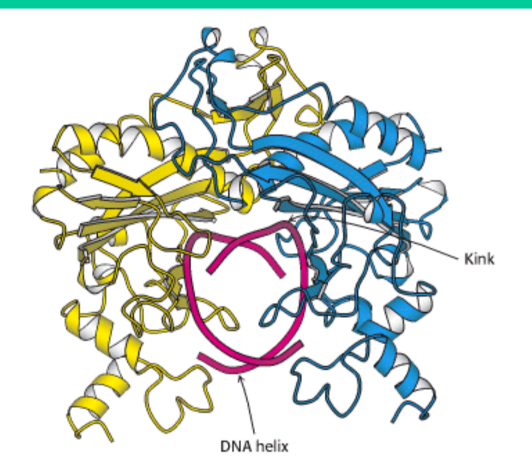
## Direct hydrolysis by nucleophilic attack at the phosphorous atom



3'OH and 5'  $PO_4^{3-}$  is produced. Mg<sup>2+</sup> is required for the catalytic activity of the enzyme.

It holds the water molecule in a position where it can attack the phosphoryl group and also helps polarize the water molecule towards deprotonation .

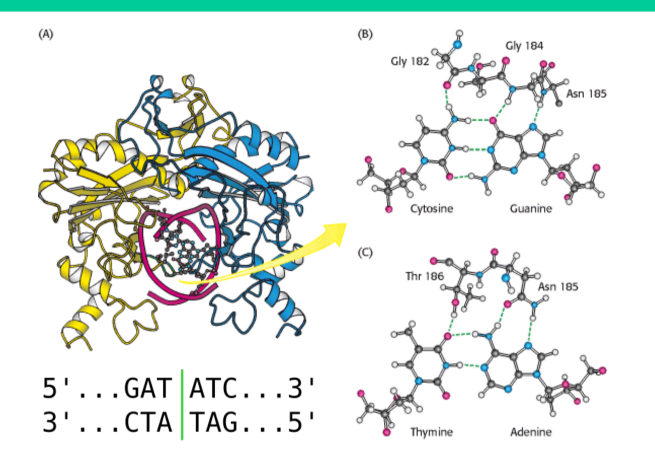
## **Structure - Function of EcoR V endonuclease**



- Consists of two subunits dimers related by two fold rotational symmetry.
- Binds to the matching symmetry of the DNA molecule at the restriction site and produces a kink at the site.

5'...GAT ATC...3' 3'...CTA TAG...5'

### **Structure - Function of EcoR V endonuclease**



Hydrogen bonding interactions between EcoRV and its DNA substrate

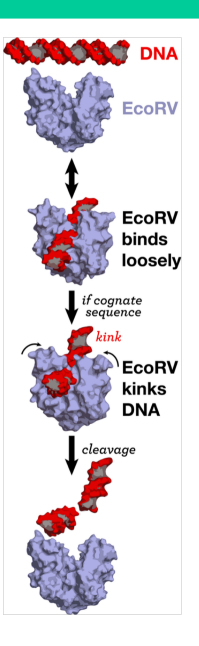
Like EcoRI, EcoRV forms a **homodimer in** solution before binding and acting on its recognition sequence. Initially the enzyme binds weakly to a non-specific site on the DNA. It randomly walks along the molecule until the specific recognition site is found.

EcoRV has a high specificity for its target DNA sequence.Binding of the enzyme induces a conformational change in the DNA, bending it by about 50°. DNA bending results in the **unstacking of the** bases, widening of the minor groove, and compression of the major groove. This brings the phosphodiester linkage to be broken closer to the active site of the enzyme, where it can be cleaved. Cleavage occurs within the recognition sequence, and does not require ATP hydrolysis. EcoRV is the only type II restriction endonuclease known to cause a major protein-induced conformational change in the DNA.

## **Structure - Function of EcoR V endonuclease**



A comparison of cognate and non-specific DNA in the EcorV-DNA complex.



