Chapter 43 BACTERIAL GENETICS: CLONES

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PRE-LECTURE ASSIGNMENT

- 1. Quickly review notes for the previous lecture.
- 2. Suggested readings:
 - a. General genetics textbooks Snyder and David: Chap. 26, pp. 401-407.

Srb and Owen: Chap. 24, pp. 534-537. Winchester: Chap. 23, pp. 318-321.

b. Additional references Bryson, V., and Szybalski, W. 1955. Microbial drug resistance. Adv. in Genet., 7: 1-46.

Lederberg, J., and Lederberg, E. M. 1952. Replica plating and indirect selection of bacterial mutants. J. Bact., 63: 399-406.

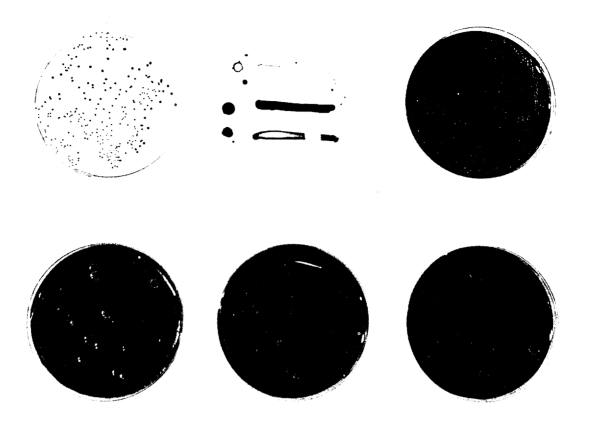
Luria, S. E., and Delbrttck, M. 1943. Mutations of bacteria from virus sensitivity to virus resistance. Genetics, 28: 491-511. Reprinted in "Papers in microbial genetics", selected by J. Lederberg. 1951. Madison: University of Wisconsin Press.

LECTURE NOTES

- A. Motivations for research with bacteria are:
 - 1. their importance in general ecology, agriculture, and disease;
 - 2. the very large populations which are easily handled in the laboratory;
 - 3. their simple cell structure.
 - a. <u>Escherichia coli</u> contains two or four nuclei, chemically defined as masses of DNA, each containing about six million nucleotide units.
 - b. DNA content of a mouse cell is about five billion units.
 - c. Although the morphological mechanism of nuclear division in bacteria is still controversial, the exact replication of

DNA occurs each cell division.

- B. <u>Vegetative (asexual) reproduction</u>
 - 1. This is the most important means for increasing bacterial numbers.
 - 2. A <u>clone</u> (see also Chap. 35) is a population of individuals all derived from a single cell by vegetative reproduction.
 - 3. Barring mutation or genetic recombination, all clonal members are genetically identical.
 - 4. Bacteria multiply rapidly.
 - a. <u>E. coli</u> divides about each half hour.
 - b. One such cell, in suitable nutrient medium, will produce a population of $N = 2^{2t} = 2^n$ individuals in t hours, or n generations.
 - c. Thus, from a single ancestor, 30 generations (requiring 15 hours) would produce about 10 billion organisms.
- C. <u>Methods for isolating a single bacterium</u>
 - 1. Directly, by the tedious but exact procedure of micromanipulation
 - 2. Indirectly, by <u>dilution</u>
 - a. When a fluid suspension of bacteria is sufficiently diluted, a sample spread on agar will contain relatively few bacteria.
 - b. Each such cell will be located on the agar at random and give rise to a visible clonal colony (Fig. 43-1, top left plate).
 - 3. Indirectly, using the simple <u>innoculating</u> <u>loop</u>
 - a. A sample of a broth culture is streaked upon fresh agar.
 - b. At some places single cells will have been deposited some distance apart, yielding separate colonies (Fig. 43-1, top right plate).
- D. Typing bacteria by clonal phenotype
 - 1. Individual cells show few morphological variations -- like presence or absence of flagella.
 - 2. Because there is not enough material per





cell, a bacterium is typed from the physiological and biochemical behavior of the clone to which it gives rise.

- 3. The top right plate in Fig. 43-1 has an eosin, methylene blue, agar medium containing lactose.
 - a. The top half was streaked with wild-type organisms, which ferment lactose via beta galactosidase, generating colored, dark, clones.
 - b. The bottom half was streaked with an ultraviolet induced lactose-negative mutant which produces light clones.
- E. Origin of bacterial mutants
 - 1. Bacteria are intimately exposed to their chemical environment.
 - 2. Mutations from lactose-negative to lactosepositive could be detected by appearance of dark colonies in the lower half of the top right plate in Fig. 43-1.
 - 3. Would the medium in that plate have induced the mutations, or would these have occurred anyway?
 - 4. The same question may be asked also whenever a mutant form has a selective advan-

tage on the culture medium employed for its detection.

- 5. For example, streptomycin-sensitive cells do not form colonies when plated on agar containing streptomycin. But a streptomycin-resistant mutant which occurs among them will form a visible colony.
- 6. Are the mutants <u>pre-adapted</u> or <u>post-adapted</u> with respect to the detecting medium? Are they spontaneous or medium-induced?
- F. <u>Pre-adaptiveness and spontaneous nature</u> of bacterial mutations
 - 1. Fluctuation test (Luria and Delbrück)
 - a. The number of mutants within a clone will depend upon the amount of time there is for multiplication before the test for them is made.
 - b. On the post-adaptive view, the number of mutants in different samples tested will fluctuate and form a normal distribution because of the random occurrence of mutations in the final generations exposed to the testing medium.
 - c. On the pre-adaptive view, the number of mutants in different samples should form

a skewed distribution. For a very few samples should contain a very large number of mutants because mutation had occurred early in clonal life, long before exposure to the testing medium.

