

UNIVERSITY OF MINNESOTA

THE MEDICAL SCHOOL

MINNEAPOLIS 14

DEPARTMENT OF BACTERIOLOGY
AND IMMUNOLOGY

December 5, 1957

Dear Josh and Esther:

Welcome back! I know that you will be very busy with the house and the laboratory for the next few weeks.

I was very interested to hear that you met Sermonti. I have accordingly written to him encouraging him to come to the U.S., offering space in our lab for a few days to the full year and telling him what limited financial aid was available and asking if I might explore other means of support on his behalf. Which Bonner - John Tyler or David - is assisting Sermonti at present. I feel I should tell him - i.e. Bonner - of my action.

It is finals week here - so the part time help is absent - as well as the graduate students. This leaves me alone - which is alright until the supply of clean petri dishes gives out.

We have not succeeded in obtaining a routine method for inducing our ~~streptococci~~ lysogenic streptococci. They come out in autolysogenic cultures and can be induced by superinfection but neither are suitable. Nor have we been able to lysogenize.

We got excited for a few days about a transduction but turned out to more nearly resemble adenine auxotroph in *Brucella*.

The anomalous heterokaryons are still there. Also there is a compatibility system for heterokaryon formation. Looks like there may be a recombination system there underneath the anomalous heterokaryons and heterokaryotic selection.

We also have recombination between two *Streptomyces* phages. We use 4 hot ranges and 1 plaque marker. Select for two hot ranges and then score for the unselected. Trouble is that mixed lawns won't work - no replica plating - so we are doing it by brute force.

We have lots of minor past times which I will save until we get together.

Lois and children are fine. Will write again soon.

Gay la Bucky