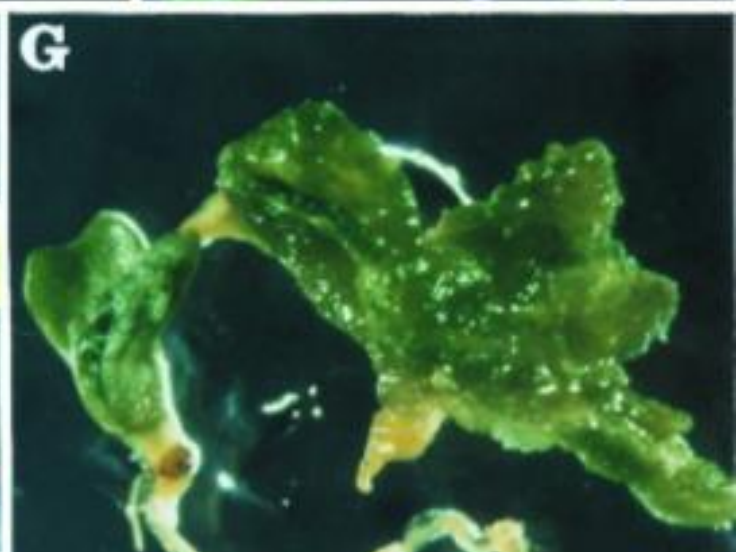
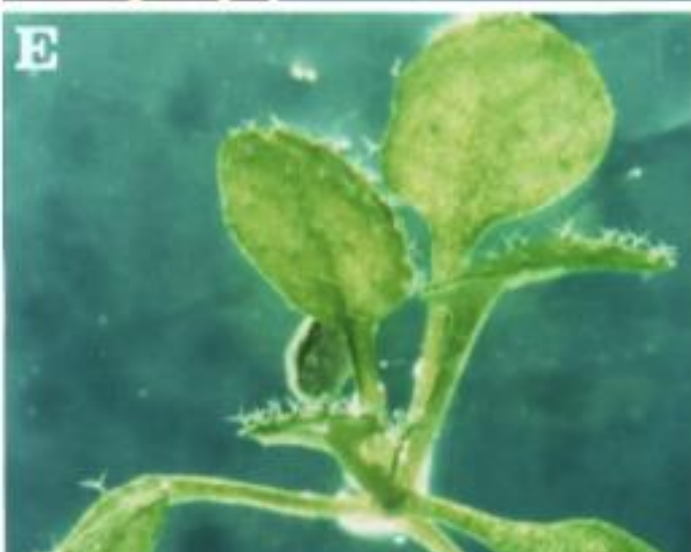




**Forward  
& Reverse Genetics**



# Strategie molecolari per studiare la funzione di un gene

↑ Aumento espressione del gene endogeno:

- forte costitutiva
- ectopica
- inducibile

↓ Riduzione/silenziamento del gene endogeno:  
• AntiSENDO

- RNAi
- co-soppressione

X Knock Out del gene.....

- Genetica Forward
- Genetica Reverse

# **Knock Out del gene.....**

**mutagenesi sito-specifica**

**v/s**

**mutagenesi inserzionale “random”**

**In piante, Drosophila, mammifero**

**bassa frequenza di ricombinazione omologa**

**Quindi.....mutagenesi random**

**O.....Genome editing: Talen, Crisp/Cas9**

# GENETICA FORWARD E REVERSE

**Obiettivo: assegnare una funzione ad ogni gene**

## **Forward Genetics:**

il metodo classico per determinare la funzione di un gene si basa sull'isolamento e analisi di mutazioni in quel gene: mutagenesi sistematica delle sequenze geniche per produrre collezioni di **mutazioni (prevalentemente recessive) di tipo “loss of function”**. Utilizzando varie tecniche di clonaggio si risale alla sequenza genica.

## **Reverse Genetics:**

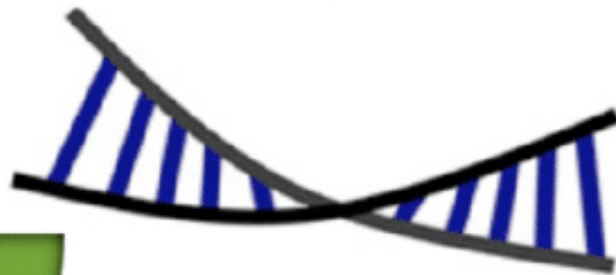
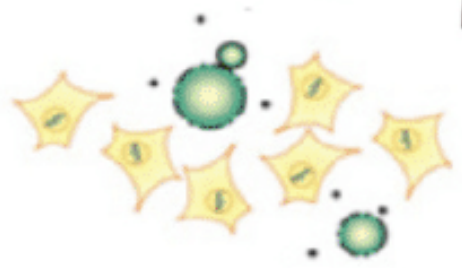
La disponibilità della sequenza del genoma di molte specie ha reso possibile un diverso approccio per la determinazione della funzione di un gene. A partire da una sequenza genica è possibile isolare la linea mutante in cui l'elemento inserzionale è all'interno della sequenza d'interesse. Dallo studio fenotipico del mutante, si può dedurre la funzione del gene.

Forward Genetics

Mutagenesis,  
QTL mapping,  
Positional  
cloning, etc...

Discover  
Gene  
underlying  
Phenotype

Known Phenotype



Known Gene

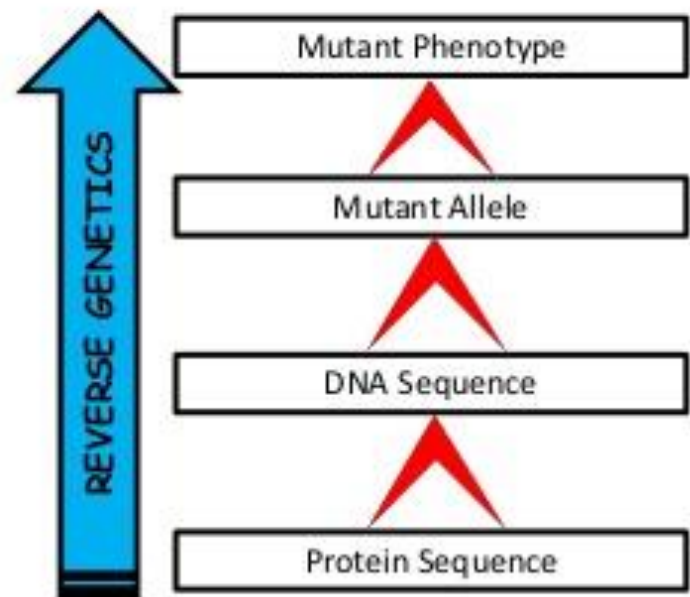
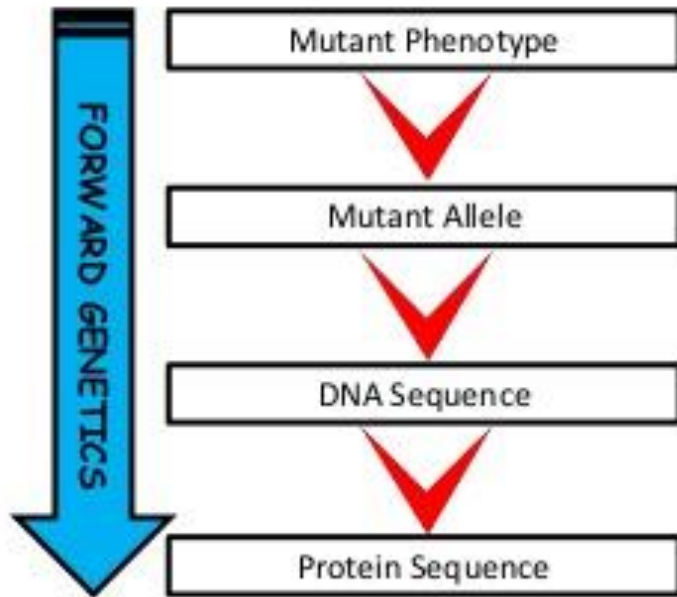
Ectopic expression,  
Gene silencing,  
Gene targeting,  
TILLING, etc...

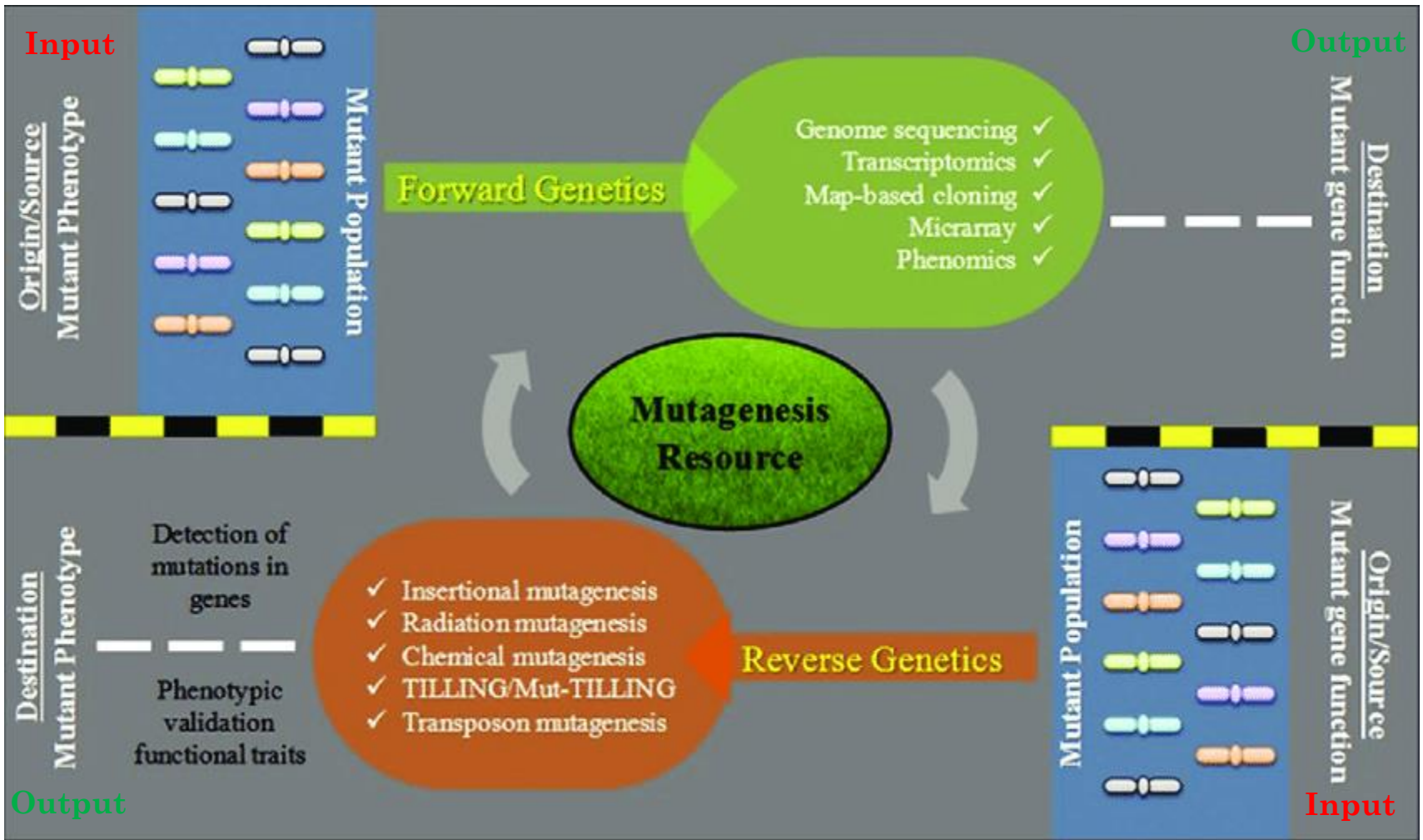
Phenotype  
Resulting  
from  
Alteration

Reverse Genetics



# Forward Vs Reverse Genetics





## GENES AND PHENOTYPES

**Gene:** a functional unit of inheritance, usually corresponding to the segment of DNA coding for a single protein.

**Genome:** an organism's set of genes.

**locus:** the site of the gene in the genome



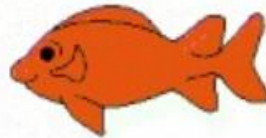
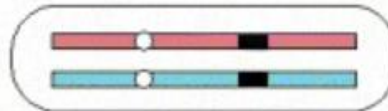
**alleles:** alternative forms of a gene



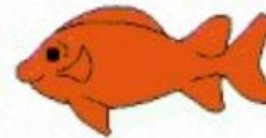
**GENOTYPE:** the specific set of alleles forming the genome of an individual

**PHENOTYPE:** the visible character of the individual

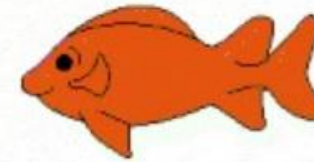
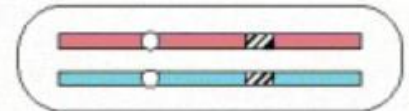
homozygous A/A



heterozygous a/A



homozygous a/a



**Wild-type:** the normal, naturally occurring type



**Mutant:** differing from the wild-type because of a genetic change (a mutation)

allele A is **dominant** (relative to a); allele a is **recessive** (relative to A)

In the example above, the phenotype of the heterozygote is the same as that of one of the homozygotes; in cases where it is different from both, the two alleles are said to be co-dominant.

