Transposon mutagenesis: past, present and future approaches

14.11.2023

Martina Pasqua



PhD in Cellular Biology and Development

Department of Biology and Biotechnology «Charles Darwin» University La Sapienza of Rome



Transposon mutagenesis

PAST

- History and importance of transposons
- Transposon as molecular tool to study bacterial genetic
- ✤ A classical experiment of transposon mutagenesis

PRESENT

- Trasposon Insertion Sequencing approaches
- TraDIS

FUTURE

- TraDIShigella infecting human organoids Project:
 - A new tool to study host-pathogen interaction: enteroids model
 - Combination of TraDIS approach and human intestinal organoids infection
 - Project workflow: preliminary results and future steps

Transposon mutagenesis

PAST

- History and importance of transposons
- Transposon as molecular tool to study bacterial genetic
- A classical experiment of transposon mutagenesis

PRESENT

- Trasposon Insertion Sequencing approaches
- TraDIS

FUTURE

- TraDIShigella infecting human organoids Project:
 - A new tool to study host-pathogen interaction: enteroids model
 - Combination of TraDIS approach and human intestinal organoids infection
 - Project workflow: preliminary results and future steps

Transposons: what are they?



The transposase recognizes the inverted repeats at the end of the transposon and also recognize the target sequence, in which it makes a double-strand break and insert the transposon.

Sequenze IS invertite

Transposons: How do they move?



Transposons: history and importance

Transposons were originally discovered as "controlling elements" in maize by Barbara McClintock in the mid-1940s.





Trends in Biochemical Sciences Volume 26, Issue 7, 1 July 2001, Pages 454-457



Forum

From controlling elements to transposons: Barbara McClintock and the Nobel Prize

Why did it take so long for Barbara McClintock (Fig. 1) to win the Nobel Prize? In the mid-1940s, McClintock discovered genetic transposition in maize. She published her results over several years and, in 1951, gave a famous presentation at the Cold Spring Harbor Symposium, yet it took until 1983 for her to win a Nobel Prize. The delay is widely attributed to a combination of gender bias and gendered science. McClintock's results were not accepted, the story goes, because women in science are marginalized, because the idea of transposition was too far-fetched and because her scientific style was too intuitive, too holistic and too feminine to be believed. In the 1960s and 1970s, transposable elements were isolated in bacteria whose amenability to genetic manipulation facilitated both detailed molecular studies of the transposition process as well as the development of transposons as molecular tools.

"Transposons can be used as tools to manipulate the genes of bacteria, phage or plasmids in ways which are otherwise difficult or impossible" Kleckner et al., 1977 Journal of Molecular Biology Volume 97, Issue 4, 5 October 1975, Pages 561-564, IN15, 565-575



Mutagenesis by insertion of a drug-resistance element carrying an inverted repetition *

Nancy Kleckner, Russell K. Chan[†], Bik-Kwoon Tye[‡], David Botstein Show more

https://doi.org/10.1016/S0022-2836(75)80059-3

Get rights and conten

A novel genetic element, which carries genes conferring tetracycline resistance (flanked by a 1400 base-pair inverted repetition), is capable of translocation as a unit from one DNA molecule to another. The *tet*^R element, which is found in nature on a variety of R-factors, was acquired by bacteriophage P22 (producing P22Tc-10 and P22Tc-106) and has now been observed to insert into a large number of different sites on the *Salmonella* chromosome. Insertion of the *tet*^R element is mutagenic when it occurs within a structural gene, and polar when it occurs within an operon. Insertion of the element is usually precise, occurring without loss of information on the recipient DNA molecule. Excision, on the other hand, is usually *not* precise, although excisions precise enough to restore a gene function can always be detected at low frequencies. Both insertion and excision processes are independent of the *recA* function.

Transposons importance in bacteria



Transposon mutagenesis in bacteria



genetic mechanism?

Making **transposon jump in vitro** in order to loss the function of the gene(s) responsible for that phenotype

Which is the gene disrupted by transposon insertion?

Localize the transposon position

Making transposons jump in vitro

In vitro transposition reaction requires:

- Transposon terminal inverted repeats
- Purified transposase
- DNA target substrate
- Reaction buffer

The in vitro transposition reaction, that can proceed with high efficiency, have been used to generate genomewide insertion mutations in a diversity of bacteria.

