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This information should be identified as an IDE supplement referencing the IDE number above, and must be submitted in triplicate to:

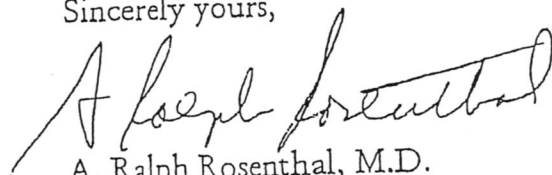
IDE Document Mail Center (HFZ-401)
Center for Devices and Radiological Health
Food and Drug Administration
9200 Corporate Boulevard
Rockville, MD 20850

If you do not provide this information within 45 days from the date of this letter, we may take steps to propose withdrawal of approval of your IDE application.

You are reminded that prior to a request for expansion beyond 150 subjects, you should provide adequate responses to deficiencies 5 - 16 in our letter of October 3, 1997.

If you have any questions, please contact Everette T. Beers, Ph.D. at (301) 594-2018.

Sincerely yours,



A. Ralph Rosenthal, M.D.

Director

Division of Ophthalmic Devices

Office of Device Evaluation

Center for Devices and Radiological Health

Enclosure:

Tables for Stability of Manifest Refraction

FDA 0 0033