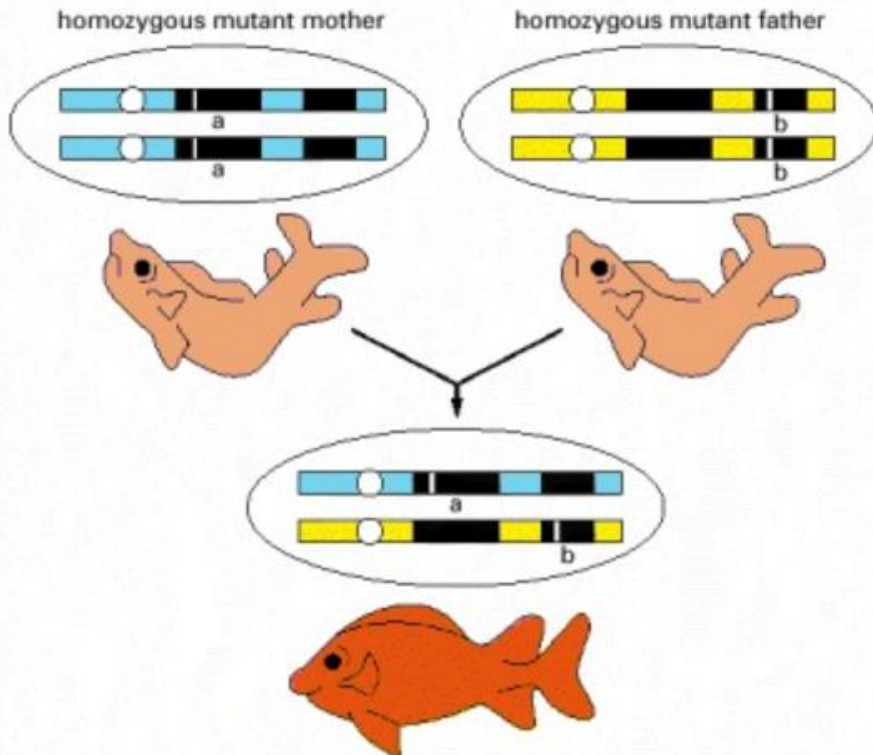


# Test di Complementazione tra alleli mutanti

## TWO GENES OR ONE?

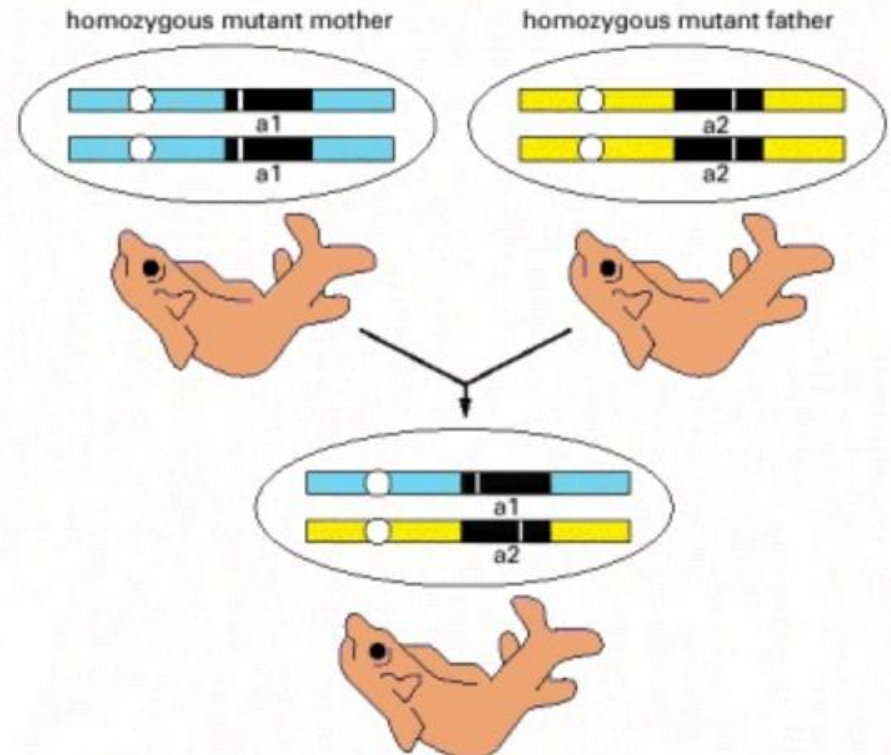
Given two mutations that produce the same phenotype, how can we tell whether they are mutations in the same gene? If the mutations are recessive (as they most often are), the answer can be found by a **complementation test**.

### COMPLEMENTATION: MUTATIONS IN TWO DIFFERENT GENES



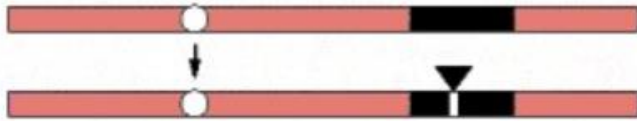
In the simplest type of complementation test, an individual who is homozygous for one mutation is mated with an individual who is homozygous for the other. The phenotype of the offspring gives the answer to the question.

### NONCOMPLEMENTATION: TWO INDEPENDENT MUTATIONS IN THE SAME GENE

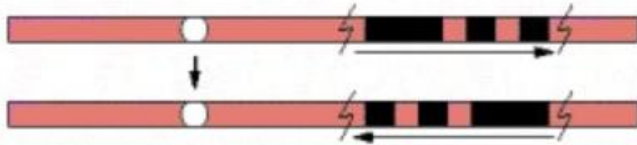


# Loss of Function vs Gain of Function mutations

## TYPES OF MUTATIONS



**POINT MUTATION:** maps to a single site in the genome, corresponding to a single nucleotide pair or a very small part of a single gene



**INVERSION:** inverts a segment of a chromosome

**lethal mutation:** causes the developing organism to die prematurely.

**conditional mutation:** produces its phenotypic effect only under certain conditions, called the *restrictive* conditions. Under other conditions—the *permissive* conditions—the effect is not seen. For a *temperature-sensitive* mutation, the restrictive condition typically is high temperature, while the permissive condition is low temperature.

**loss-of-function mutation:** either reduces or abolishes the activity of the gene. These are the commonest class of mutations. Loss-of-function mutations are usually *recessive*—the organism can usually function normally as long as it retains at least one normal copy of the affected gene.

**null mutation:** a loss-of-function mutation that completely abolishes the activity of the gene.



**DELETION:** deletes a segment of a chromosome



**TRANSLOCATION:** breaks off a segment from one chromosome and attaches it to another

**gain-of-function mutation:** increases the activity of the gene or makes it active in inappropriate circumstances; these mutations are usually *dominant*.

**dominant negative mutation:** dominant-acting mutation that blocks gene activity, causing a loss-of-function phenotype even in the presence of a normal copy of the gene. This phenomenon occurs when the mutant gene product interferes with the function of the normal gene product.

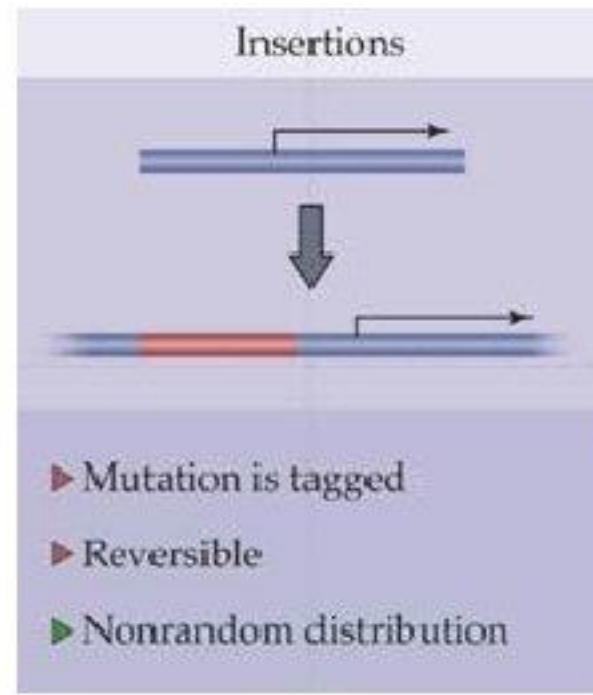
**suppressor mutation:** suppresses the phenotypic effect of another mutation, so that the double mutant seems normal.

An *intragenic* suppressor mutation lies within the gene affected by the first mutation; an *extragenic* suppressor mutation lies in a second gene—often one whose product interacts directly with the product of the first.

# Mutagenesi inserzionale mediata da elementi trasponibili

Facile mappatura dell'inserzione

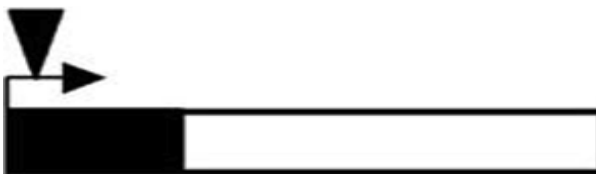
Rapido clonaggio del gene mutato





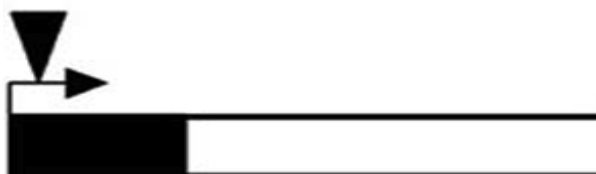
**Posizione**  
ORF o promotore

**Risultato**  
KO



Promotore o  
3' UTR

espressione  
ridotta



Promotore

espressione  
aumentata



ORF

non nullo



ORF o promotore  
multipli

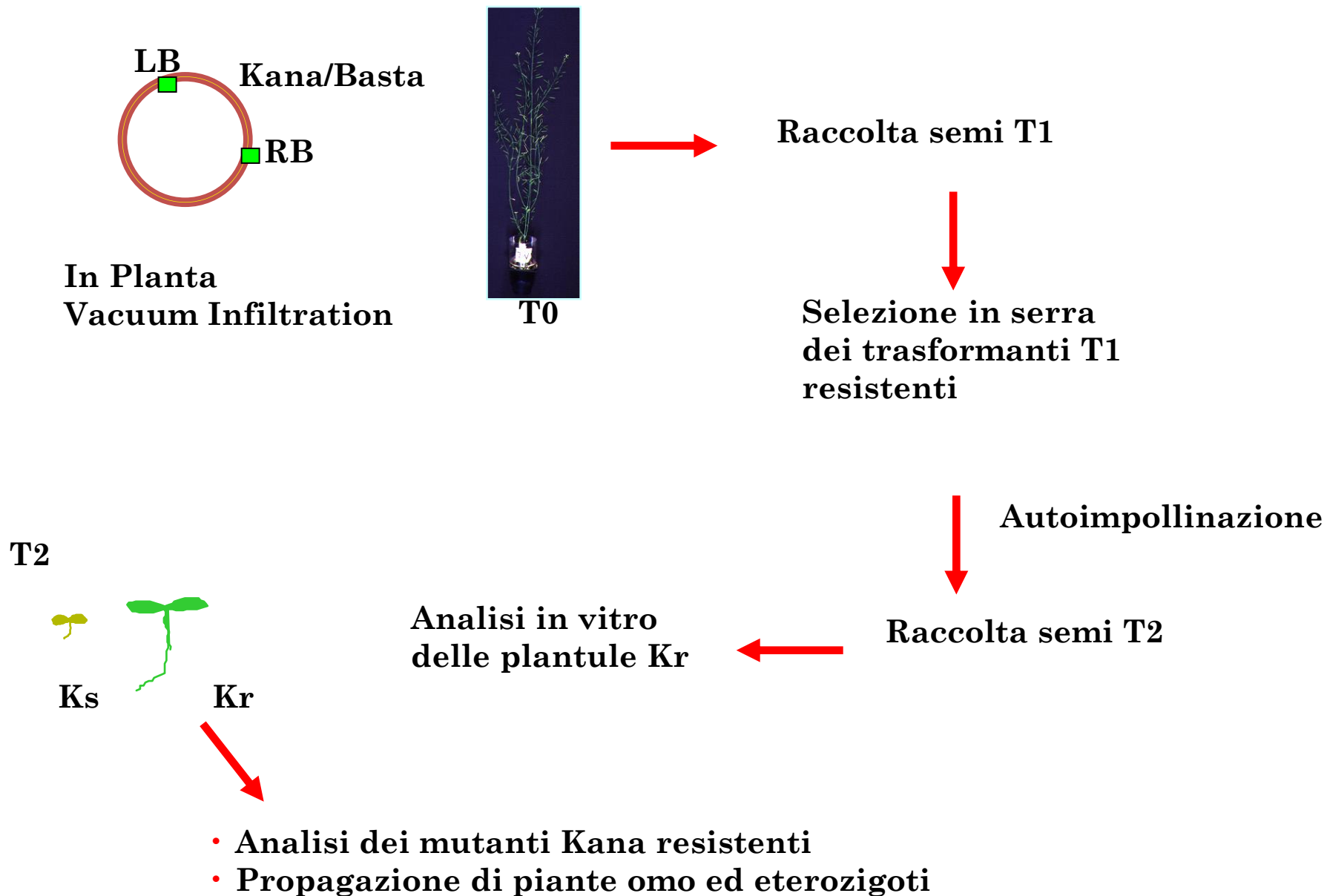
KO



ORF o  
promotore

Riarrangiamento  
cromosomico

# Strategia di Isolamento Mutanti Inserzionali





**Ma il T-DNA è correlato  
(responsabile del) con il fenotipo mutato???**

- **Analisi di segregazione  
del marcatore e eventualmente del fenotipo mutante  
(inserzione T-DNA in 1 o più loci??)**
- **Caratterizzazione molecolare del sito di inserzione  
(T-DNA singolo, a tandem oppure inserzione più complessa??)**
- **Clonaggio delle “Flanking Sequences”**

