

Chapter 8 Molecular approaches to study virulence factors

Approaches used to study new virulence factors

Traditional approaches

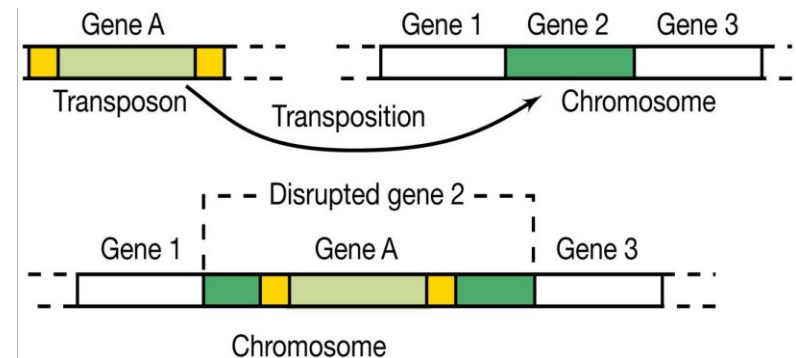
Biochemical: purification of virulence factors (diphtheria and cholera toxins)

Molecular genetics: transposon mutagenesis, reporter gene fusion

Finding genes that are expressed *in vivo*

Signature tagged mutagenesis (STM)

In vivo Expression technology (IVET)



Genomic method for identifying virulence genes

Genomic Subtractive Hybridization (GSH)

Massively parallel methods of DNA sequencing (Next-generation sequencing-NGS)

In Vivo Induced Antigen Technology (IVIAT)

Measuring Infectivity and Virulence

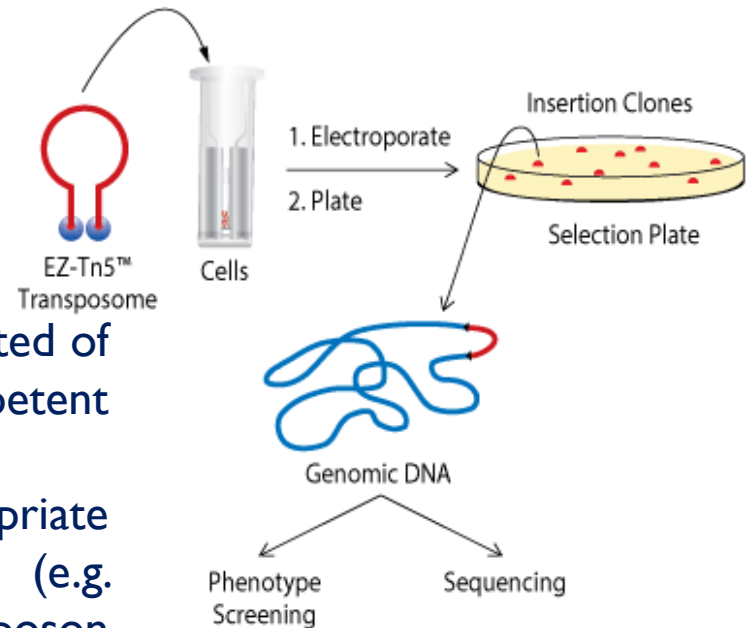
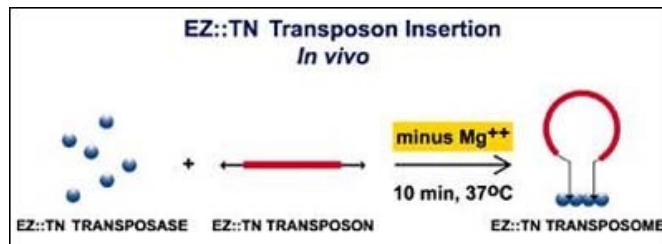
Animal models of infection

Cell and organ cultures.

Transposon mutagenesis

An approach to generate strains with attenuated virulence compared to the wt parent strain and to identify genes responsible of the attenuated virulence is that to create a library of random mutants.

Library of mutants: obtained by chemical mutagenesis, UV irradiation, **transposon insertion** and a biological assay to screen for those that still grow in the medium but with attenuated virulence.



In vivo mutagenesis: transposon of choice completed of transposase (fig.) can be electroporated into competent bacteria of choice.

Transformed bacteria are selected on the appropriate medium containing the selective marker (e.g. kanamycin). Surviving colonies contain a transposon randomly inserted into their genomic DNA.



Transposon mutagenesis protocol

Transposon with selectable marker is introduced into genome of virulent (e.g. invasive) bacterial cells

↓ *Plate on medium that selects for transposon marker*

Collection of n colonies each of which has a transposon in different site in the chromosome (library)



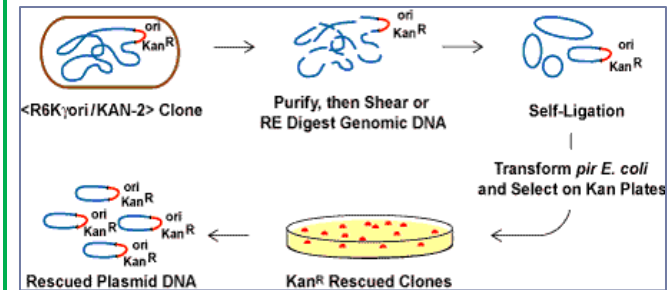
Screen n colonies for a **certain phenotype** (e.g. lost capability to invade tissue culture cells, to survive in macrophages...)



Identify the gene (x) interrupted by transposon insertion



Complementation testing by introducing the gene x to restore the wt phenotype



Limitations: mutant have to be tested individually for altered virulence. In vitro screening only. Transposon insertions have often a polar effect; genes have not to be essential;

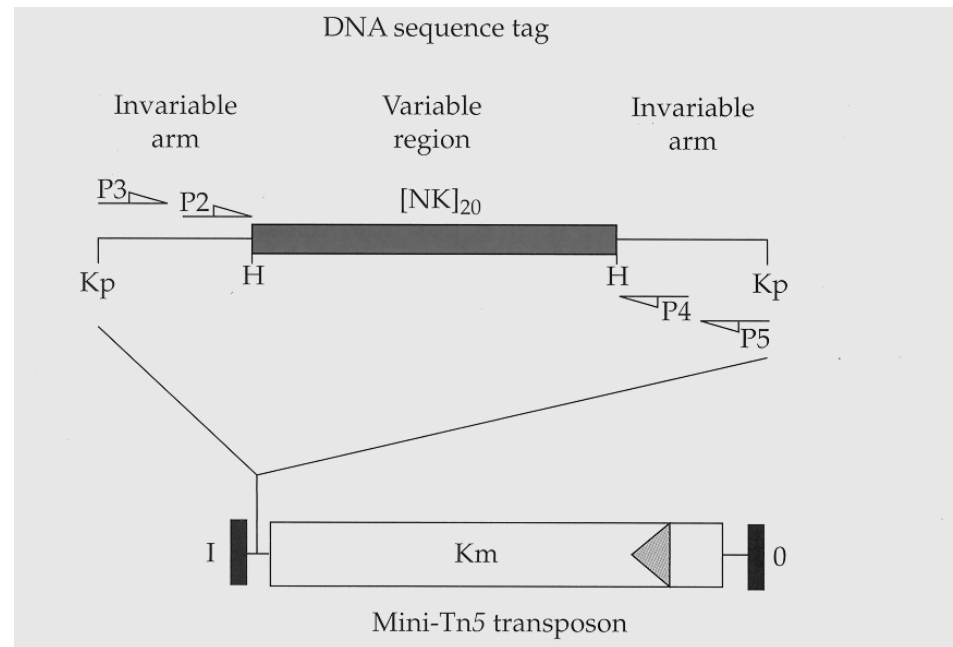
Finding Genes that are expressed *in vivo*: STM

Signature-tagged mutagenesis (STM): approach which combines *in vitro* transposon mutagenesis with *in vivo* selection using an animal model of infection.