Microorganism	Significance	Transposon	Reference
Campylobacter jejuni	Food-borne pathogen	Tn552	26, 66
Erwinia carotovora	Plant pathogen	Mu	87
Escherichia coli	Model bacterium for genetic analysis	Tn5 Mu	47 87
Haemophilus influenzae	Pulmonary infectious agent	<i>mariner</i> Tn7	2, 3 56
Helicobacter pylori	Gastric infections and ulcers	mariner	55
Mycobacterium spp.	Opportunistic pathogen	Tn552	13, 79
Neisseria meningitidis	Meningitis agent	Tn10	145
Proteus vulgaris	Opportunistic pathogen	Tn5	47, 70
Pseudomonas sp.	Opportunistic pathogen	Tn5	70
Rhodococcus sp.	Opportunistic pathogen	Tn5	37
Saccharomyces cerevisiae	Model lower eukaryotic for genetic analysis	Tn5	47
Salmonella typhimirium	Food-borne pathogen	Tn5 Mu	47, 70 87
Streptococcus pneumoniae	Pneumonia agent	mariner	2
Streptomyces coelicolor	Antibiotic producer	Tn5, mariner	44
Synechocystis sp.	Photosynthetic cyanobacterium	ND ^a	12
Xylella fastidiosa	Plant pathogen	Tn5	54
Yersinia enterocolitica	Systemic infectious agent	Mu	87

TABLE 1 Microbial genomes mutagenized using in vitro transposition reactions

^aND, not described

Example of classical transposons mutagenesis application

Ferric Uptake Regulator Fur Is Conditionally Essential in *Pseudomonas aeruginosa*

Martina Pasqua,ª Daniela Visaggio,^b Alessandra Lo Sciuto,ª Shirley Genah,ª Ehud Banin,^c Paolo Visca,^b Francesco Imperiª

Background

Ferric Uptake Regulator (Fur) depletion makes *Pseudomonas aeruginosa* cells severely defective in colony growth on solid media.

4

<u>Aim</u>

Investigate the mechanism(s) underlying the inhibitory or toxic effect of the lack of Fur-mediated repression on colony development.

\mathbf{r}

Method

by performing transposon mutagenesis screening in order to select transposon insertion derivatives of the fur mutant able to grow on MH agar plates.

Example of classical transposons mutagenesis application

<u>Result</u>

the screening of almost 30,000 transposon insertion mutants led to identify 3 clones whose colony growth phenotype resembled that of the wild-type strain .



Limitits of a classical transposon mutagenesis approach



The necessity to assess the phenotype of each mutant individually requires considerable amount of labor and time thus limiting the total number of mutants that could be screened.

Transposon mutagenesis

PAST

- History and importance of transposons
- Transposon as molecular tool to study bacterial genetic
- ✤ A classical experiment of transposon mutagenesis

PRESENT

- Trasposon Insertion Sequencing approaches
- TraDIS

FUTURE

- TraDIShigella infecting human organoids Project:
 - A new tool to study host-pathogen interaction: enteroids model
 - Combination of TraDIS approach and human intestinal organoids infection
 - Project workflow: preliminary results and future steps

Present: applications of transposon mutagenesis





With the advent of high-throughput molecular biology approaches such as rapid nucleotide sequencing, an enormous advance in the use of transposon mutagenesis as powerful genetic instruments was made.



It is now possible to sequence many transposon mutants simultaneously allowing genome-wide analyses.

Transposon Insertion Sequencing

Is the most recent incarnation of transposon-based genomic analyses.

Is a group of similar techniques that combine transposon mutagenesis with massively parallel sequencing (MPS)

Transposon insertion sequencing (TIS)

It requires:

- 1. The construction of a transposon insertion library
- 2. Growth of the library in defined in vitro or in vivo conditions
- 3. MPS of the transposon junctions of the population at the start and at the end of the experiment
- 4. Define the frequency of each mutant in the population in order to quantify the fitness of each gene in each condition.