# Analisi di segregazione

Fenotipo «selvatico»				
<u>T2 Plants</u>	aa(Kr/Kr)	Aa(Kr/Ks)	AA(Ks/Ks)	Total
	96	193	94	383
	1	2	1	

autoimpollinazione

<u>T3 Plants</u>	aa(Kr)	Aa(Kr/Ks)	Ks	Total
a	24	47	26	97
b	25	46	24	95
c	27	55	28	110
e	121	0	0	121

aa gene inattivo in omozigosi

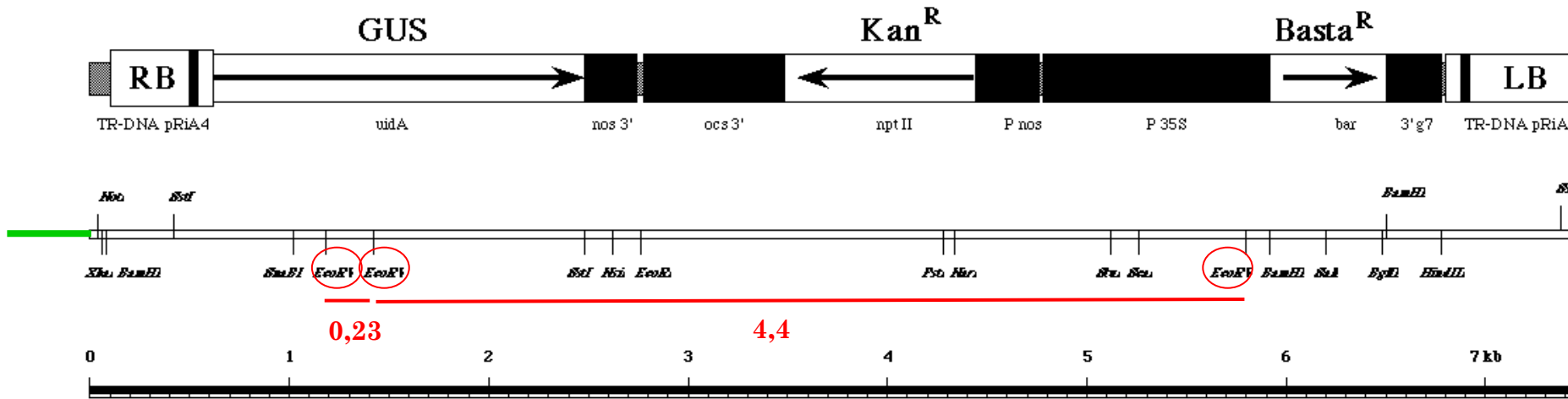
Aa una copia gene inattiva, una selvatica

**Ma il T-DNA è correlato  
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- **Analisi di segregazione  
del marcatore e del fenotipo mutante  
(inserzione T-DNA in 1 o più loci??)**
- **Caratterizzazione molecolare del sito di inserzione  
(T-DNA singolo, a tandem oppure inserzione più complessa??)**
- **Clonaggio delle “Flanking Sequences”**

pGKB5 T-DNA

X



probe

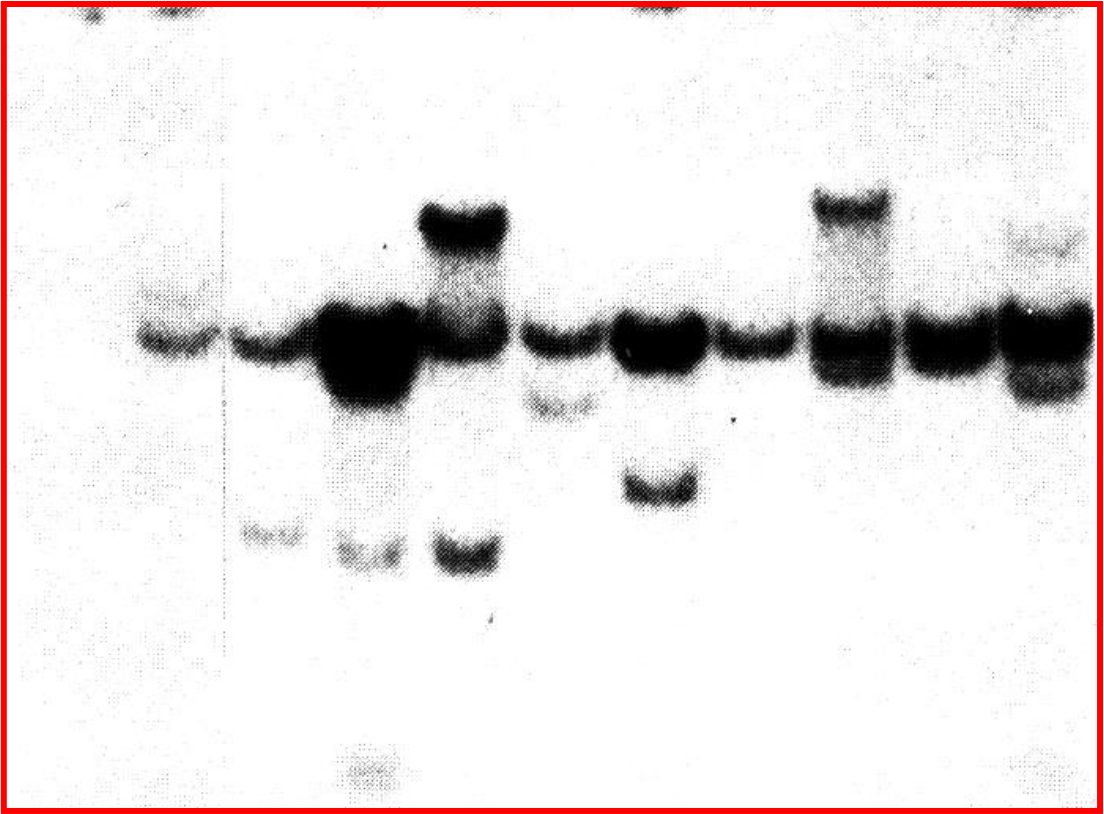
**Digestione EcoRV:**

**0.23 Kb e 4,4 Kb interne al T-DNA**

**X Kb fino al sito EcoRV sulla flanking sequence**

**Caratterizzazione  
molecolare  
del sito di  
inserzione**

**NT**



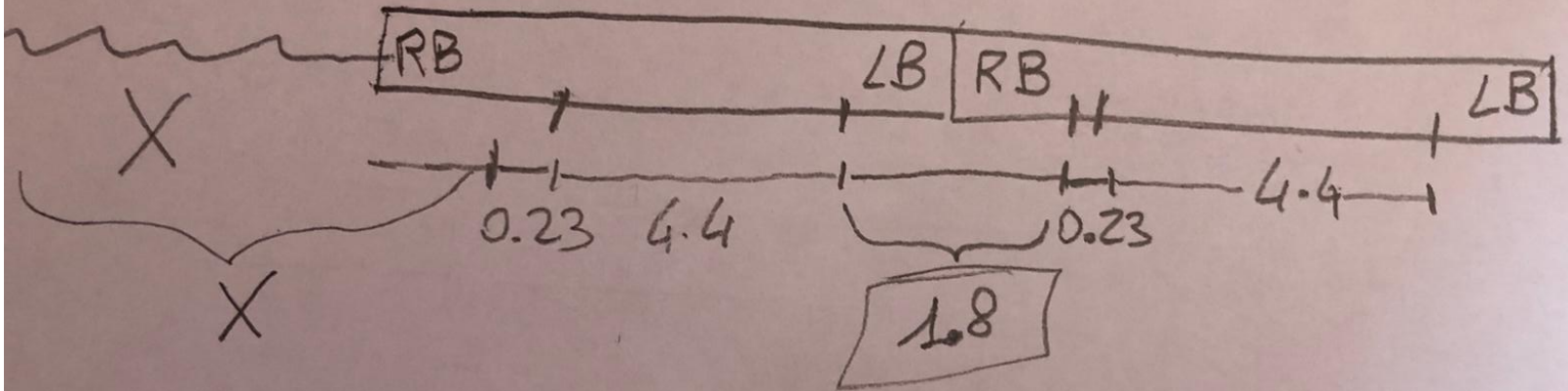
← 4,4 Kb

← 1,8 Kb  
RB-LB IR

0,23 Kb →



$F_s$



Probe

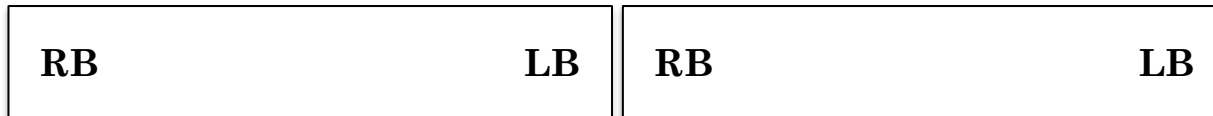


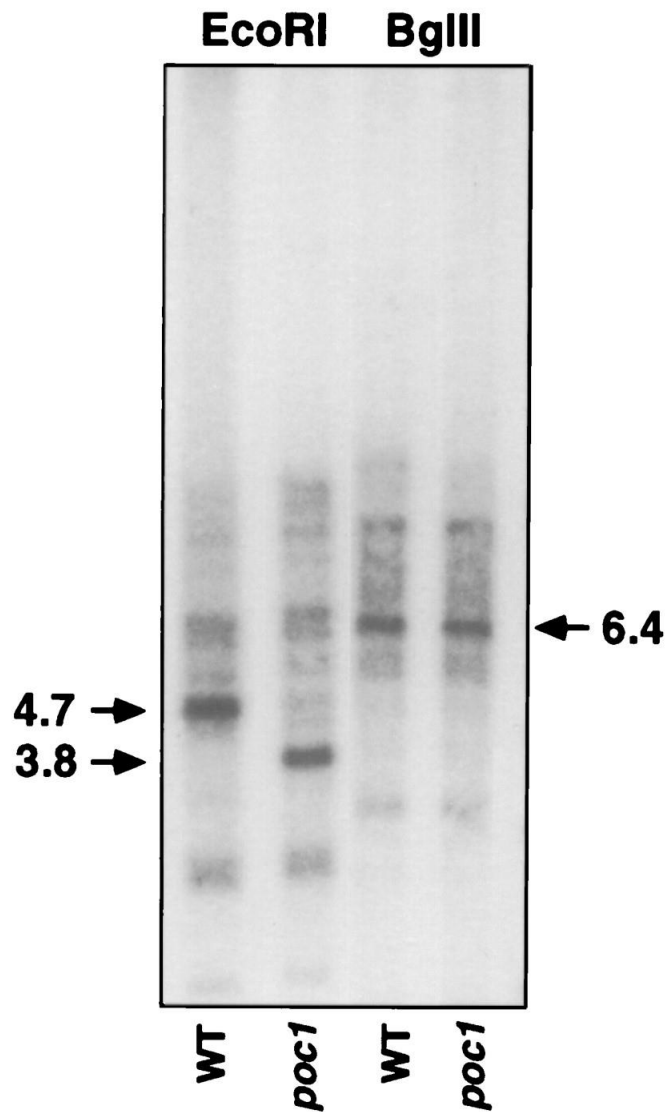
Probe

## Coda-Coda



## Testa-Coda

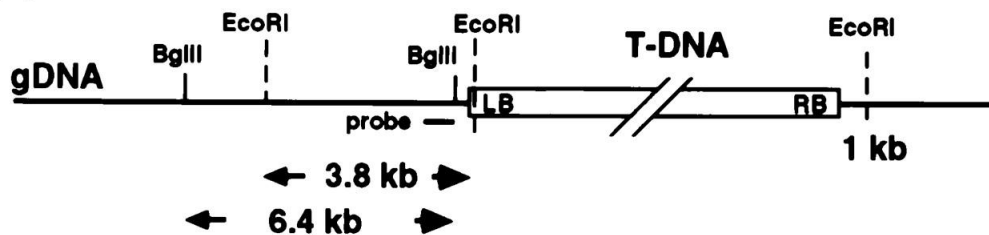


**A**

Identification of the genomic region flanking the T-DNA inserted in *poc1*.

