

Genetic Basis of Variation in Bacteria

- I. Organization of genetic material in bacteria
 - a. chromosomes
 - b. plasmids
- II. Genetic variation: Source
 - a. point mutation
 - b. DNA rearrangements
- III. Genetic variation: Transmission
 - a. transformation
 - b. transduction
 - c. conjugation
- IV. Genetic variation: Implications for pathogenesis

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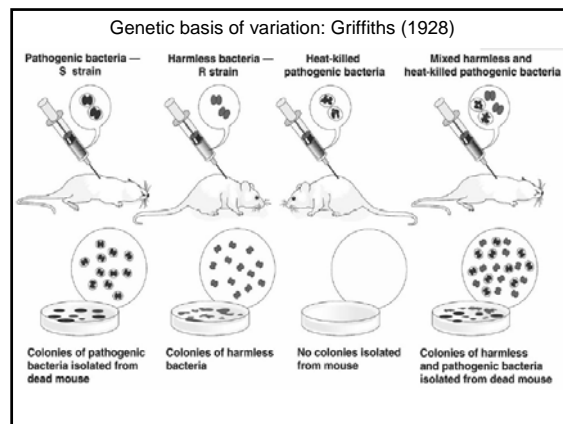
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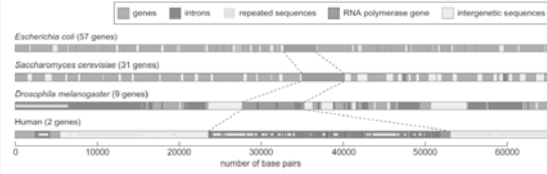


Genetic basis of variation: Avery et.al (1944)



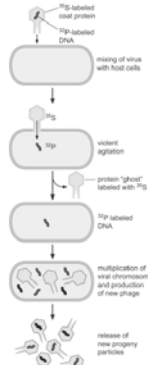
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Organization of genetic material in bacteria: chromosomes



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Genetic basis of variation: Hershey and Chase (1952)

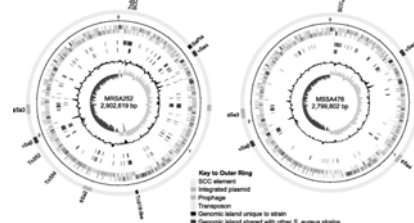


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Organization of genetic material in bacteria: chromosomes

Complete genomes of two clinical *Staphylococcus aureus* strains: Evidence for the rapid evolution of virulence and drug resistance

Matthew J. C. van den Broek¹, Edward J. Keif¹, Julia A. Lindberg¹, Sharon J. Heuvelink¹, Michelle R. J. Dwyer¹, Mark C. Wright¹, Tim J. Foster¹, Colin E. Moore¹, Laurence Hens², Rebecca Allbar¹, Andrew Barnes¹, Nathan Barron¹, Stephen B. Baxby¹, Carol Billington¹, Tracy Chittenden¹, Carol Churcher¹, Louise Clark¹, Craig Curran¹, Ann Dwyer¹, Jon Dwyer¹, Linda Good¹, Thomas Hahn¹, John Hain¹, Barbara Harris¹, Sarah Hattersley¹, Simon Heffernan¹, Kate Jolley¹, Keith D. Jolley¹, Nicola Loman¹, Alexandra Linn¹, Rebecca Moxley¹, Sharon Neill¹, Karen Murray¹, Douglas Ormond¹, Michael A. Quill¹, Elin Rukkenstein¹, Kim Sutherland¹, Mandy Sweeney¹, Sarah Taylor¹, Mark Townsend¹, Kim Warren¹, Sally Whitehead¹, Karl G. Wilson¹, Brian G. Spratt¹, and Julian Parkhill¹*



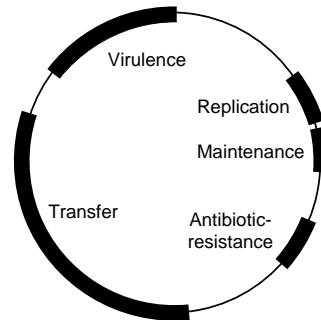
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PNAS