It allows to identify genes **essential for the survival in the host** by screening for mutants that lost the ability to grow in the host (**negative selection**).

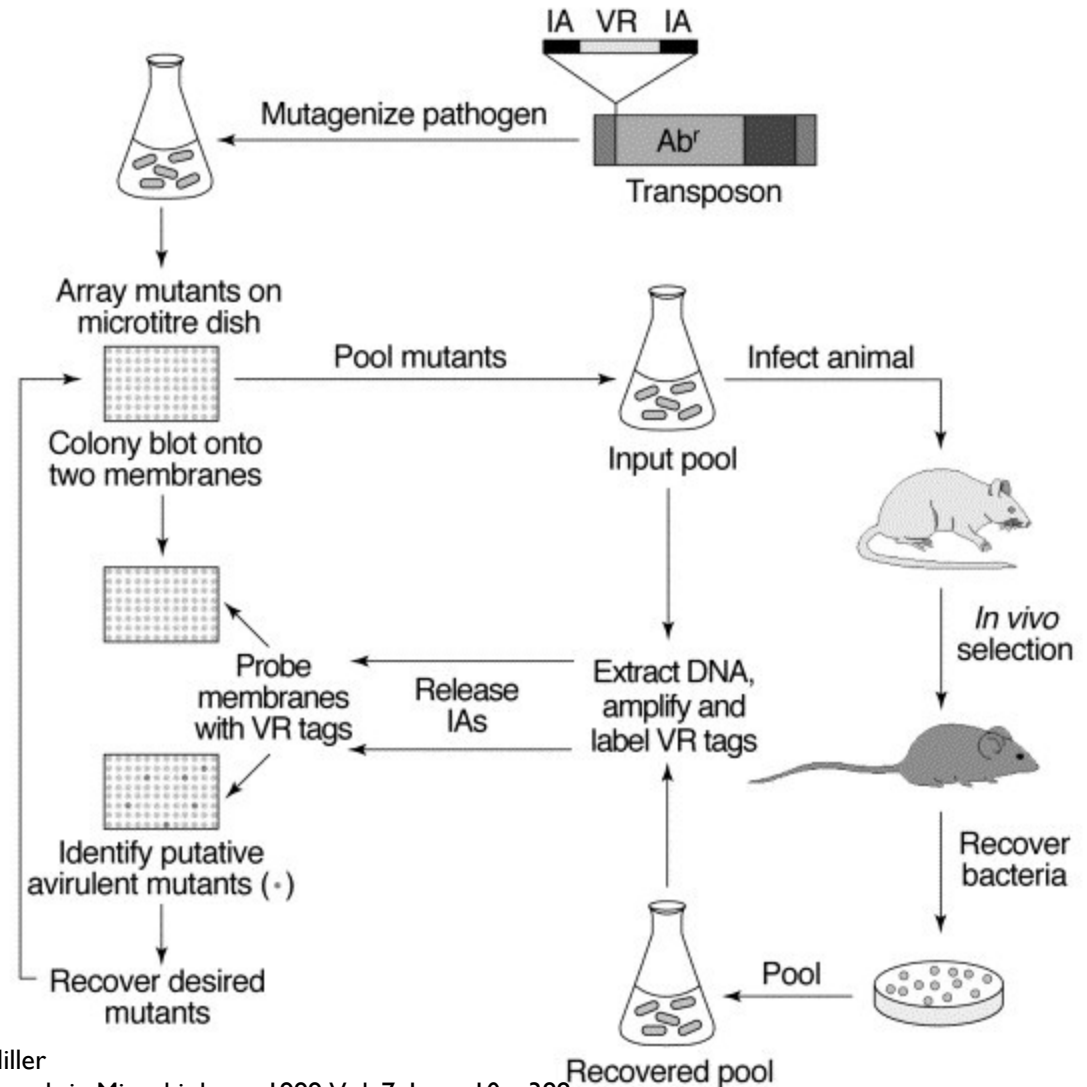
Generation of transposon variants:

STM uses a **mixture of transposon variants** to generate a library of mutants each with a different variant inserted. Procedure: variants are generated from a single transposon by cloning a mix of random oligomers (40 bp) into a transposon so that each has own **individual tag (barcode)** the oligomer it carries (transposon library).



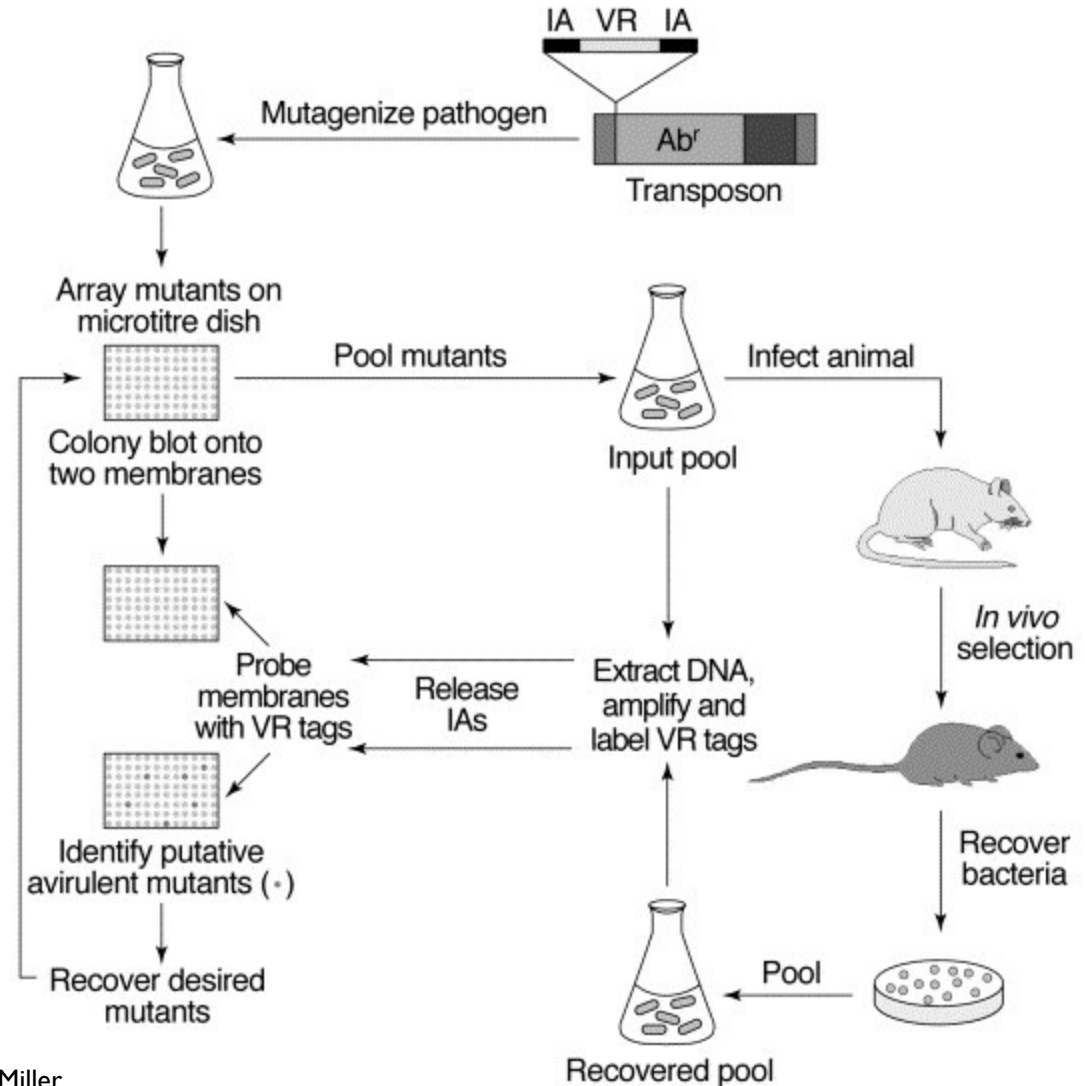
Protocol for Signature-tagged mutagenesis

- The transposon library is inserted into the target bacteria by transformation or electrophoration and selection of the transformants performed by the selective marker (Km^r).
- Individual colonies (representing each a different random mutation) are saved on masterplate (array mutants on microtitre).
- DNA is transferred on two different replicate membranes copy of the masterplate
- Mutants are pooled, grown in vitro and the culture is splitted in 2 parts:
 - I part is the input pool of mutants (control), II part is used to inoculate an animal and perform the *in vivo* selection (recovered pool).



