10/1/56 Kalelson

Problem: If the units (mutons) of an allele each produce a small part of an ensyme and they are put together in a particular sequence, it is understandable why the mechanism is stopped if one muton in the same chromosome is defect. The problem is whether, in case the defect muton is supplied from a virus (λ), some synthesis of somplete ensyme can take place, i.e., without recombination to a cisform. In other words, is it possible to devise more sensitive methods than cell division and coloniformation for the study of ensyme synthesis? According to Jacob, at least 10 molecules of glucose are meded to make one bacterial cell divide, whereas less than 2 x 10 molecules are needed to produce 50 phages. The following method is proposed.

Kurahashi has found that Lederberg's number W3091 and W3091 of $K_{12}\lambda$ are PGal unidyl transferase-less. Cross \$\lambda\$ 3091 with K_{12} 3091, wash and starve; then induce with U.V. light, take an aliquot containing about 10^3 cells and plate on galactose agar which has in advance been inoculated with a very large amount of Gal + E. coli to which \$\lambda\$ is virulent. In a population of about 1000 $K_{12}\lambda$ there

should be between 0 and 5 big plaques due to recombinants. If the transduced but non-recombined K₁₂ also can produce ensyme, but much less effectively, a number of several hundred small plaques should appear after an induction period of 1 to 3 hours, for instance. Since many enzymes have rate constants (number of moles substrate reacted per mole ensyme protein) amounting to between 10⁶ to 10⁹ substrate molecules per hour, only between 10 and 1000 protein molecules of galactokinase need be produced during 1 hour to make enough UDFGall to start a small plaque. This is a true trigger reaction in the sense that one induced K₁₂ being lysed will lysogenize about 5 to 50 surrounding sensitive E. coli and so on.

Controls: Transduced W3091/W3091 (\lambda) or W309h/W309h, i.e.

'crosses' of same muton should not give any plaques. An experiment
of the same type, as well as on controls, could actually see be run
on the mutants of galacto kinase. The outlined techniques might also
be used on mixtures of protoplasts inside a cellophane container,
using heavy inoculation both of induced K₁₂\lambda, Gal- and of sensitive

E. coli, gal+. Jitis resum Andrews

This principle should also be considered for use in trying to pick up phenotypic segregation of β-glucuronidase in spermatosoos from hybrid mice. Spermatoscom are incubated with glucose-glucuronide at pil h.3. About 200 spermatesoes are spread on agar without any carbon source and plated with starved v_*v_* induced $x_{*2}\lambda$ as indicator strain (heavy bacterial inoculate). The spersatosco is the feeder of the $K_{1,2}\lambda_{\bullet}$. In a hybrid population one should observe two types of plaques: large and small (difference in size: 1 to 3 or more). In homosyg., GG (i.e. high f-glucuronidese titer) only large plaques should be present, in homosyg., gg (low glucuronidase titer) only small plaques should appear. If the spermatozoo population of a single individual show only small fluctuations, such a technique should be able to reveal phenotypic segregation in heteroxygetes (Gg) i.e., 50% large plaques and 50% small. One prerequisite which is important is that the substrates used, whether it is galactose or cellobiuronic acid, must be very pure. Any trace of glucose would

disturb greatly.

^{*} Another variation of this experiment would be to use K_{12} gal + $(\vec{s} \ \lambda)$ as the indicator strain on the galactose agar and induce with U.V. several times after plating. This would in effect man the Gal+sensitive after lysogenisation. If \vec{s} cell strain which is sensitive to without any induction, if would obviously be preferable).