- d. The frequency of cultures containing jackpots of mutants demonstrated these were pre-adaptive and spontaneous in origin.
- 2. <u>Three clone-sampling procedures</u> are available.
 - a. A single streptomycin-sensitive clone is plated on agar to produce a large number of colonies. <u>Each colony is individually</u> streaked across a streptomycincontaining line in the agar. All clones will grow except in the streptomycin region, but if enough clones are tested one will grow there also -- being a spontaneous, pre-adaptive, streptomycin-resistant mutant (see Fig. 43-1, top center plate).

This method is too laborious to test the pre-adaptation hypothesis.

b. <u>Replica plating of separate colonies</u> An agar plate containing up to a thousand separate colonies is pressed on velvet so that a sample of each colony is left on it. The velvet is then used as a master to plant a corresponding pattern of growth on a series of additional agar plates.

The three lower plates in Fig. 43-1 show the master (left) and two of its subsidiary plates prepared this way.

A master plate not containing streptomycin can be used to make a first copy, also on drug-free agar, and then additional copies on plates containing streptomycin. On the streptomycin plates only the resistant colonies will grow.

This also is too laborious for testing the pre-adaptation hypothesis.

c. <u>Replica plating of unseparated colonies</u> A billion or so organisms plated on agar will form small clones so closely spaced as to show continuous growth. Replicas can be made as already described.

Subsidiary streptomycin-containing plates will show growth where there are drug-resistant mutants. On the preadaptive view one can return to the corresponding site on the master plate and obtain a sample which is richer in drugresistant mutants than is a sample taken from another part of the plate. This result has been found.

- d. In all of these methods only a sample of each colony is exposed to the medium that tests for mutants, making it possible to prove the testing medium has not played a direct role in producing the mutants.
- G. Detection of bacterial mutants
 - 1. Spontaneous mutants
 - a. can be selected for, using deleterious agents.
 - b. to nutritional independence can be detected easily among nutritional mutants plated on media lacking the required nutrients.
 - 2. Very low mutation rates can be measured with these techniques. The lowest rate so far detected is one per one billion divisions for mutation from streptomycin sensitivity to resistance in <u>E. coli</u>.
 - 3. Mutagen-induced mutants also can be detected through the use of these techniques.
- H. Induced mutation
 - 1. X-rays and many other agents are mutagenic in bacteria.
 - 2. Novick and Szilard showed that purines like caffeine, adenine, and guanine increase, while their ribosides decrease, bacterial mutation rate.
 - 3. No mutagen is known at present which produces a given mutation at will.
 - 4. This reflects the fact that each gene must contain all four of the nucleotides in DNA, different genes having these in different arrangements.
 - 5. A specific mutagen would have to recognize specific assemblages of nucleotides, being itself as complicated, chemically and structurally, as the gene it mutates.
 - 6. Spontaneous mutation is in many respects an incident of the normal metabolism of the cell.

POST-LECTURE ASSIGNMENT

- 1. Read the notes immediately after the lecture or as soon thereafter as possible, making additions to them as desired.
- 2. Review the reading assignment.
- 3. Be able to discuss or define orally or in writing the items underlined in the lecture notes.
- 4. Complete any additional assignment.

QUESTIONS FOR DISCUSSION

- 43. 1. What advantages do bacteria have as material for genetic study?
- 43. 2. What disadvantages do bacteria have as genetic material?
- 43. 3. If division occurred once an hour, how many bacteria would be produced
 - a. after 4 hours, starting with one bacterium?
 - b. after 3 hours, starting with four bacteria?
 - c. after n-1 hours, if on the nth hour there were 2ⁿ?
- 43. 4. What proportion of a clone would be mutant if one cell produced by the third division underwent a mutation, but was adaptively unchanged?

What would you expect to find in this clone if there was selection for or if there was selection against the mutant?

- 43. 5. Discuss the advantages and disadvantages of various techniques for obtaining a clone from a single bacterium.
- 43. 6. Bacterial clones have been compared to the soma produced by zygotes of multicellular organisms.

Discuss whether or not this view is justified or potentially fruitful.

- 43. 7. What is the virtue of the necessity of using biochemical traits in most mutation studies with bacteria?
- 43. 8. What morphological traits of clones would be of use in bacterial genetics?
- 43. 9. What disadvantage has the use of the clone for typing the parental cell?
- 43.10. Does the post-adaptive view of the origin of bacterial mutations ever apply? Explain.
- 43.11. Suppose from a single clone of streptomycin-sensitive <u>E. coli</u> approximately 100 bacteria are placed in each of 100 test tubes containing drug-free broth. When each test tube contains about one billion individuals its contents are poured on the surface of nutrient agar medium containing streptomycin.

- a. Describe the kind of result which would prove mutations to streptomycin resistance were pre-adaptive.
- b. What kind of result would prove neither the pre-adaptive nor the post-adaptive hypothesis of mutant origin?
- 43.12. How could you show that the streptomycin resistance seen in the two central streaks in the top center plate of Fig. 43-1 was not induced by the exposure to streptomycin?
- 43.13. Explain how you would proceed to detect and isolate independent mutations from methionine-requiring to methionine-independence using the techniques of replica plating
 - a. separated colonies, and
 - b. unseparated colonies.
- 43.14. How would you proceed to detect bacterial mutations from wild-type to threonine-requiring? from threonine-requiring to threonine-independence?
- Design a specific experiment which would test whether X-rays induce mutations in bacteria.
- 43.16. Design an experiment using semi-solid culture medium to detect and collect mutations to motility.
- 43.17. Discuss Lederberg's statement that spontaneous mutation is in many respects an incident of the cell's normal metabolism.