Organization of genetic material in bacteria: chromosomes

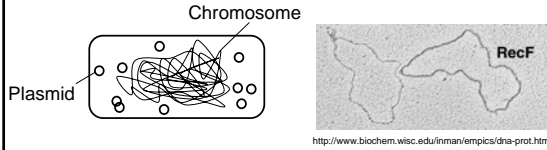
- **Most** bacteria contain a single chromosome (+ extrachromosomal elements)
- **Some** bacteria have been found also to contain 2-3 replicons which can be considered either megaplasmids or minichromosomes e.g. 3.0 Mb and 0.9 Mb replicons in *Rhodobacter sphaeroides*
- A **few** bacterial genera contain >1 chromosome e.g. 2.1 Mb and 1.2 Mb chromosomes in *Brucella*
- **Some** bacteria harbour large replicons essential for survival in a specific ecological niche but not under laboratory conditions e.g. 1.4 Mb and 1.7 Mb replicons in *Rhizobium meliloti* are required for plant symbiosis

Organization of genetic material in bacteria: plasmids

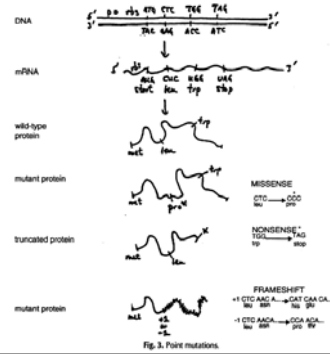


Organization of genetic material in bacteria: plasmids

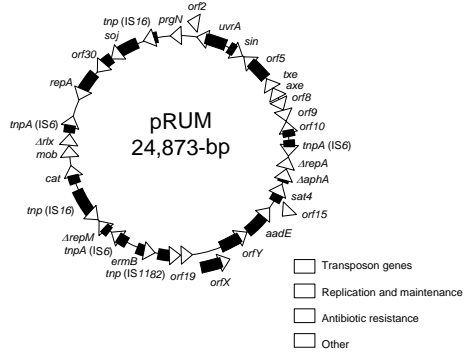
- Extrachromosomal
- Circular or linear
- 2 kb to hundreds of kb in size
- Non-essential
- May carry 'supplemental' genetic information or may be cryptic
- Employ host functions for most of DNA metabolism



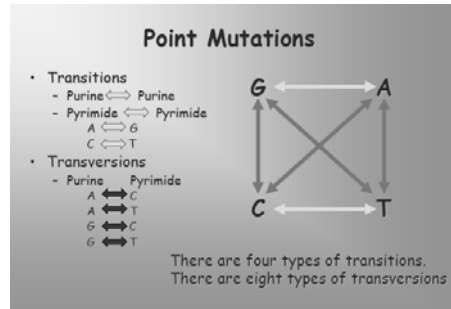
Sources of genetic variation: point mutations



Organization of genetic material in bacteria: plasmids



Sources of genetic variation: point mutations



Organization of genetic material in bacteria: plasmids

Plasmid	Host	Plasmid size (kb)	Relevant feature
pT181	<i>Staphylococcus aureus</i>	4.4	Tetracycline resistance
ColE1	<i>Escherichia coli</i>	6.6	Colicin production and immunity
pGKL2	<i>Kluyveromyces fragilis</i>	13.5	Killer plasmid
pAMβ1	<i>Enterococcus faecalis</i>	26.0	Erythromycin resistance
pSK41	<i>Staphylococcus aureus</i>	46.4	Multidrug resistance
pBM4000	<i>Bacillus megaterium</i>	53.0	rRNA operon
pZ59	<i>Staphylococcus aureus</i>	28.0	Metal ion resistance
pSLT	<i>Salmonella enterica</i> subsp. <i>typhimurium</i>	93.9	Virulence determinants
pMT1	<i>Yersinia pestis</i>	101.0	Virulence determinants
pADP-1	<i>Pseudomonas</i> sp.	108.8	Atrazine (herbicide) catabolism
pWW0	<i>Pseudomonas putida</i>	117.0	Aromatic hydrocarbon degradation
pBoxis	<i>Bacillus thuringiensis</i> subsp. <i>israelensis</i>	137.0	Mosquito larval toxicity
pX01	<i>Bacillus anthracis</i>	181.7	Exotoxin production
pSOL1	<i>Clostridium acetobutylicum</i>	192.0	Solvent production
pSymb	<i>Sinorhizobium meliloti</i>	1683.3	Multiple functions associated with plant symbiosis