- Experiments are performed with pooled transposon libraries
- Critical tool to help interpret the mounting levels of genome sequencing data being generated
- Sensitive enough to detect even minor changes in mutant fitness
- Precise enough to be able to assay not only genes but also intergenic regions, promoter regions and essential protein domains within coding regions

Transposon insertion sequencing workflow



Experimental parameters that vary among TIS studies:

- selected transposon;
- complexity of the transposon libraries generated (number of independent mutants per library);
- the constraints imposed by the experimental conditions chosen;
- reliability with which representative DNA libraries are created and sequenced;
- downstream data normalization and statistical methods involved in TIS analysis.

Transposon sequencing methods



Transposon sequencing methods



TraDIS approach

Systems biology

The TraDIS toolkit: sequencing and analysis for dense transposon mutant libraries

Lars Barquist^{1,2}, Matthew Mayho¹, Carla Cummins¹, Amy K. Cain¹, Christine J. Boinett¹, Andrew J. Page¹, Gemma C. Langridge¹, Michael A. Quail¹, Jacqueline A. Keane¹ and Julian Parkhill^{1,*}

¹Wellcome Trust Sanger Institute, Hinxton, Cambridge CB10 1SA, UK and ²Institute for Molecular Infection Biology, University of Würzburg, Würzburg D-97080, Germany

Transposon-insertion sequencing screens unveil requirements for EHEC growth and intestinal colonization

Alyson R. Warr^{1,2°}, Troy P. Hubbard^{1,2°}, Diana Munera^{1,2°¤a}, Carlos J. Blondel^{1,2°¤b}, Pia Abel zur Wiesch^{1,2¤c}, Sören Abel^{1,2¤c}, Xiaoxue Wang^{1,2¤d}, Brigid M. Davis^{1,2}, Matthew K. Waldor^{1,2,3}*

Simultaneous assay of every Salmonella Typhi gene using one million transposon mutants

Gemma C. Langridge,^{1,6} Minh-Duy Phan,^{1,6} Daniel J. Turner,^{1,6} Timothy T. Perkins,¹ Leopold Parts,¹ Jana Haase,² Ian Charles,³ Duncan J. Maskell,⁴ Sarah E. Peters,⁴ Gordon Dougan,¹ John Wain,⁵ Julian Parkhill,^{1,7} and A. Keith Turne

RESEARCH ARTICLE

Combining *Shigella* Tn-seq data with gold-standard *E. coli* gene deletion data suggests rare transitions between essential and non-essential gene functionality

Open Access

Nikki E. Freed¹², Dirk Burnann² and Olin K. Slander^{1,3*}

Genome-Wide Identification by Transposon Insertion Sequencing of *Escherichia coli* K1 Genes Essential for *In Vitro* Growth, Gastrointestinal Colonizing Capacity, and Survival in Serum

References

- **Bourque** G, Burns KH, Gehring M, Gorbunova V, Seluanov A, Hammell M, Imbeault M, Izsvák Z, Levin HL, Macfarlan TS, Mager DL, Feschotte C. Ten things you should know about transposable elements. Genome Biol. 2018 Nov 19;19:199.
- Cain AK, Barquist L, Goodman AL, Paulsen IT, Parkhill J, van Opijnen T. A decade of advances in transposon-insertion sequencing. Nat Rev Genet. 2020 Sep;21(9):526-540.
- **Chao** MC, Abel S, Davis BM, Waldor MK. The design and analysis of transposon insertion sequencing experiments. Nat Rev Microbiol. 2016 Feb;14:119-28.
- **Comfort** NC. From controlling elements to transposons: Barbara McClintock and the Nobel Prize. Trends Biochem Sci. 2001;26:454-7.
- Hayes F. Transposon-based strategies for microbial functional genomics and proteomics. Annu Rev Genet. 2003;37:3-29.
- **Kleckner** N, Chan RK, Tye BK, Botstein D. Mutagenesis by insertion of a drug-resistance element carrying an inverted repetition. J Mol Biol. 1975;97:561-75.
- **Kwon** YM, Ricke SC, Mandal RK. Transposon sequencing: methods and expanding applications. Appl Microbiol Biotechnol. 2016;100:31-43.
- **van Opijnen** T, **Camilli** A. Transposon insertion sequencing: a new tool for systemslevel analysis of microorganisms. Nat Rev Microbiol. 2013;11:435-42.
- Yin Y, Zhou D. Organoid and Enteroid Modeling of Salmonella Infection. Front Cell Infect Microbiol. 2018;8:102.