Dear Spicer:
I don't know whether ay note2of the 18th April was a proper reply to yours of the 23d, or whether. I had thanked you for the H serums received some long time ago. If not, let the present suffice.

Nothing yery spectaculat bes happened during the last few months. The weather is getting $b^{\prime} l y$ hot, of courso, büt we'll havo to survive it. We did not get to the CSSH symposiun, but ăre expected a visit from Hayes later on. Tom Nēlson and I have been studying ellmination patterns more closely, in diploids from Het F+ x F-, and from diploid F $+x$ haploid F-. Along the lines of our discussions, here, the results seem to require that
 tunity for crossing-over. E.G., from $\mathrm{Mal}+\mathrm{S}^{3} \mathrm{~F}+\times \mathrm{Mal-S} \mathrm{~S}^{\mathrm{F}}$-, the diploids (selectod as pbototrophic, Lac $+/-$ etc) are about $85 \% \mathrm{Mal}-\mathrm{S}^{\mathrm{r}}, 12 \% \mathrm{Mal}+\mathrm{S}^{s}$, $28 \mathrm{Mal}+\mathrm{S}^{5}$, but none of them diploid for Mal or S. The latter items are inconsistent with the pre-elimination of Mal-S from some of the F+ gametis.

Not a thing on somatic antigen transduction, except fortuitous possibilities. In one case (not many trials) DM-x abortus equi gave a IV V XII a:enx. Eidwards is tryire to convince me that this has to be a coincidental transduction of $V$ and "diphasicity"; selection was only in enx motility agar, but the evidence that this is a transduction of either character is not yet very tacur strong. The enx serum (from ab, -equi, isn't it, might have had some IV XII) or there may have been more subtle selection for a spontaneous Form Variation. There have also been some losses and gains of I in flagellar transductions to paraA and durazzo, but these are of even less certain significance (though Kauffmann seems to be convinced by one of his own examples). Have you seen some papers by Iseki (Proc. Japan Acadery 2953) on El-E2 transformations? Some Japanse had tried to repeat Fdwards and Bruner's expts., using bolied cells for absorbing the sera (as we Ciscussed on the highway to New Orleans), and could get E2 to E1, but not the converse. On the other hand, E2 is supposed to have a phage that conver蛙 Fl to E2 by infection (all the time, not as in transduction). Edwards is digging out his old intertransformed anatum-cambridge, etc., and will I hope check on this. You may have seen by now a ms. by Edwards and myself on the flagellar transductions- he sentbone to Joan Taylor.

Are you still Interested in George Boole? The Bover edition of Laws of Thought is available (hardcover, $3 t \$ 4.50$ less $10 \%$ )- would it help you if I sent it? How are you making out to get me a Weatherburn?

Still begging, I wonder if you have gotten round to making those IV and $\nabla$ sera you oncs mentioned, and if so cculd I prevail on your generosity for some (to use in cleaning up the ab-equi, which is rather messy now. Edwatds has gone over the funzendorf story, and thinks $c^{\prime}$ is the Cl somatic antigen (as everyone else suspected). I haven't checked on the thermostability which was my only evidence against. Anti-Cl is remarkably offective in inhibiting motility if this ds so, and it might make a good stagatm system if we could do any transductions in it. So far, no. Can you enlighten me as yet on the dublin cultures, and especially the muntant you loft behind (PC-1,-2,-3;AT-2,8,14) or should I chuck tham out?

One last fayor: do you think you could spuggle out a photo of your hab. colleaguas? I'd sspecially like to soe what Felix and J. T. look like.

The most intoresting experimants around hore in a loag time are in near prospect. We've finally beon able to work up some' Gal-/Ged-, Lp ${ }^{3} / \mathrm{Lp}^{3}$, Gal,/Gal-, etc., diploids for, qise ellandeeus expts on the localization of lambde-infectioh, Gal-transdnctioh, ete. Skaprls oxpts. on motility/ $F$ are rather up In the air. Mo tility idself seems to have nothine to do with it, and the procedure may well be just a roundabout means of quaran-

The results are not nearly as consistent as we'd like. Some of the Fare (at least not readily) not reinfectable, but we have too few $F$ - from other sources to have any real basis for comparison. Skaar is continuing with it, would probably prefer not to talk too loudly about it until it is somewhat less confused.

How your heaith are you well over your Brucella? How did it ever happen? Me are, I mot danit slighty concerned dot to hear more directly from you.

2tre are looking forward to see Bernstelnthere arter ahile. I hope he will hate a chonce to yhit you beloth he leates. He is rater concerped ábout dollar problem; I.tought. Jou might be dble to advise (or at Ieast reassure) bis

 fellowship. Larry is also, rightly, pleased to have mon a mational selence Foundation Fellowship, which makes him much better off. Dave's pland are not sottied. Whare etisl hoplag ror remotoilling to improve of epace; we have ecoupled all the bpade inside the aoor afthe labding, batit will take some work to improve it to full efficiency. If you should meet any other candidates
 sume next year ( $54-55$ ); of perhaps boonet.

Do you remember'Bots Rotgan (splegeliman's, former student). He's coming up as a postidoctorig fellow to work (hotug at the Enzype Institute) on what is Intiled In the mactivation of lactase when E. 8011 celis are proken.

