

July 9, 1968

Dr. Joyce G. Small
Clinical Director of Research and
Laboratories
Larue D. Carter Memorial Hospital
1315 West 10th St.
Indianapolis, Indiana 46207

Dear Joyce,

You are correct. My associates and I did measure the changes in laboratory test data during our drug study program, and a copy of the report is enclosed.

Since completing that study we have become even more concerned that any drug study should have concurrent laboratory controls since established textbook figures are inaccurate when applied to an individual institution. We have designed a laboratory form and set up programs for the analysis of the data. Our first results were quite good, but we learned that some corrections should be made in the form itself and we have involved a private consulting firm in New York to revise our data analysis and provide the results not only for ourselves but for other institutions that may be interested. In the event that Biodata Inc. can interest some of the pharmaceutical concerns collecting data from various centers to send them the data sheets then a central bank of laboratory results, hopefully both for drug data and for placebo control will be available for analysis.

A copy of the MIP form is enclosed and the new one will be sent to you when it is available.

My best regards.

Sincerely yours,

Max Fink, M.D.
Professor of Psychiatry

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