Protocol for Signature-tagged mutagenesis (II)

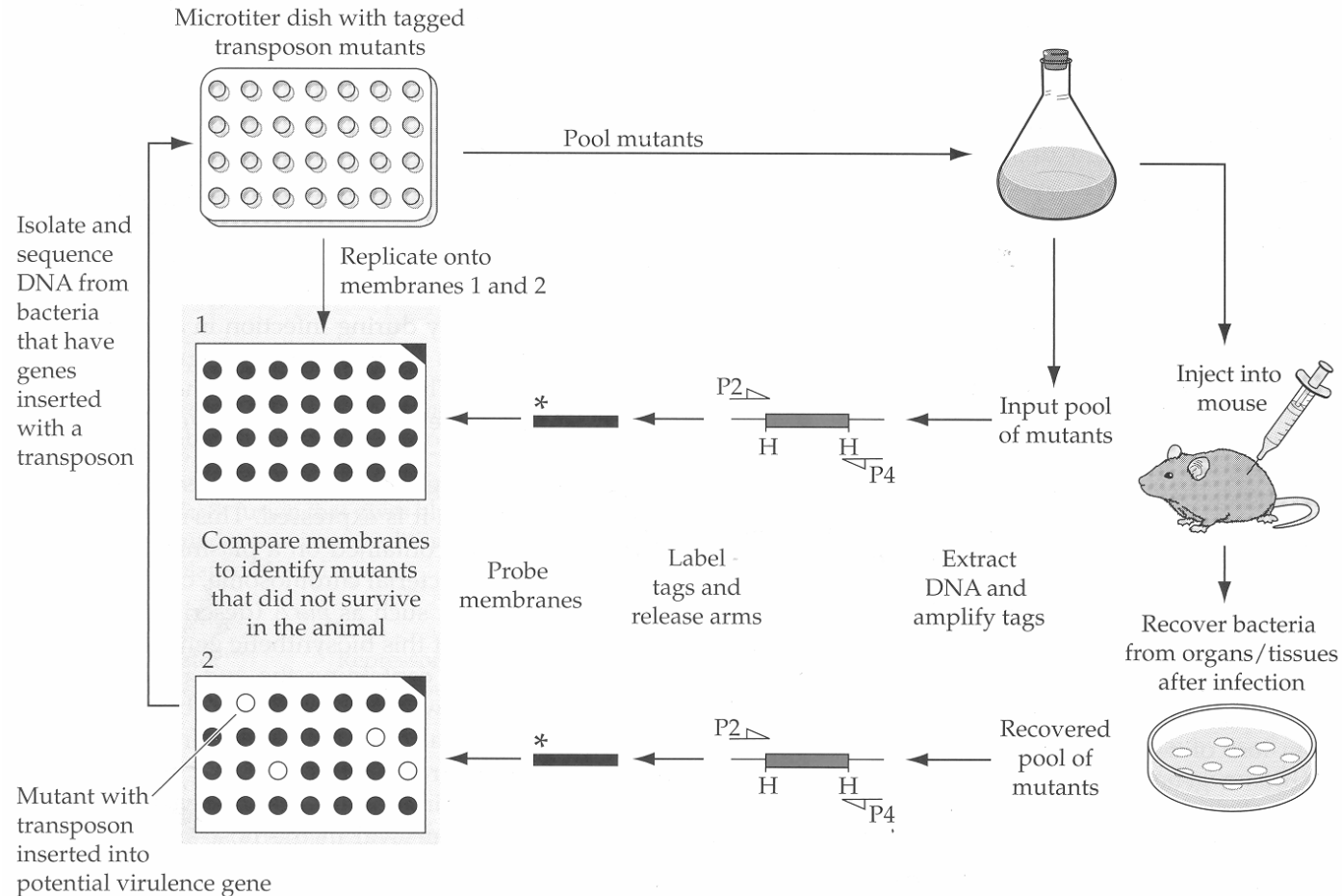
- After a time (depending from the type of infection) an organ or blood is recovered and bacteria plated in vitro.
- DNA is extracted from the both pools of bacteria and amplified using the DNA primers (labeled nucleotides) designed on the invariant oligomer tag sequence.
- Restriction enzyme digestion remove the tags sequence from the amplicons.
- The amplified DNA is used to probe the membrane containing replicates of DNA from the original collections.
- Those mutants from the output that do not hybridize with the mixture of probe tags represent **mutants lost during the infection.**



Miller

Trends in Microbiology, 1999 Vol. 7, Issue 10, p388

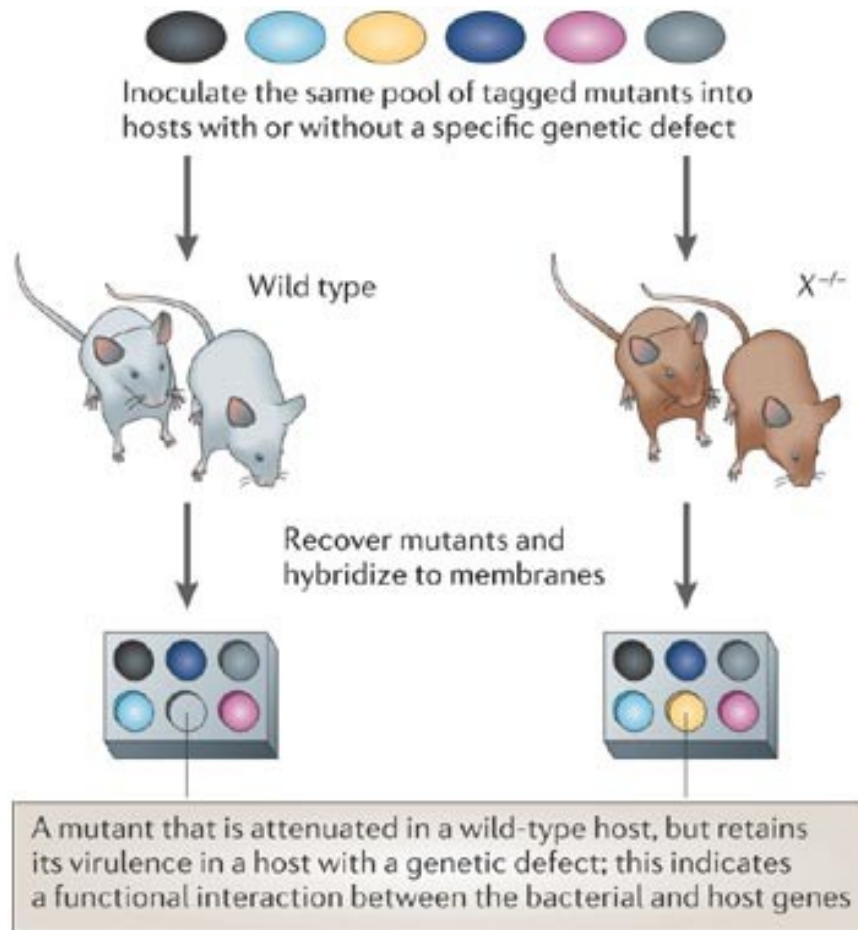
Signature-tagged mutagenesis



Mutants identified as “lost” in the host may arise from genes required for growth in the host:

- 1) needed to acquire, biosynthesize or metabolize a nutrient;
- 2) virulence genes needed for colonization, invasion, or dissemination in the host.