Sources of genetic variation: point mutations

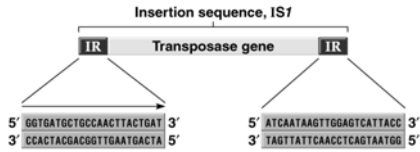
Mutation phenotypes

- Silent mutation (synonymous), no change in amino acid
AGG → AGA, both codons specify Arginine
- Missense mutation (replacement; nonsynonymous), change in amino acid
- Nonsynonymous missense (or radical replacement)
UUU (Phe) → UCU (Ser); Phe is hydrophobic and Ser is polar
- Nonsense mutation, premature termination of translation
CAG (Gln) → UAG (Stop)
- Frameshift, addition or deletion of base pairs, not in a multiple of three, within the coding region of a gene.

Sources of genetic variation: DNA rearrangements

Insertion sequence (IS) elements:

1. Simplest type of transposable element found in bacterial chromosomes and plasmids.
2. Encode only genes for mobilization and insertion.
3. Range in size from 768 bp to 5 kb.
4. IS1 first identified in *E. coli*'s galactose operon is 768 bp long and is present with 4-19 copies in the *E. coli* chromosome.
5. Ends of all known IS elements show **inverted terminal repeats (ITRs)**.



Sources of genetic variation: DNA rearrangements

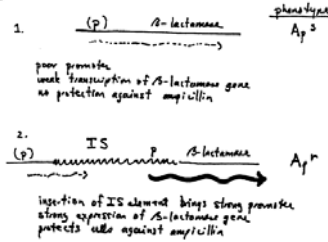
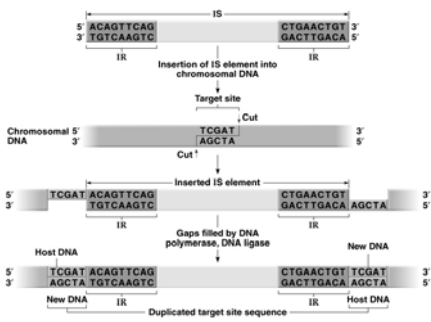


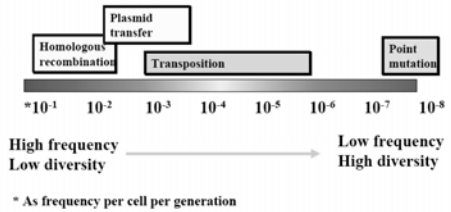
Fig. 4 (of Jeff). Activation of a gene by IS element transposition.

Sources of genetic variation: DNA rearrangements

Integration of IS element in chromosomal DNA.



Sources of genetic variation: frequency of occurrence



Sources of genetic variation: DNA rearrangements

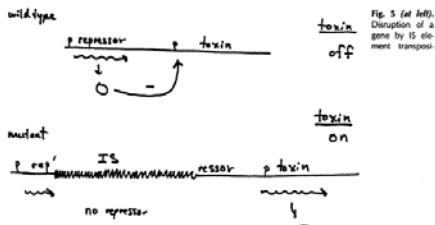
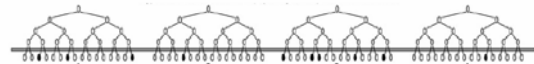


Fig. 5 (of Jeff). Disruption of a gene by IS element transposon.