(A) Southern blot analysis of total genomic DNA from wild type (WT) and *poc1* digested with *EcoRI* or *BglII* and hybridized with a **0.5-kb probe from the cloned region flanking the T-DNA left border (LB)**.

(B) Diagram of the T-DNA insert within the *poc1* genomic DNA (gDNA) illustrating the locations of the restriction sites, the size of the restriction fragments generated, and the fragment used as the probe.

**B**

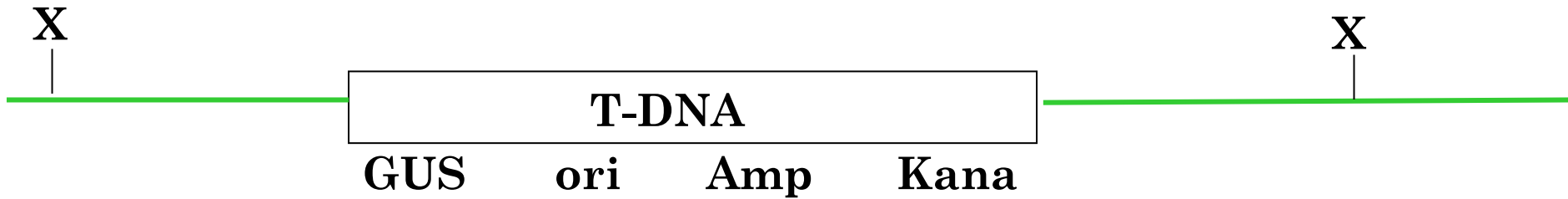
**Ma il T-DNA è correlato  
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- **Analisi di segregazione  
del marcatore e del fenotipo mutante  
(inserzione T-DNA in 1 o più loci??)**
- **Caratterizzazione molecolare del sito di inserzione  
(T-DNA singolo, a tandem oppure inserzione più complessa??)**
- **Clonaggio delle “Flanking Sequences”**

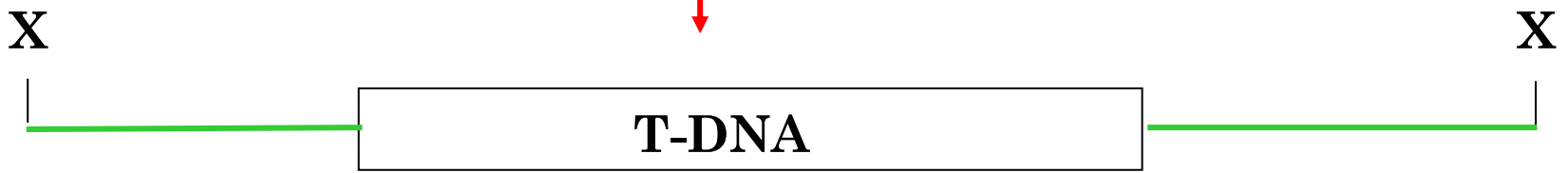
# Strategie di clonaggio delle “Flanking Sequences”

- **Plasmid Rescue**
- **Inverse PCR (IPCR)**
- **Libreria genomica del mutante  
(probe T-DNA)**

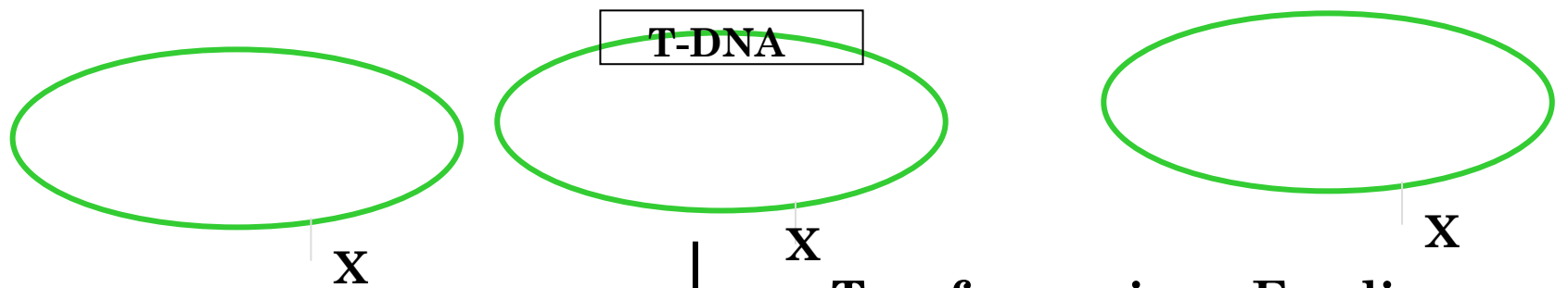




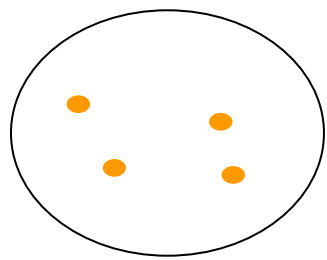
**Digestione**



**Ligazione intramolecolare**



**Trasformazione E.coli**



**Selezione colonie AmpR, su LB+Amp**

# Strategie di clonaggio delle “Flanking Sequences”

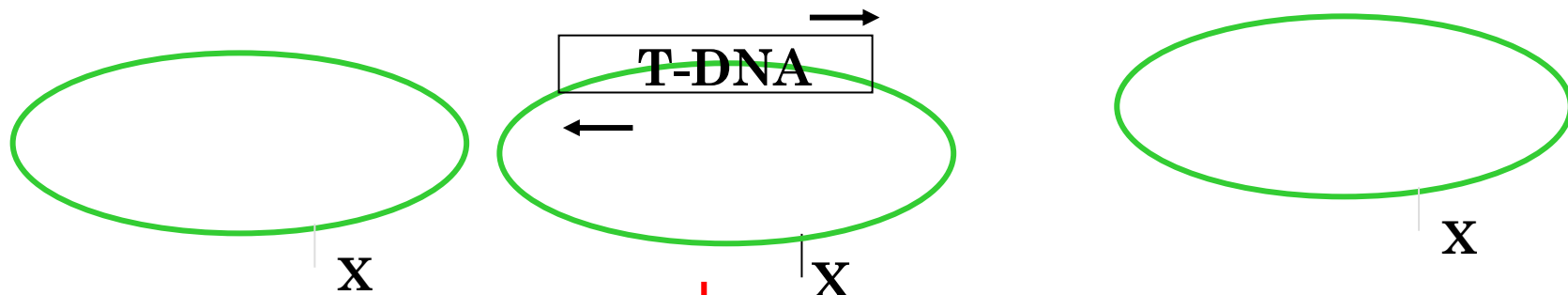
- **Plasmid Rescue**
- **Inverse PCR (IPCR)**
- **Libreria genomica del mutante  
(probe T-DNA)**



**Digestione**



**Ligazione intramolecolare**



**PCR**



# Gene Targeting in Mice/hES cells

**Il Gene Targeting è mediato da Ricombinazione omologa**

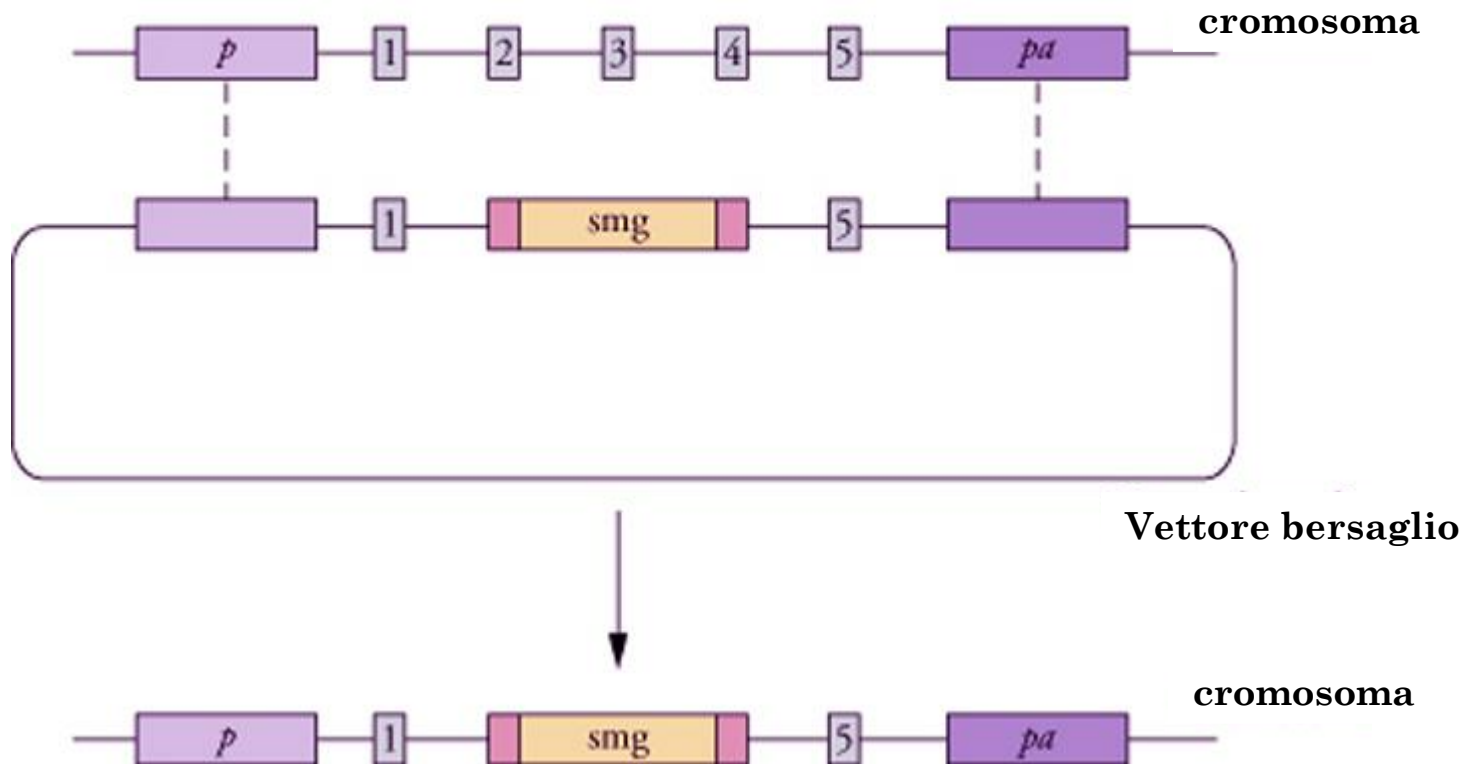
Nelle cellule di mammifero la ricombinazione omologa è un evento  
**molto raro** (a differenza del lievito)

La frequenza di questo evento aumenta se il grado di omologia di sequenza tra il DNA introdotto (esogeno) e il gene bersaglio (endogeno) è molto elevata

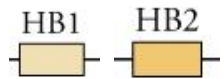
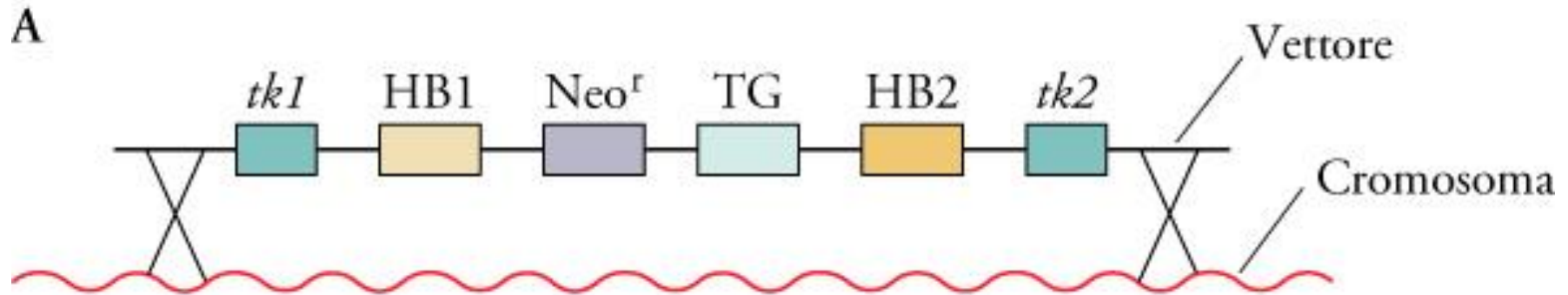
Per questo il clone di DNA esogeno è una sequenza isogenica  
(derivante dallo stesso ceppo murino)

## Gene targeting mediante ricombinazione omologa

Comporta l'introduzione di una mutazione in un gene specifico e può essere considerata come una forma di "site-directed in vivo mutagenesis" e quindi è utilizzata per lo studio della funzione genica



# SELEZIONE POSITIVA-NEGATIVA cellule ES



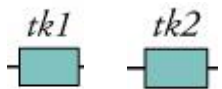
Seq di DNA per RICOMBINAZIONE OMOLOGA nel genoma bersaglio