Detection of functional interaction between the bacterial and host genes



An STM variant could be performed with wt and a defective mice for a immune genes.

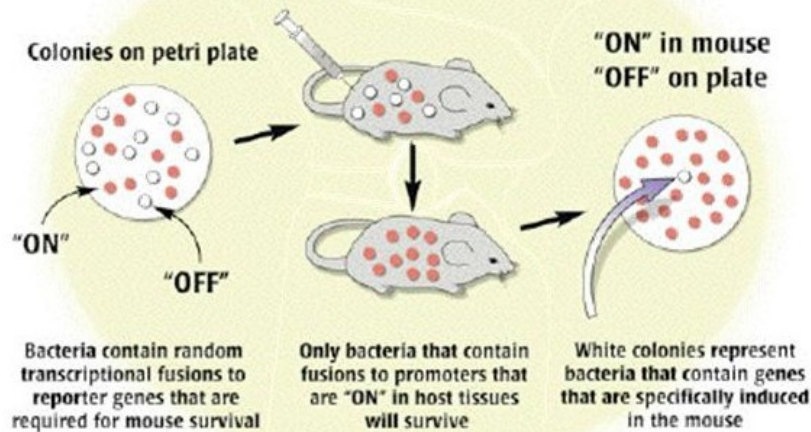
The recovery of bacterial mutants in the knock-out mice but not in the wt indicates a functional interaction between the bacterial and host genes.

In Vivo Expression Technology (IVET)

Virulence genes are usually subjected to **differential regulation** under selective conditions (e.g. ON when in the host). IVET: a method for identifying genes that are expressed when bacteria are infecting an animal host.

It has been developed to select **promoters** of genes or operons (**promoter trap technology**) that are **turned on** when the bacteria infect a host organism and turn off when bacteria are grown on laboratory medium.

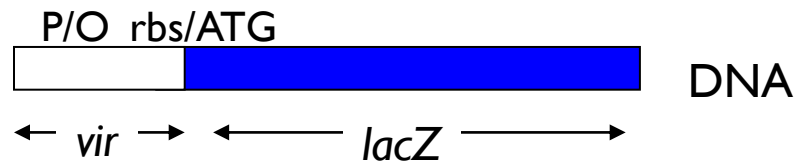
Host Specific Induction of Bacterial Virulence Genes



Experimentally characterization of the regulatory expression properties of (virulence) genes can be performed using **transcriptional fusion**:

Measuring gene regulation: transcriptional fusions

Transcriptional fusions: hybrid gene with promoter and regulatory regions of the gene of interest + structural gene (*reporter gene*) coding for a reporter enzyme that can be easily detected using several techniques (colorimetric, spectrophotometric, fluorescence or luminescence techniques).



Very popular reporter genes: β -galactosidase (*lacZ*), β -glucuronidase (*iudA*), GFP (*gfp*), Chloramphenicol acetyltransferase (*cat*) alkaline phosphatase (*phoA*) firefly luciferase (*luc*), luciferase (*lux*).



IVET: in vivo Expression Technology

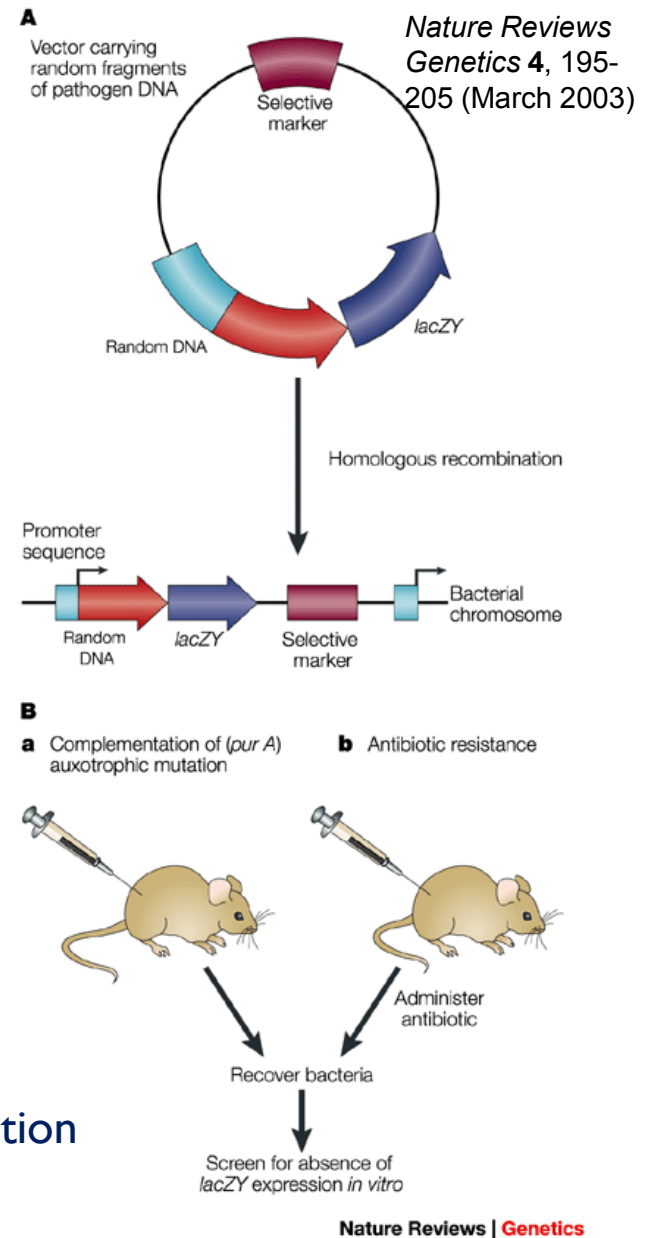
IVET vectors contain a random fragment of the chromosome of the pathogen (red) and a promoter-less gene that encodes a **selective marker** that is required for survival (dark red) (e.g. $\Delta purA$ = a purine auxotrophs unable to grow in the mice).

Random integration of the IVET vector into the pathogen chromosome is performed by insertion-duplication mutagenesis to create a pool of recombinant pathogens

Pooled clones are then inoculated into the mouse

Only those bacteria that contain the selective marker (auxotrophic mutation, or antibiotic resistance) fused to a **gene that is transcriptionally active** in the host are able to survive.

The inclusion of *lacZY* gene (blue) allows post-selection screening for promoters that are only active *in vivo*.