Transmission of genetic variation: Luria-Delbruck test

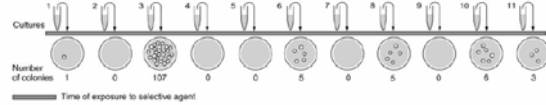
1. Resistance by mutation is a physiological response



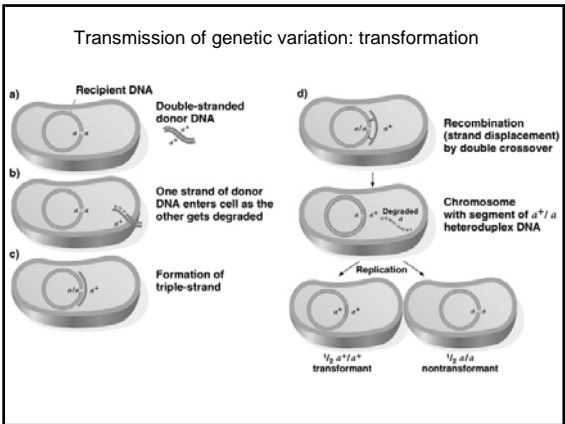
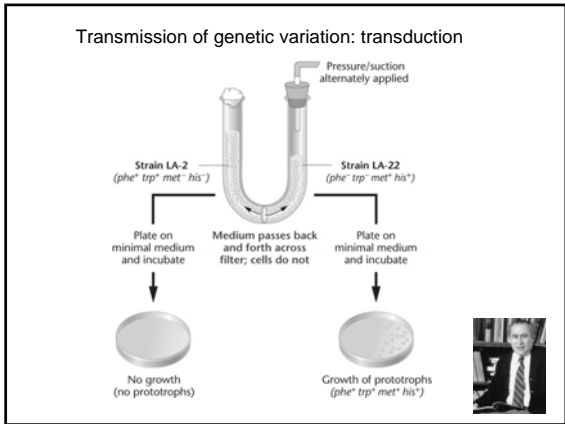
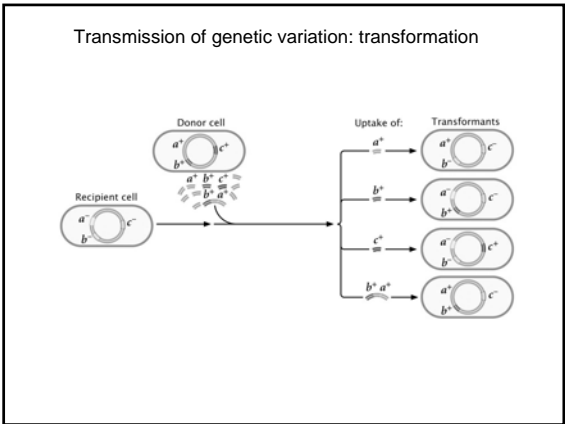
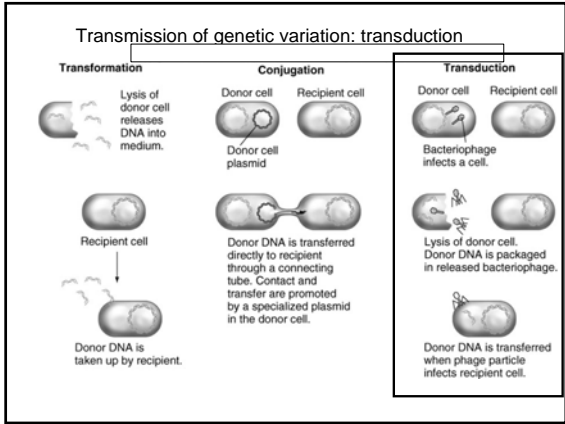
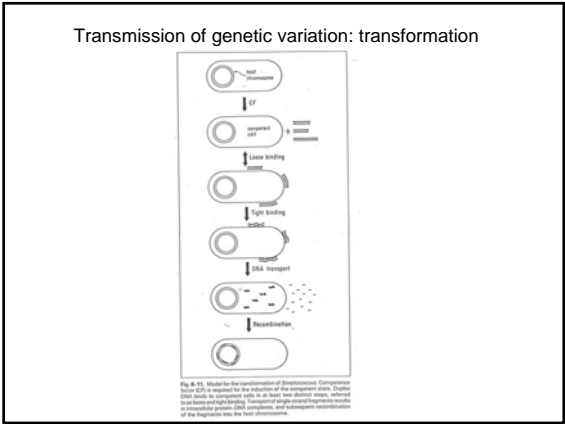
2. Resistance by mutation arises randomly in time