TRANSGENE di interesse

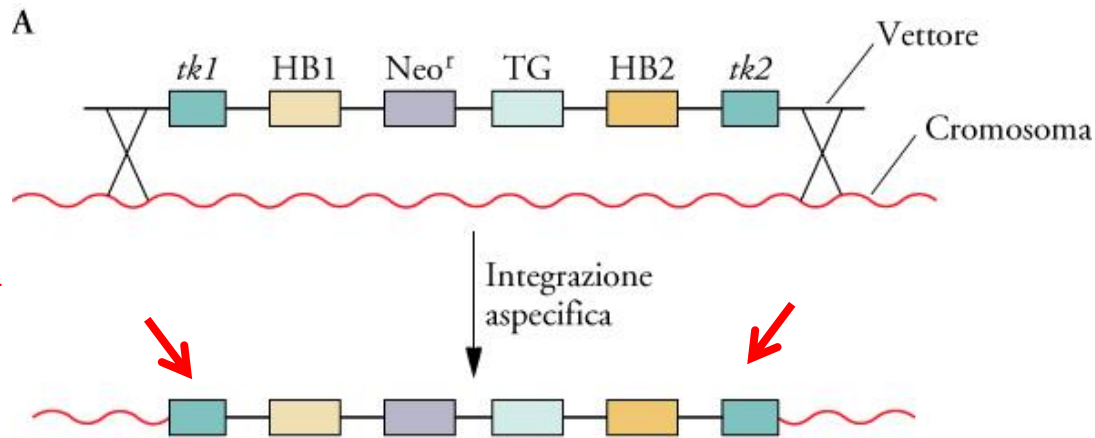


Gene che conferisce resistenza alla G418

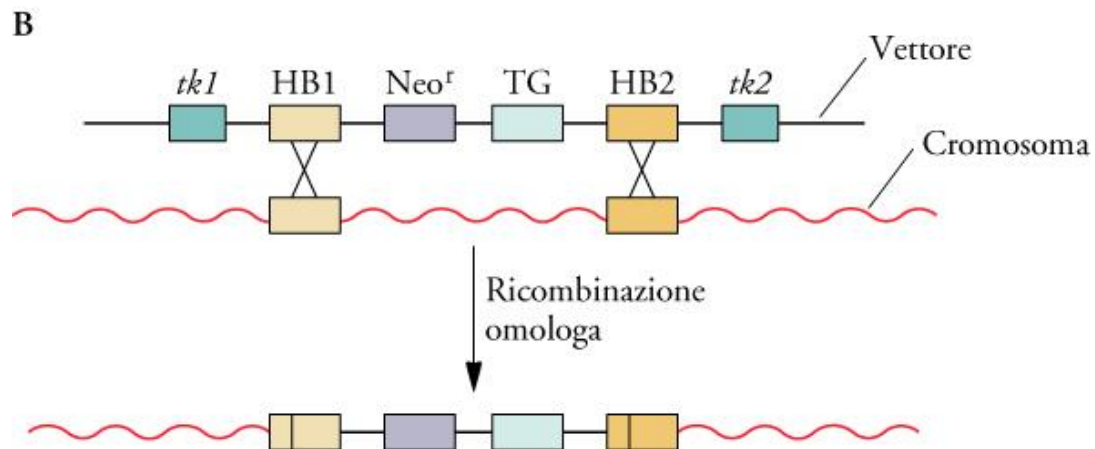


Geni della Timidina Kinasi (virus herpes simplex). In presenza di GANCICLOVIR: MORTE

# SELEZIONE POSITIVA e NEGATIVA



ASPECIFICA



SPECIFICA

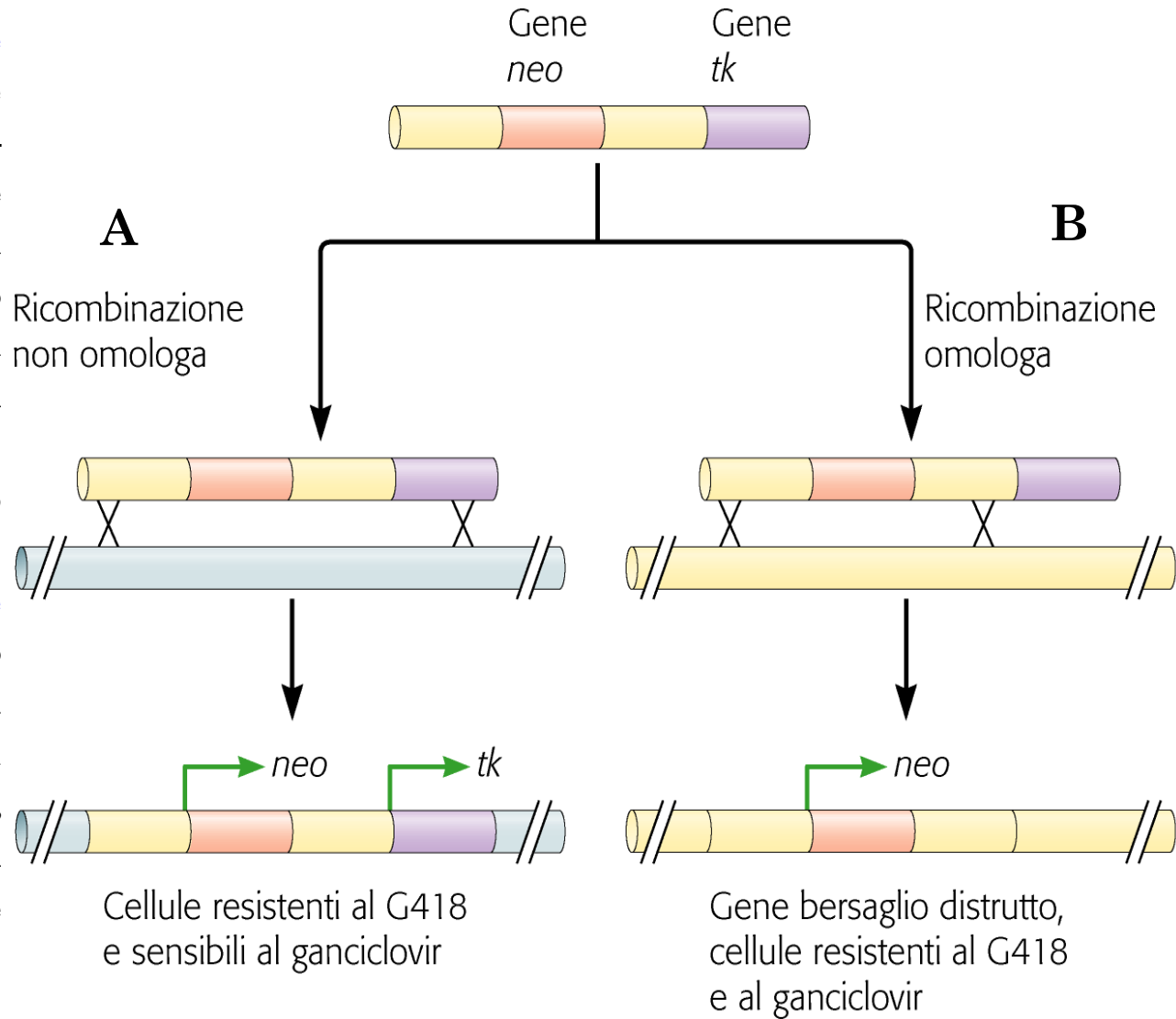
- 1) Aggiunta di **G418**: SELEZIONE POSITIVA di cellule che hanno INTEGRATO il DNA
- 2) Aggiunta di **GANCICLOVIR**: SELEZIONE NEGATIVA di cellule con INTEGRAZIONE ASPECIFICA



## La selezione positiva/negativa

**A: risultato di integrazione aspecifica.** Si selezionano le cellule positivamente per la resistenza a G-418, conferita dal gene Neo, e negativamente in presenza del gancyclovir, che viene trasformato dalla timidina chinasi in composti tossici. Le cellule contenenti il vettore inserito random (A) conterranno il gene TK, e saranno uccise in presenza di gancyclovir

**B: risultato di ricombinazione omologa.** Il prodotto del doppio crossover tra i blocchi omologhi HB1 e HB2 non contiene i due geni della timidina chinasi (tk1 e tk2), quindi sono resistenti al gancyclovir, e anche al G-418 grazie alla presenza del gene Neo.

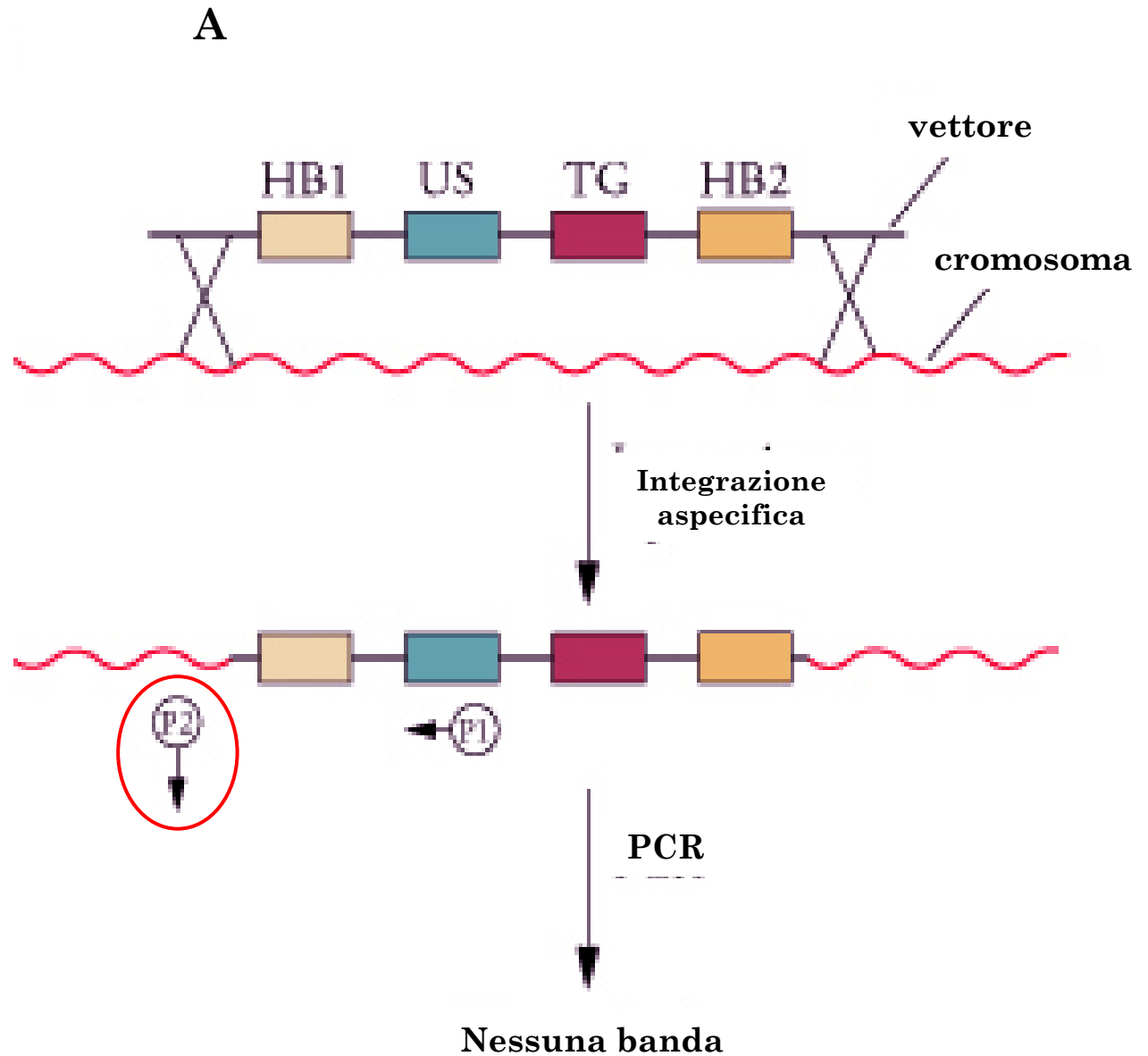


**Analisi dell'integrazione mediante PCR**

**A: integrazione aspecifica**

Uno dei primers (P2) non è in grado di appaiarsi al sito cromosomico P2. P1 invece si ibrida ad una sequenza US che si trova sul DNA esogeno.