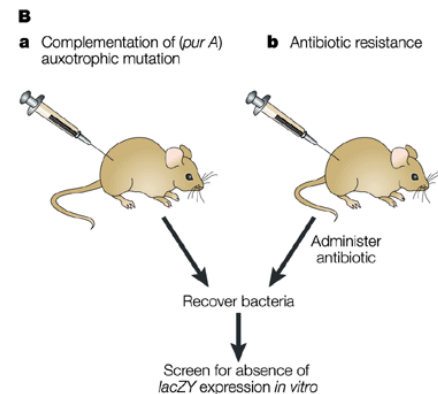
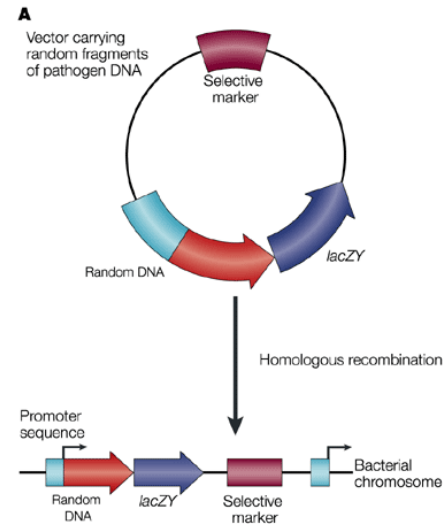


Protocol for the detection of *S. enterica Typhimurium* genes that are turned on in the mouse

The two main types of IVET promoter trap strategies are (a) the complementation of auxotrophic mutation and (b) the expression of antibiotic resistance.

Only those bacteria that contain the selective marker fused to a gene that is transcriptionally active in the host are able to survive.

After a suitable infection period, bacteria that express the marker are isolated from the spleen or other organs. The inclusion of *lacZY* gene (blue) allows post-selection screening for promoters that are only active *in vivo*.



Nature Reviews | Genetics

$\Delta purA$ = a purine auxotrophs unable to grow in the mice

STM vs IVET to identify virulence genes

STM and IVET have been successfully used for the identification and characterization of several genes of different pathogens.

STM advantages/limits:

- It is applicable to a wide range of bacteria. Selection is fast (when the transposon library is made).
- Transposon are not available for all bacteria and insertions are not completely random.

• IVET advantages/limits

- It is a selection that identify only promoters not single genes. More work is needed to examine putative virulence downstream genes
 - A high transcription level *in vivo* is mandatory
 - Genes products that are activated in the host by a posttranscriptional mechanism may be missed.
 - Auxotrophic strains of different pathogens are not common.
-



In Vivo Induced Antigen Technology (IVIAT)

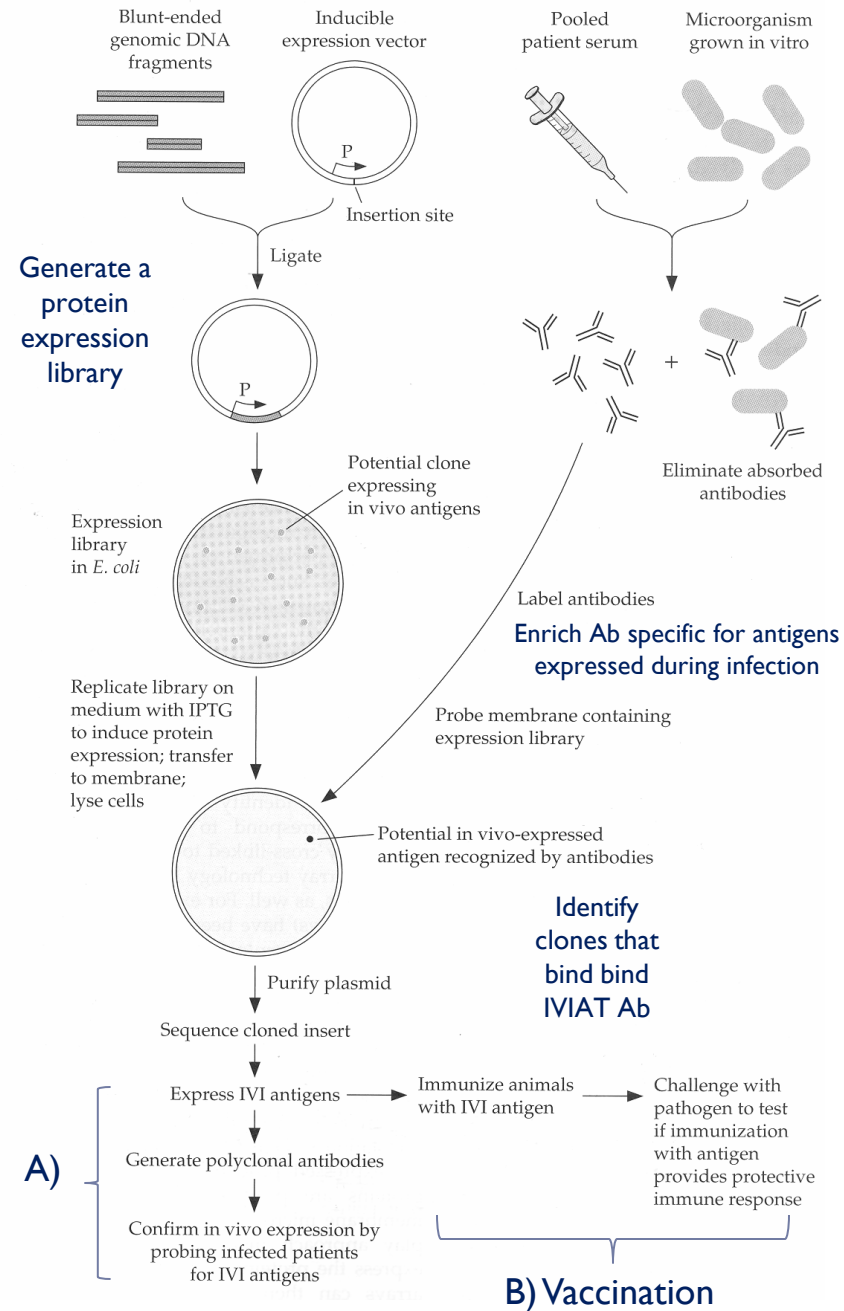
To identify in vivo-expressed antigens.

Allow to identify pathogen-associated factors that elicit immune response: potential candidates for new vaccines and diagnostic tools.