(b) Fluctuation test results



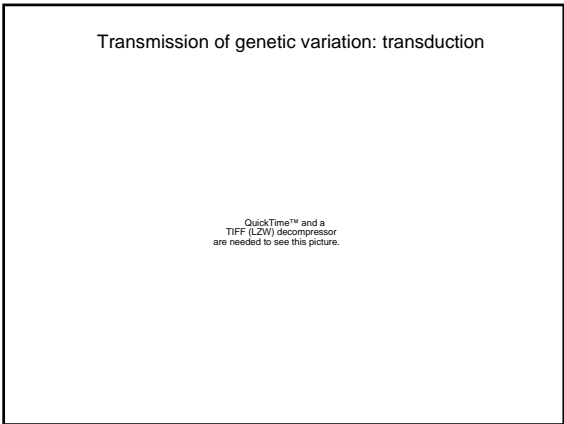
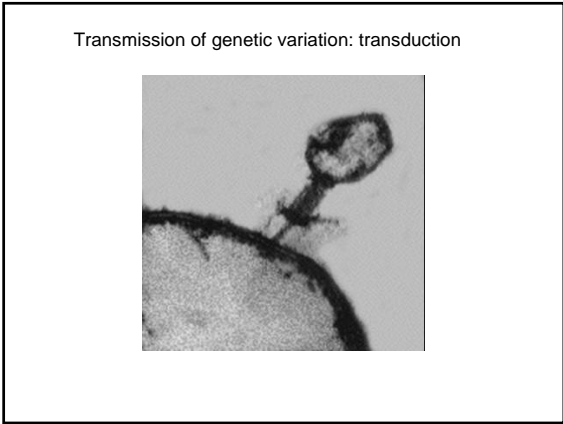
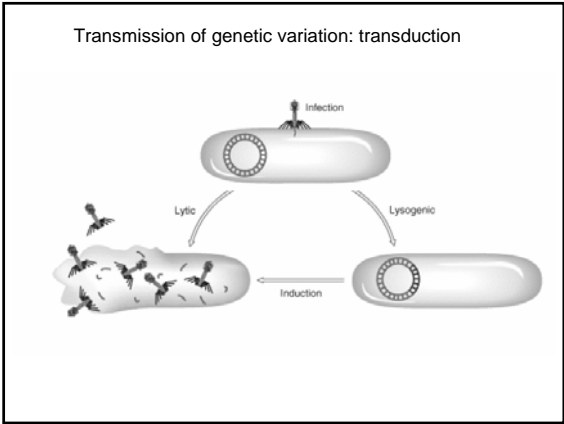
Results fit with expectations if random mutation occur at random.