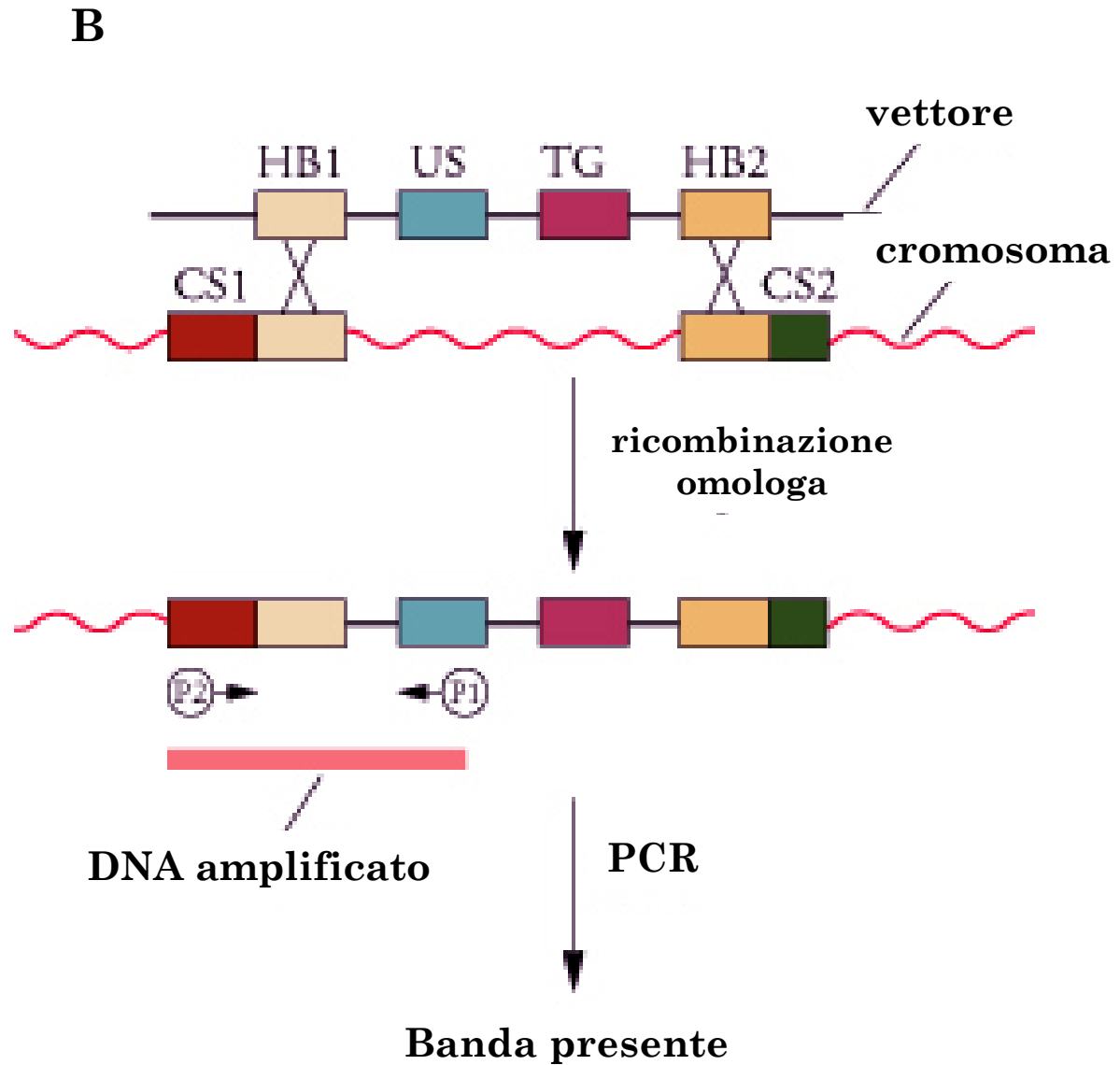
Quindi non si ha amplificazione



## Analisi dell' integrazione mediante PCR

B: ricombinazione omologa

La ricombinazione omologa tra le sequenze di DNA HB1 e HB2 del DNA esogeno che sono complementari ai siti cromosomici CS1 e CS2, crea regioni di ibridazione per P1 e P2 che si trovano ad una distanza predeterminata. L' amplificazione così genera il prodotto di dimensioni attese.



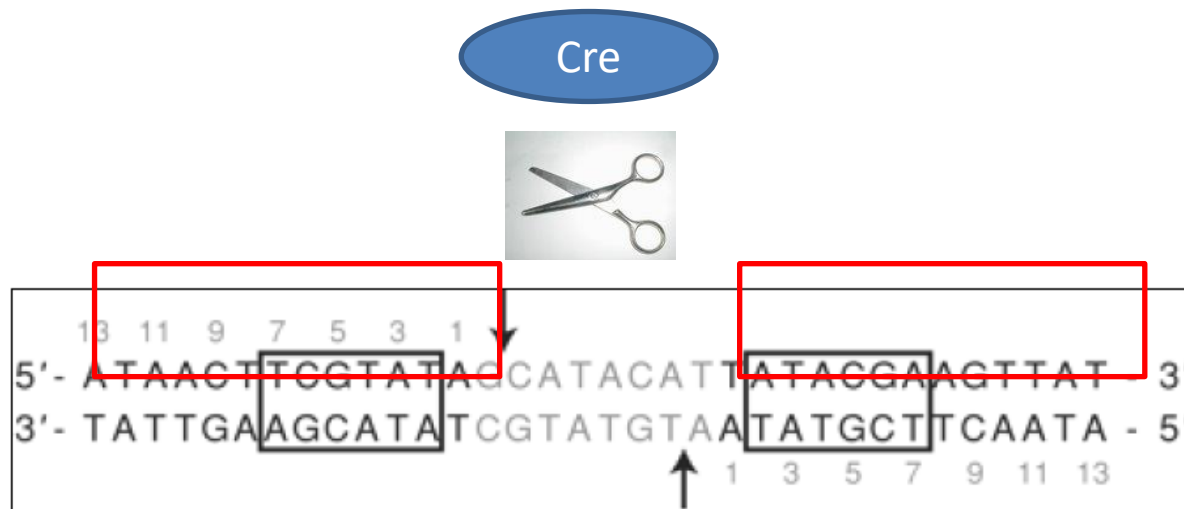
# Cre-Lox System

Il sistema Cre/Lox deriva dal fago P1

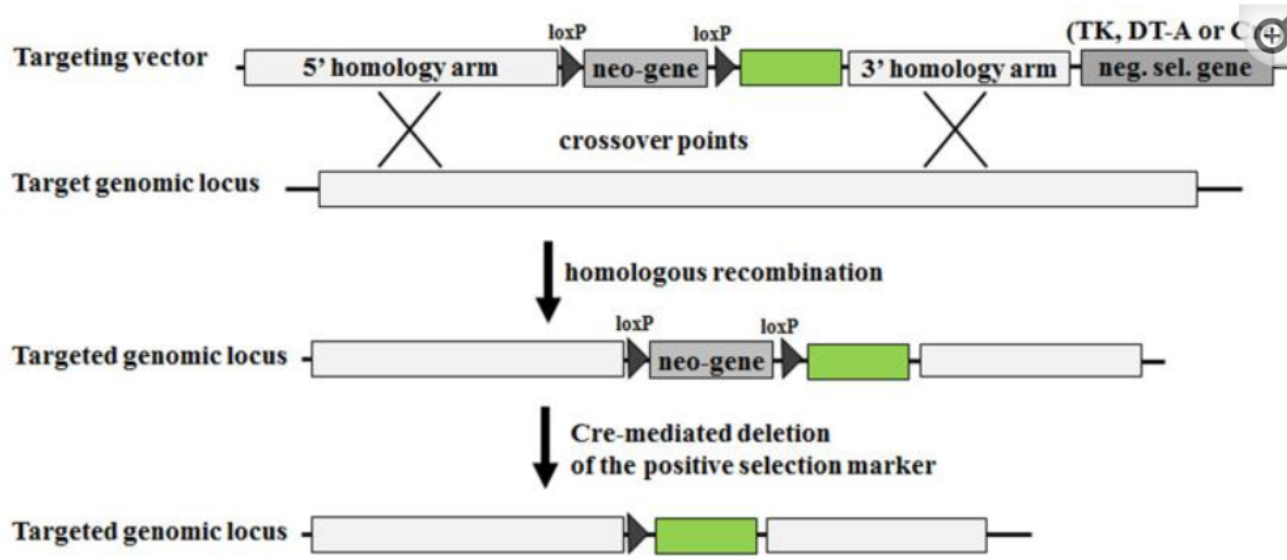
Il sistema è detto “floxing” in quanto consiste di sequenze fiancheggianti il gene e la ricombinazione mediata da Cre

**Cre:** Ricombinasi sequenza-specifica (catalizza la ricombinazione tra siti Lox)

**LoxP:** Locus di crossover (2 palindromi 13bp + regione centrale di 8nt)



la ricombinasi CRE riconosce i siti LoxP e ricombina le sequenze di DNA adiacenti

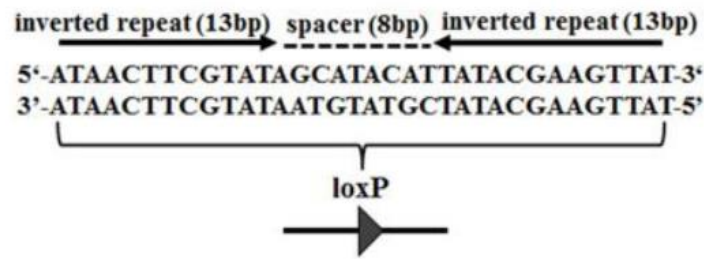


Typical gene targeting strategy

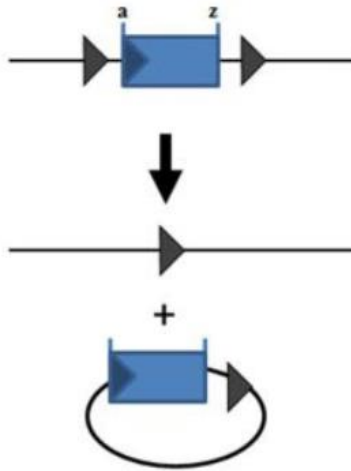
## Typical gene targeting strategy

A targeting vector is composed of three basic units: (i) a 5' homology arm; (ii) a gene marker for positive selection (e.g. neomycin resistance gene (neo)); (iii) a 3' homology arm; and (iv) a negative selection marker (neg. sel. Marker), such as thymidine kinase, diphtheria toxin fragment A (DT-A), or, if the positive selection marker is flanked by loxP sites, Cre recombinase gene (Cre). Furthermore, any desired DNA sequence of interest (here green box) can be inserted between the homology arms of the targeting vector, in order to introduce it into the target genome by homologous recombination. Homologous recombination between the targeting vector and the target cognate chromosomal region results in the disruption of one genomic copy of the targeted genomic locus and loss of the vector's negative selection marker gene. Crossover points are depicted by "X". The floxed (loxP sites flanked) positive selection marker gene can be removed by expressing Cre recombinase in the recombinant ESCs or by crossing the chimeric mice with Cre-expressing transgenic mice

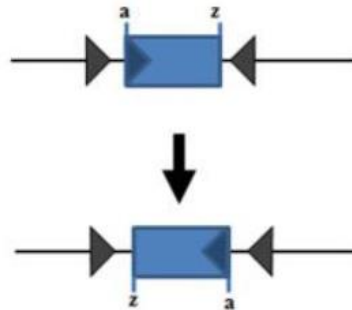
A)