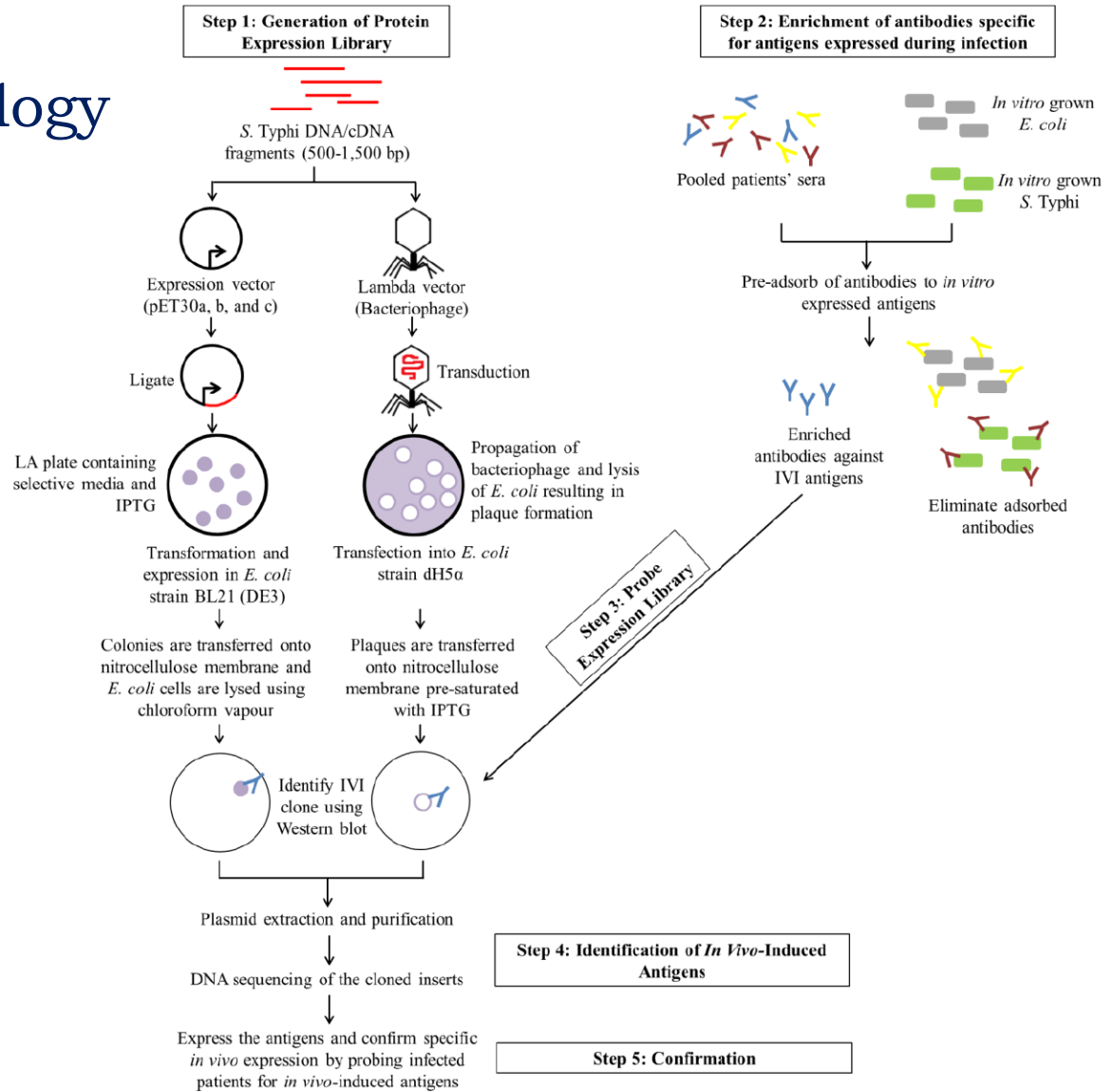
Used **pooled sera from patients** that have been exposed to a certain pathogen to probe an **expression library** obtained from the pathogen and cloned in *E. coli* (or in a phage, phage display).

Proteins recognized by the patient serum are potential **in vivo-expressed antigens**

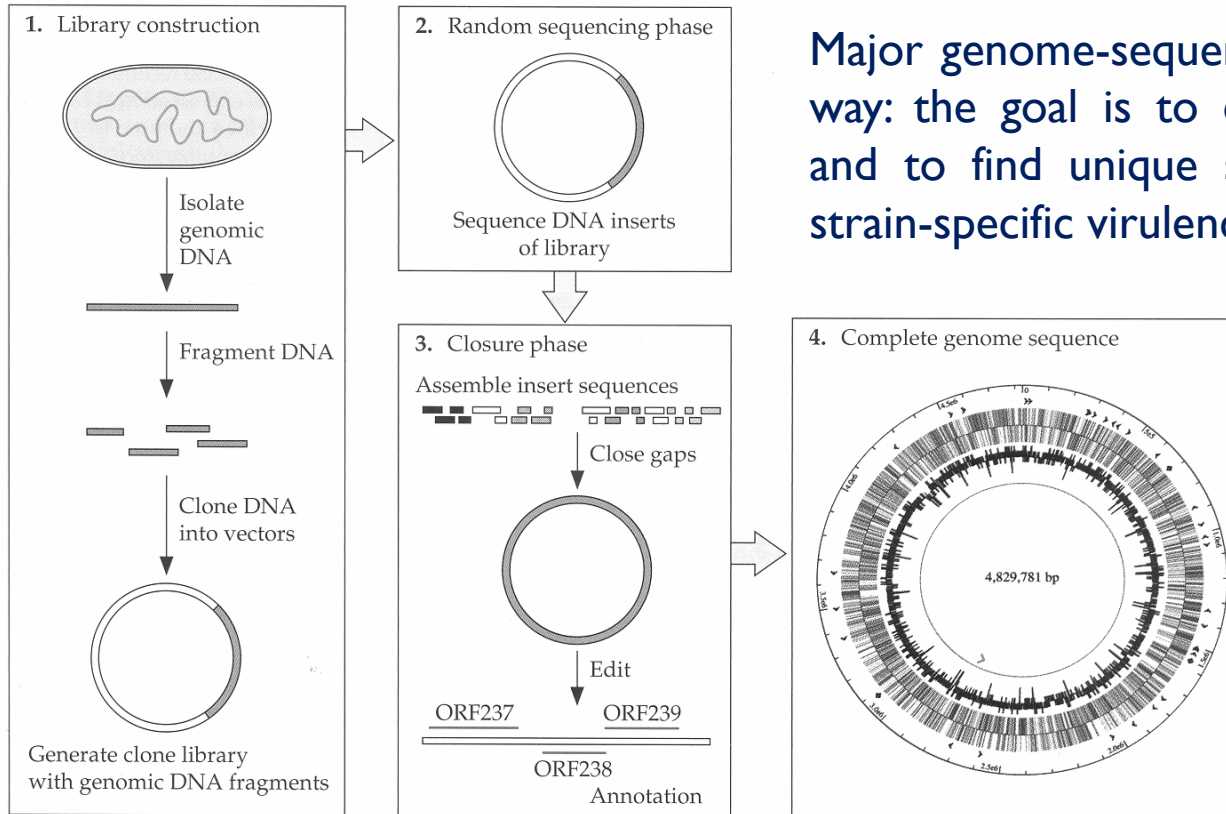
in vivo induced antigens (IVI) are then (A) inoculated in the animals to confirm their antigenicity; and (B) to test the effectiveness of the protective immune response of animals challenged with the pathogen (vaccination).



In Vivo Induced Antigen Technology (IVIAT)



Step in whole-genome shotgun sequencing



Major genome-sequencing projects are under way: the goal is to compare different strains and to find unique sequences that might be strain-specific virulence genes.

Steps in whole-genome shotgun sequencing

In shotgun sequencing, DNA is broken up randomly into numerous small segments, which are sequenced to obtain *reads*. Multiple overlapping reads for the target DNA are obtained by performing several rounds of this fragmentation and sequencing. Computer programs then use the overlapping ends of different reads to assemble them into a continuous sequence.

