COMPETITIVE PROBLEMS IN THE DRUG INDUSTRY

6. T	HE UNDERSIGNED UNDERSTANDS THAT THE FOLLOWING COND NVESTIGATIONAL USE GOVERN HIS RECEIPT AND USE OF THIS	TIONS GENERALLY APPLICABLE TO NEW DRUGS FOR INVESTIGATIONAL DRUG:
	The sponsor is required to supply the investigator with full information concerning the preclinical investigation that justifies clinical pharmacology. The investigator is required to maintain adequate records of the disposition of all receipts of the drug, including dates, quantity, and use by subjects, and if the clinical pharmacology is suspended or terminated to return to the sponsor any unused supply of the drug.	Drug Administration so notified. Upon the request of a scien fically trained and specifically authorized employee of the Department, at reasonable times, the investigator will make such records available for inspection and copying. The name of the subjects red not be divulged unless the records of the particular subjects require a more detailed study of the case or unless there is reason to believe that the records do not represent actual studies or do not represent actual studies or do not represent actual results
c.	The investigator is required to prepare and maintain adequate case histories designed to record all observations and other data pertinent to the clinical pharmacology.	f. The investigator certifies that the drug will be administered only to subjects under his personal supervision or under the supervision of the following investigators responsible to him.
	The investigator is required to furnish his reports to the sponsor who is responsible for collecting and evaluating the results, and presenting progress reports to the Food and Drug Administration at appropriate intervals, not exceeding I year. Any adverse effect which may reasonably be regarded as caused by, or is probably caused by, the new-drug shall be reported to the sponsor promptly; and if the adverse effect is alarming it shall be reported immediately. An adequate report of the clinical pharmacology should be furnished to the sponsor shortly after completion. The investigator shall maintain the records of disposition of the drug and the case reports described above for a period of 2 years following the date the new-drug application is approved for the drug; or if no application is to be filled or is approved until 2 years after the investigation is discontinued and the Food and	and that the drug will not be supplied to any other investigat or to any clinic for administration to subjects. 8. The investigator certifies that he will inform any patients or persons used as controls, or their representatives, that drugs being used for investigational purposes, and will obtain the sent of the subjects, or their representatives, except where d is not feasible or, in the investigator's professional judgment contrary to the best interests of the subjects.
	Very tru	y yours,
		(Name of Investigator)
		(Address)