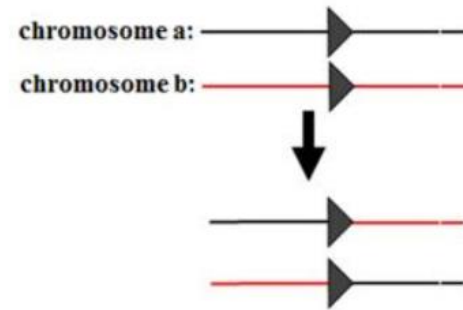
B) Cre-mediated deletion



C) Cre-mediated inversion

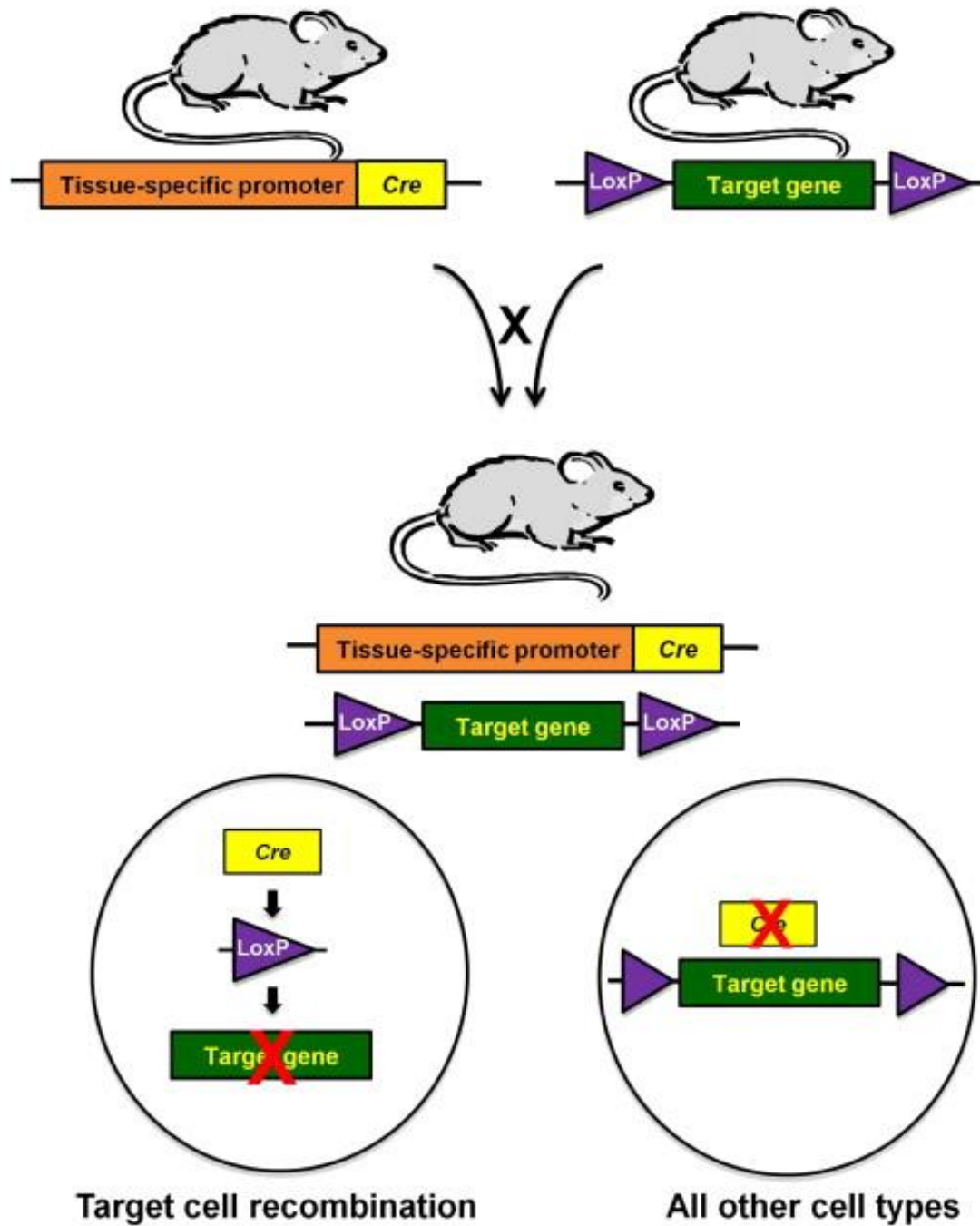


D) Cre-mediated translocation



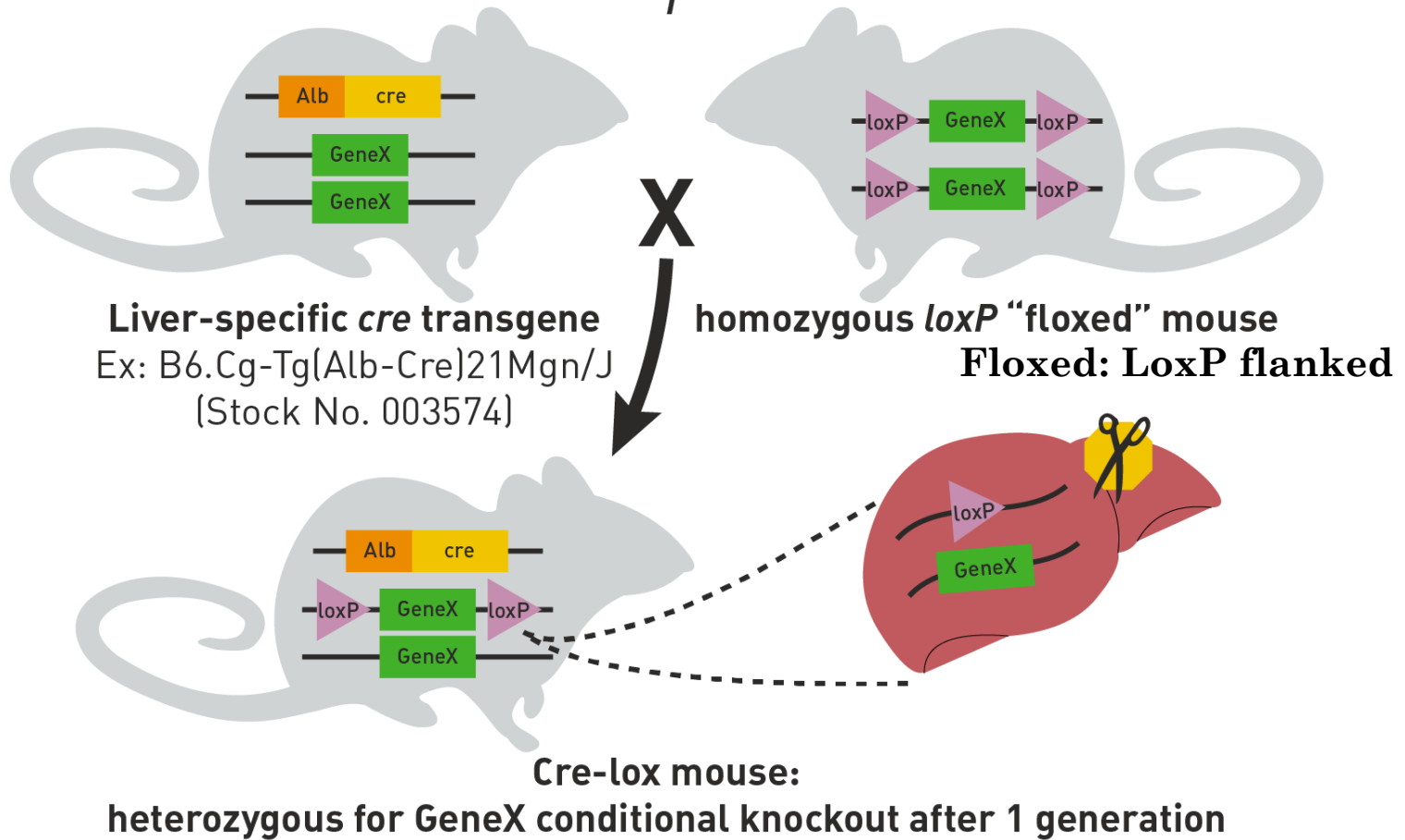
## LoxP structure and Cre recombinase-mediated recombinations

Single loxP site that contains two inverted 13 bp repeats, separated by an asymmetric 8 bp long sequence. The type of Cre-mediated recombination is dependent on the orientation and location of the loxP sites: (B) Cre excises a circular molecule from between two loxP sites placed in the same orientation; (C) Cre inverts the DNA sequence between two loxP sites positioned in opposite orientation; (D) Cre-mediated recombination between two different linear DNA molecules (e.g. chromosomes), each containing a loxP site, resulting in the exchange of the DNA regions flanking the loxP sites.

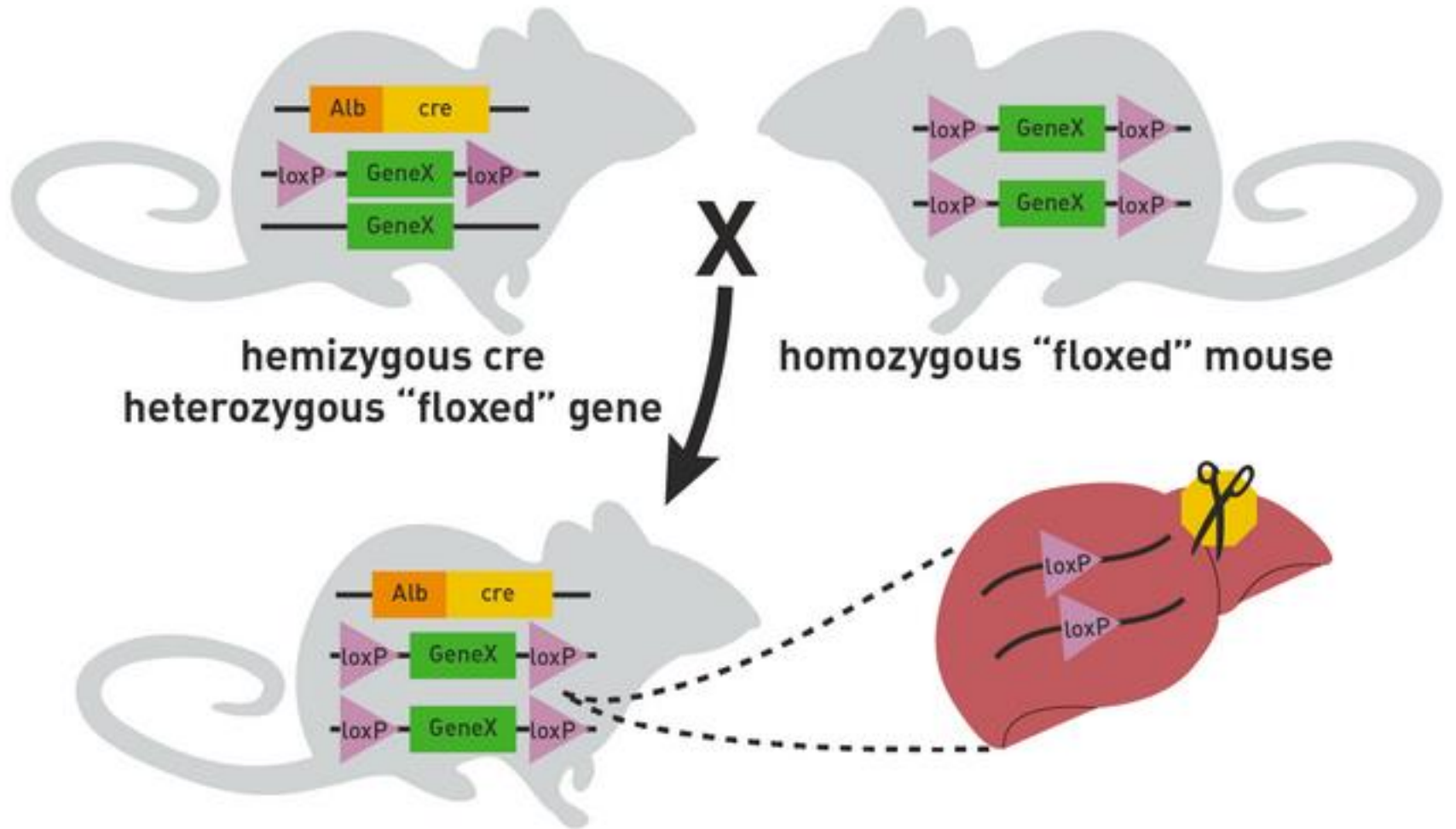




# Cre-lox *Tissue-Specific* Knockout

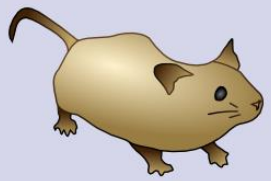


Approximately 50% of the offspring will be heterozygous for the loxP allele and heterozygous for the cre transgene. Mate these mice back to the homozygous *loxP*-flanked mice. Approximately 25% of the progeny from this mating will be homozygous for the *loxP*-flanked allele and heterozygous for the *cre* transgene. These will be your experimental mice.

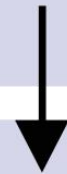
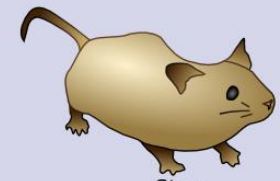