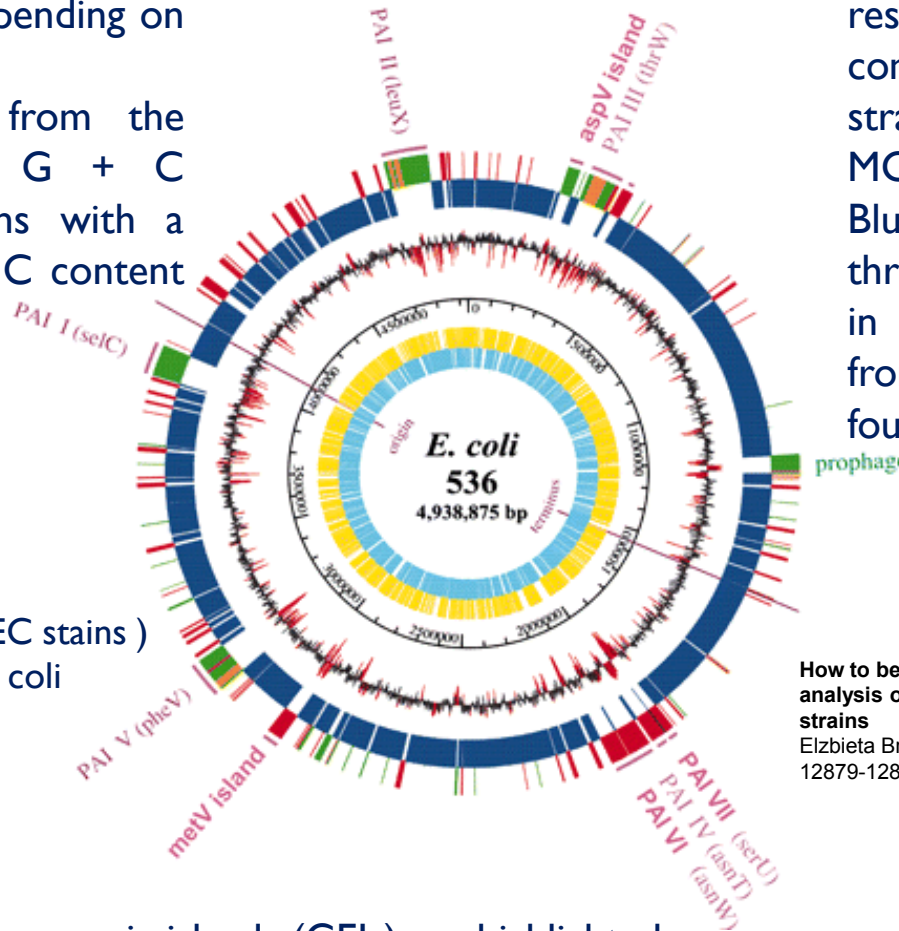


Genetic map of the UPEC strain 536 chromosome

The two inner circles represent all putative genes, depending on ORF orientation.

The fourth circle from the center shows the G + C distribution. Regions with a highly aberrant G + C content are in red.

536 and CFT073 = UPEC stains)
MGI655 = reference E. coli



The outermost circles show the result of a three-way genome comparison with the UPEC strain CFT073 and *E. coli* MGI655 (K-12) genomes.

Blue: backbone genes found in all three strains; red, genes present in 536 and CFT073 but absent from MGI655; green, genes found in 536 only.

How to become a uropathogen: Comparative genomic analysis of extraintestinal pathogenic *Escherichia coli* strains

Elzbieta Brzuszkiewicz, et al. PNAS August 22, 2006 103 (34) 12879-12884;

Pathogenicity and genomic islands (GEIs) are highlighted

Microbial genomes resource

Genome



Microbial Genomes

Microbial Genomes resource presents public data from prokaryotic genome sequencing projects. Prokaryotes are the earliest forms of life, appearing on earth 4 billion years ago. The Prokaryotes include the Archaea, which include inhabitants of some of the most extreme environments on the planet, and the Bacteria, which include both important pathogens and producers of fermented food, antibiotics, and vitamins.

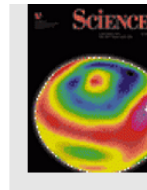
Sequencing status can be monitored at: <https://www.ncbi.nlm.nih.gov/genome/microbes/>

(10/05/2018) Eukaryotes (5657) Prokaryotes (139120) Viruses (15507) Plasmids (12745) Organelles (11722)
(16/11/2018) Eukaryotes (6817) ; Prokaryotes (170843); Viruses (21184); Plasmids (14309); Organelles (12116)

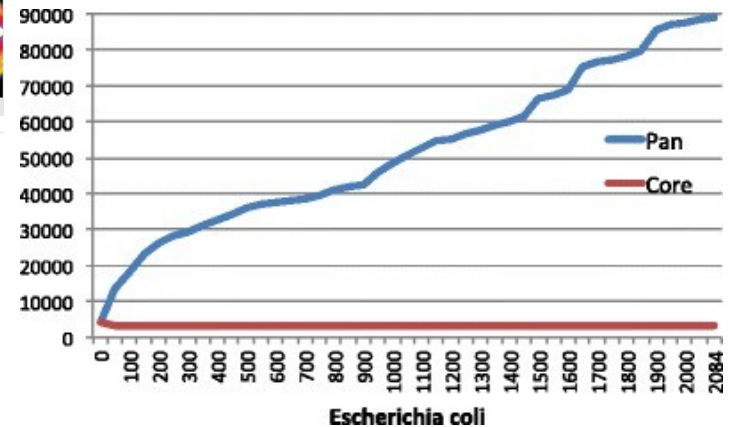
The Complete Genome Sequence of *Escherichia coli* K-12

Frederick R. Blattner*, Guy Plunkett III*, Craig A. Bloch, Nicole T. Perna, Valerie Burland, Monica Riley, Julio Collado-Vid...
* See all authors and affiliations

Science 05 Sep 1997:
Vol. 277, Issue 5331, pp. 1453-1462
DOI: 10.1126/science.277.5331.1453



Core and pan-genome of 2085 *E. coli* genomes.
Core gene families defined as those families with at least one member in at least of 95 % of genomes



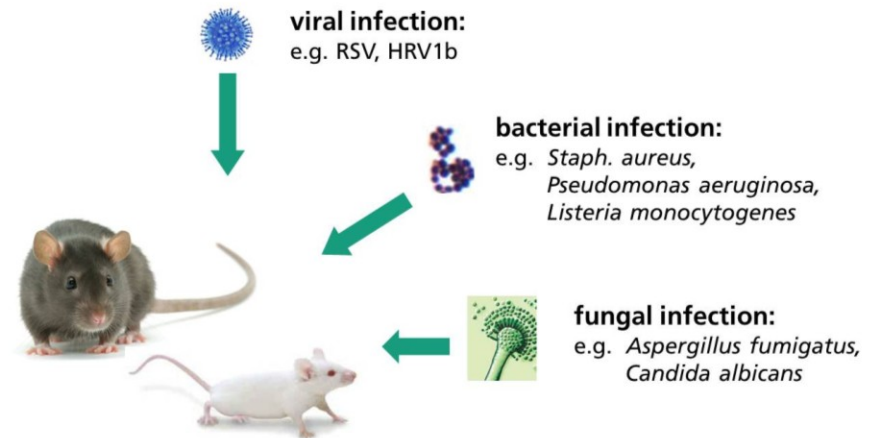
Funct Integr Genomics. 2015; 15(2): 141-161.

Measuring Infectivity and Virulence

Animal models of infections: **Gold standard for infectious diseases research.**
Usually rodents are used.

Factors that have to be taken into account:

- Differences with humans: microbiota, in part the immune system,
- Symptoms of disease could be different:
 - *i.e.* *S. enterica* in mice causes a systemic disease, *S typhi* does not infect mice.
- Use other animals: not closely related but with similar aspect of diseases:
 - Ferrets for gastric ulcers, guinea pigs as model for tuberculosis etc
 - *Caenorhabditis elegans* for *P. aeruginosa* infections etc



Ideal animal model: similar symptoms and body distribution, same route of infection. Similar course of disease

Choice of animal model: others factors.

Many factor to be considered:

- how difficult is to maintain animals

-Availability to genetically manipulated animals:

Genetically defective of some immune element

Knockout mice and knock-in mice (human gene introduced into their genome)
and germ-free animals.

Ethical considerations:

Compelling reasons for carrying out these experiments,

Approval of ethical committee is mandatory

There must be a truly strong rationale to use animals.

No alternative model can provide information equivalent.

Death of animal should not be used as an end point.