Transmission of genetic variation: transduction

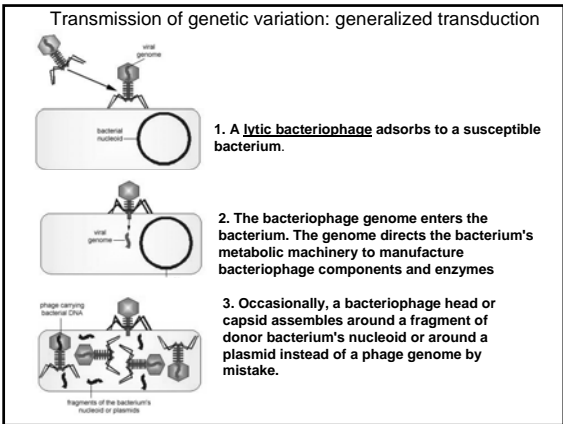
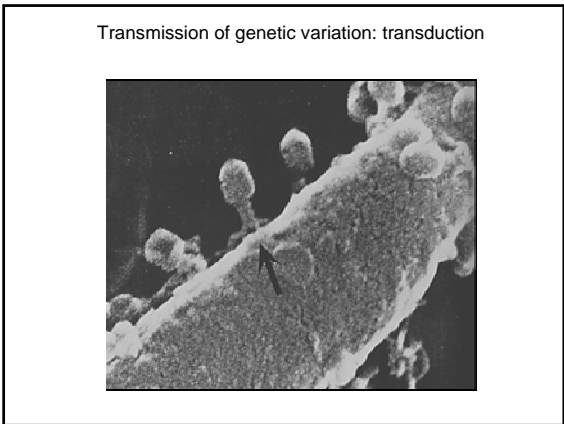
How did Zinder and Lederberg prove that the phenotype was the result of transduction?

- presence of DNAase rules out transformation
- filter prevented contact so no conjugation
- reducing filter pore size to below size of phage inhibited



Transmission of genetic variation: transduction

- There are two types of transduction:
 - generalized transduction: A DNA fragment is transferred from one bacterium to another by a lytic bacteriophage that is now carrying donor bacterial DNA due to an error in maturation during the lytic life cycle.
 - specialized transduction: A DNA fragment is transferred from one bacterium to another by a temperate bacteriophage that is now carrying donor bacterial DNA due to an error in spontaneous induction during the lysogenic life cycle



Transmission of genetic variation: generalized transduction

4. The bacteriophages are released.

5. The bacteriophage carrying the donor bacterium's DNA adsorbs to a recipient bacterium

Transmission of genetic variation: specialized transduction

3. Occasionally during spontaneous induction, a small piece of the donor bacterium's DNA is picked up as part of the phage's genome in place of some of the phage DNA which remains in the bacterium's nucleoid.

4. As the bacteriophage replicates, the segment of bacterial DNA replicates as part of the phage's genome. Every phage now carries that segment of bacterial DNA.

Transmission of genetic variation: generalized transduction

6. The bacteriophage inserts the donor bacterium's DNA into the recipient bacterium .

7. The donor bacterium's DNA is exchanged for some of the recipient's DNA.

<http://www.cat.cc.md.us/courses/bio141/lecguide/unit4/genetics/recombination/transduction.html>

Transmission of genetic variation: specialized transduction

5. The bacteriophage adsorbs to a recipient bacterium and injects its genome.

6. The bacteriophage genome carrying the donor bacterial DNA inserts into the recipient bacterium's nucleoid.

<http://www.cat.cc.md.us/courses/bio141/lecguide/unit4/genetics/recombination/transduction/specialized.html>

Transmission of genetic variation: specialized transduction

1. A temperate bacteriophage adsorbs to a susceptible bacterium and injects its genome .

2. The bacteriophage inserts its genome into the bacterium's nucleoid to become a prophage.

Transmission of genetic variation: conjugation

Transformation	Conjugation	Transduction
<p>Lysis of donor cell releases DNA into medium.</p> <p>Recipient cell</p> <p>Donor DNA is taken up by recipient.</p>	<p>Donor cell Recipient cell</p> <p>Donor cell plasmid</p> <p>Donor DNA is transferred directly to recipient through a connecting tube. Contact and transfer are promoted by a specialized plasmid in the donor cell.</p>	<p>Donor cell Recipient cell</p> <p>Bacteriophage infects a cell.</p> <p>Lysis of donor cell. Donor DNA is packaged in released bacteriophage.</p> <p>Donor DNA is transferred when phage particle infects recipient cell.</p>