F<sub>0</sub> Generation

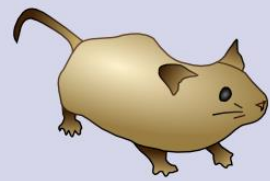
Cre Mouse



LoxP (Floxed) Mouse

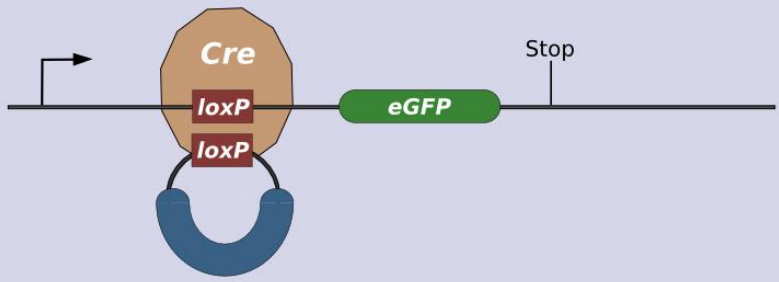


Cre LoxP Mouse



F<sub>1</sub> Generation

Cells with active Cre recombinase



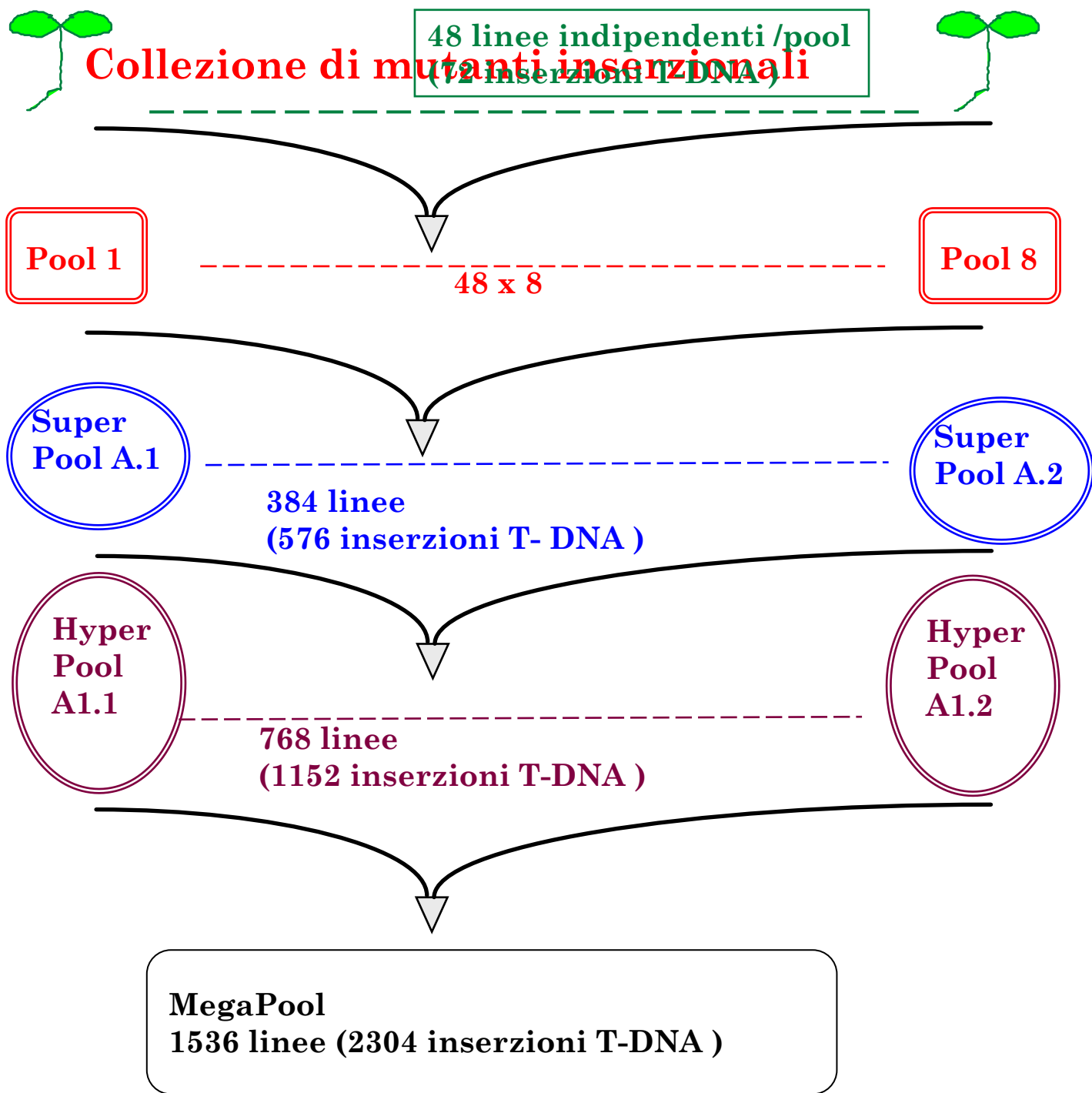
Cells lacking active Cre recombinase



**Original gene function is disrupted, a reporter gene is transcribed instead.**

**Original gene function is untouched.**

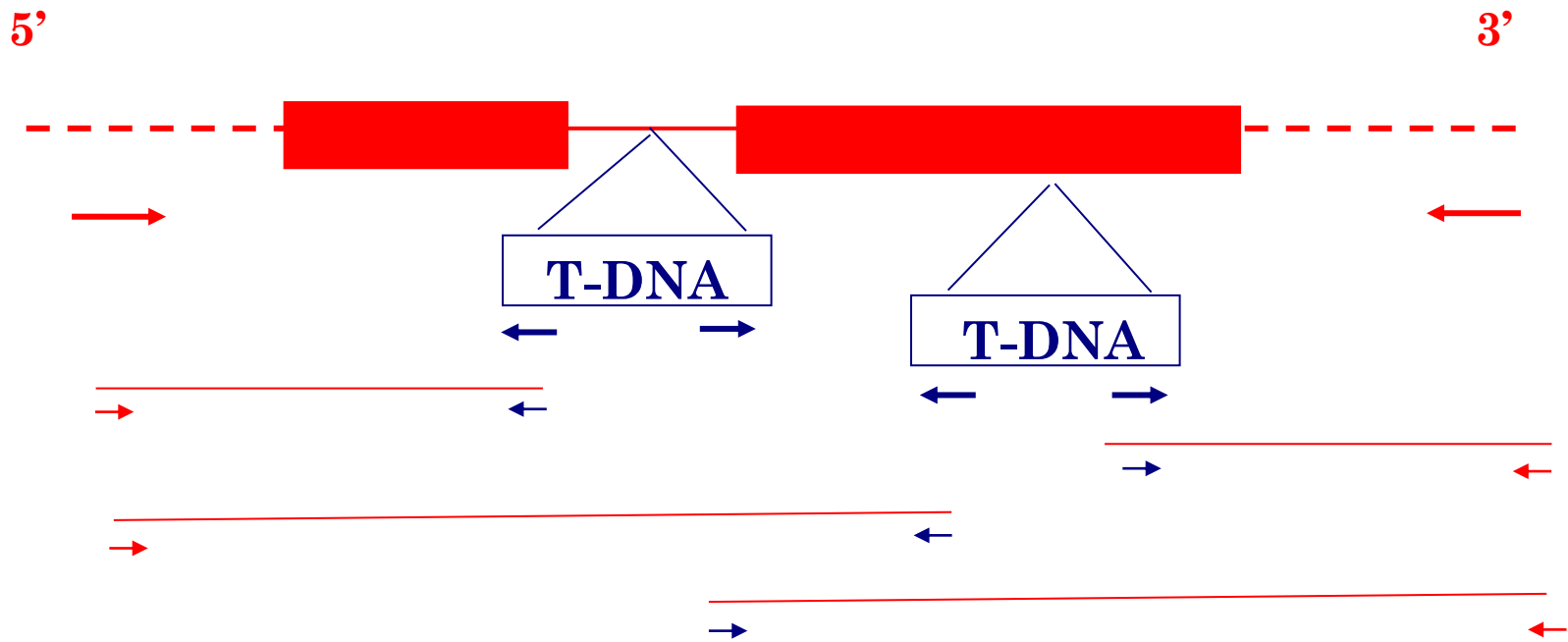
# Reverse Genetics



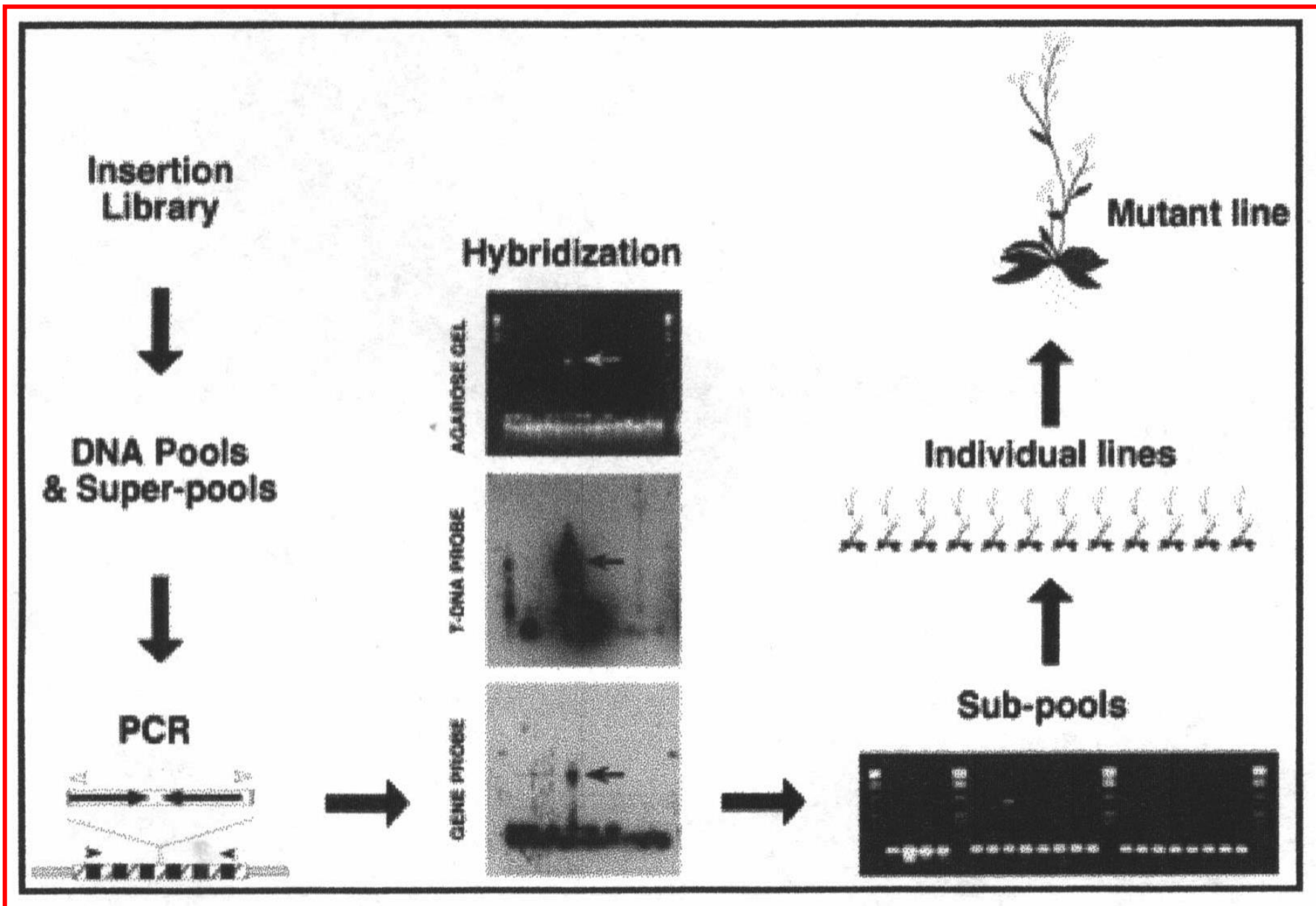
# Protocollo di Screening mediante PCR

- **Screening per PCR di n HP:**  
combinazione di un primer gene-specifico (F/R)  
ed un primer T-DNA (RB/LB)
- **Qualità dei primers:** specificità-sensibilità  
background con i primers T-DNA
- **Analisi per ibridazione (transfer “sandwich”):**  
sonda T-DNA (RB+LB)  
sonda gene-specifica
- **Segnali coincidenti con le 2 sonde:**  
PCR di conferma con primers T-DNA nested  
o con primers gene-specifici  
sequenza del frammento amplificato

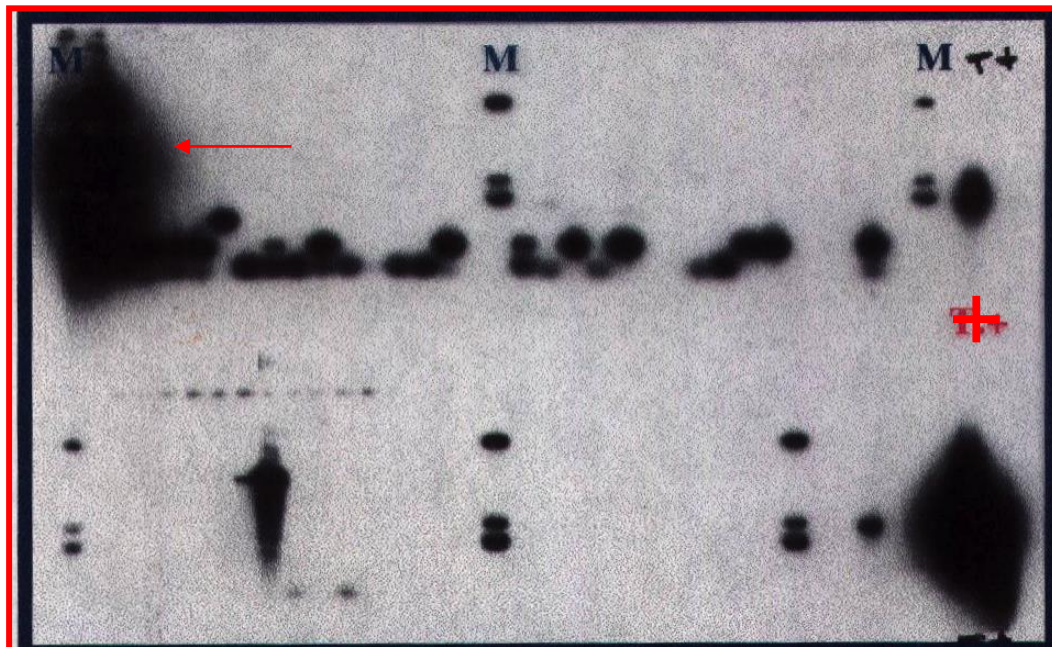
# Scelta dei Primers gene-specifici







**Sonda DAG1**



**Sonda T-DNA**