## How to set up a restriction digest

Order of solution addition	Solution	Volume(µl)
1	Nuclease free water	23.5
2	10X Buffer K	5.0
4	100 µg BSA	0.5
5	Plasmid DNA	20.0
3	ScaI(20U/ µl)	1.0
Total Volume		50.0

#### Scal buffer

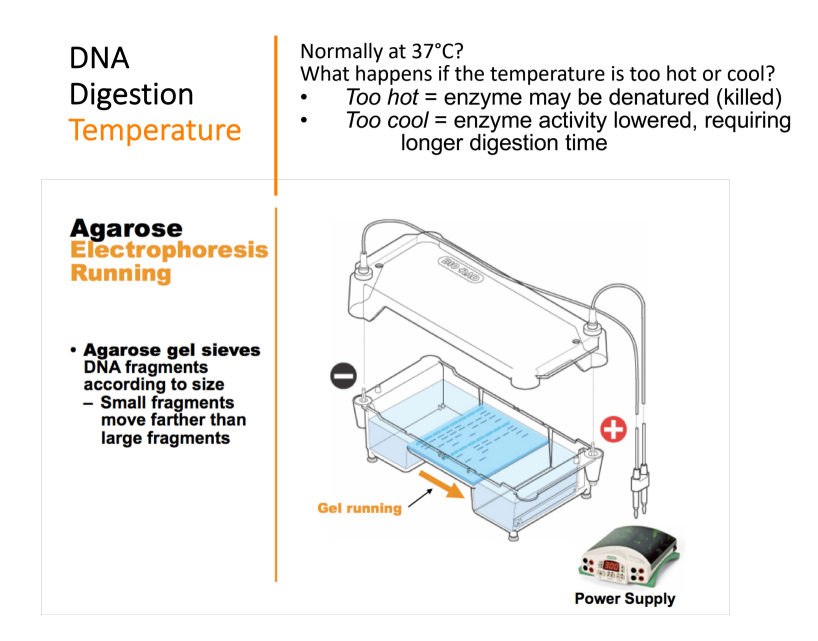
100 mM NaCl 50 mM Tris-HCl 10 mM MgCl<sub>2</sub> 1 mM DTT (pH 7.9 @ 25°C) NaCI (or other salt) provides the correct ionic strength Tris-HCI provides the proper pH

- Mg<sup>2+</sup> is an enzyme co-factor
- DDT is a reducing agent

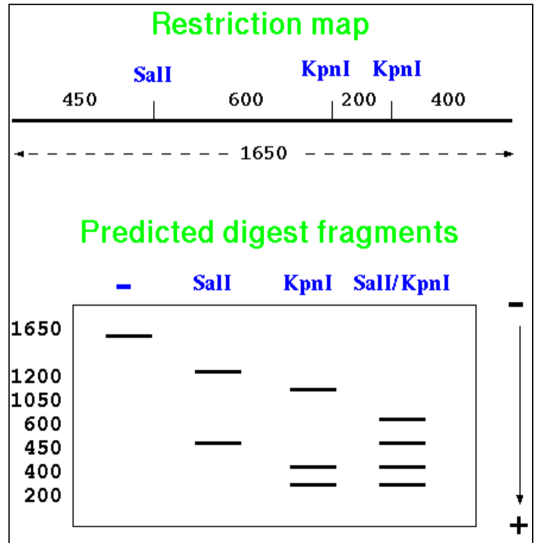
Each enzyme has defined buffer composition

DTT commonly is used as redox reagent to prevent formation of disulfide bonds in cysteine-containing proteins. Such proteins require proper formation or absence disulfide bonds for exhibiting of specific activity. DTT helps to keep cysteine-containing proteins in active state. However, if protein doesn't contain cysteins, there is no need to use DTT for its activity.

## How to set up a restriction digest



# **Restriction Digest Analysis**



- Length=1650bp
- Sall yields two fragments (1200bp and 450bp)
- *KpnI* cuts at 2 sites giving 3 fragments
- *Sall* and *Kpnl* cut 3X yielding 4 fragments