July 31, 1950.

Dr. C. B. van Niel, Hopkins Marine Station, Pacific Grove, California.

Dear Dr. van Niel:

Forthwith, but under seprate cover, I am sending two cultures from <u>Escherichia coli</u>, K-12, for recombination experiments:

58-161 B-M- (biotin, methionine) W-1177 T-L-B₁- Lac- Mal-Xyl-Gal-Ara-Mtl- V₁^r S^r (threonine, leucine, thiamin lactose, maltose, d-xylose, 1-arabinose, mannitol phage T1 streptomycine)

As we discussed, the easiest characters to classify among prototrophs would be Lac and Mal, using EMB agar with 1% and 1% sugar respectively. The other characters can be ighored. The easiest technique is as follows:

Inoculate separate cultures from slant into Pennassay broth (or any other nutrient broth without too much sugar), incubate overnight at $35-37 \frac{\text{without}}{\text{without}}$ shaking or aeration. Wash cells, mix equal aliquots of concentrated suspensions and epread on thiamin supplemented plates of minimal agar, about $10^8 - 5 \times 10^9$ per plate. Use thick plates (25 ml agar/10 cm diameter plate). Incubate at 30-37 for 48-72 hours. Pick prototrophs, etc.

EMB plates should be incubated at 37 about 20 hours. Especially with EMB Maltose, Malf tends to fade after a time.

If you wish to make up a more complicated experiment, you can also score S^{T} on the EMB plates by streaking first a solution of streptomycin, 10^{5} u/ml, and cross-streaking the bacterial suspensions after the streptomycin has dried into the agar. S is very closely linked to Mal.

I would appreciate learning how this goes. Let me know if I can help in any way.

Sincerely,

Joshua Lederberg