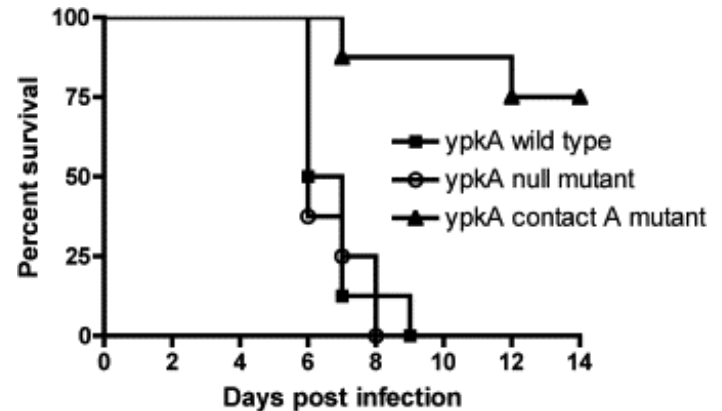
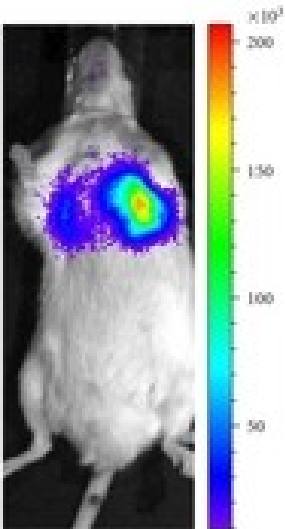


Type of in vivo Infectivity and Virulence measurements

Survival curve analysis: median survival time after infection with wt or mutant is tested for define virulence (on the right).

Common measures: LD₅₀ ID₅₀ : dose responsible to the 50% of animals dead or infected.

Measurements of colony forming units (CFU) in the blood (bacteremia) or in other tissues after mice have been sacrificed.



G. Prehna et al. Cell, 126, 2006, 869 - 880

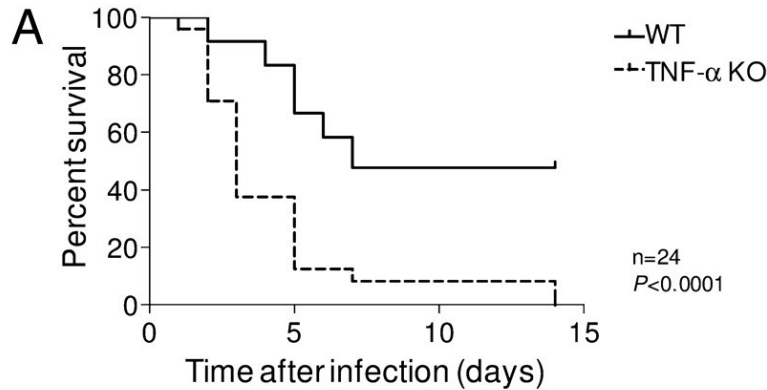
Survival-curve analysis of mice infected intragastrically with 5×10^9 CFU wt strain and mutants of *Y. pseudotuberculosis*.

In vivo imaging technology for studies of infectious disease. A bacteria luciferase operon is inserted into bacterial strains and the strain used to infect mice.

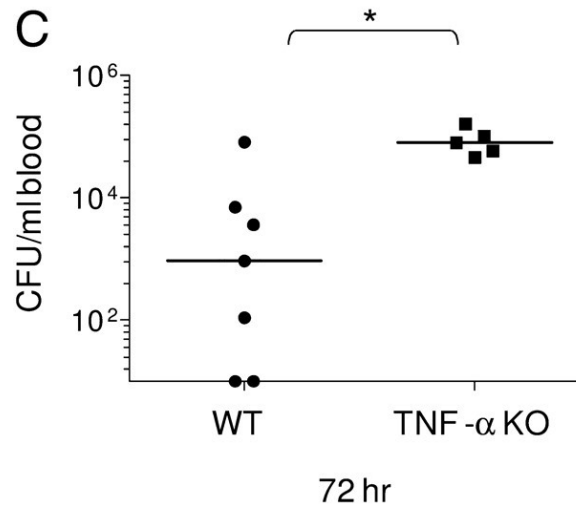
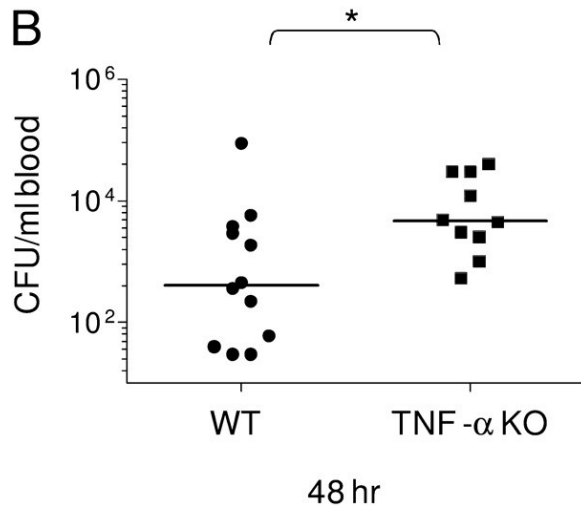
Bioluminescence imaging of invasive infection. Light emission is detected from both lung lobes indicating the establishment of bronchopulmonary infection. Bioluminescence was acquired 28 h after infection

M. Brock International Journal of Microbiology Volume 2012 (2012),

Survival-curve analysis and CFU counts of TNF- α deficient mice infected with a bacterial pathogen



WT (C57BL/6) and mutant mice deficient of the cytokine TNF- α were subcutaneously inoculated with 10^8 CFU of a strain of *group A streptococcus*, which was isolated from a patient.



TNF- α KO mice are highly susceptible to the pathogen, and showed a rapidly progressing infection and an enhanced death rate within 2–7 d after challenge.

Cell and organ culture models

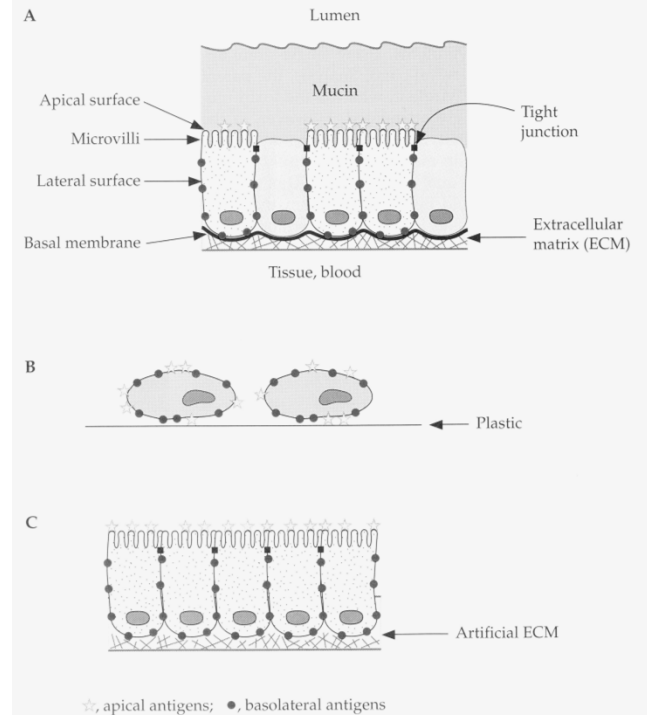
Primary cultures of mammalian cells and cell lines are used. Provide a more easily controlled system for investigating host–bacterium interactions.

Limitations: cells are no longer in the same environment. They lose many traits of the original tissues: express different markers, often they are not polarized: impossible to reproduce normal mucosa cells (difficult to mimic presence of mucus).

Addition of an extracellular substitute substrate (matrix) for the cells to grow on.

Widely used for studying: adherence properties of bacteria to host cells, attachment of bacteria, cell invasion, intracellular replication, etc.

Organ cultures: more realistic but difficult to obtain and to maintain. Development of artificial organ cultures.



Differences between cells of an actual mucosal membrane and cultured cells
A) In vivo membrane, B) nonconfluent, nonpolarized cultured cells, C) polarized monolayer of tissue culture cells attached to a semipermeable membrane