



UNITED STATES  
**NUCLEAR REGULATORY COMMISSION**  
REGION I  
475 ALLENDALE ROAD  
KING OF PRUSSIA, PENNSYLVANIA 19406-1415

June 28, 2011

Docket No. 03031689  
Control No. 575471

License No. 19-07538-05

Robert Parker, Ph.D.  
Acting Director  
Laboratory of Clinical Pharmacology  
Office of Testing and Research  
Center for Drug Evaluation and Research  
U. S. Food and Drug Administration, DHHS  
Life Sciences Building #64, Room 2022  
10903 New Hampshire Avenue  
Silver Spring, MD 20993

SUBJECT: LABORATORY OF CLINICAL PHARMACOLOGY, LICENSE AMENDMENT,  
CONTROL NO. 575471

Dear Dr. Parker:

This refers to your license amendment request. Enclosed with this letter is the amended license.

Please review the enclosed document carefully and be sure that you understand and fully implement all the conditions incorporated into the amended license. If there are any errors or questions, please notify the U.S. Nuclear Regulatory Commission, Region I Office, Licensing Assistance Team, (610) 337-5239, so that we can provide appropriate corrections and answers.

An environmental assessment for this action is not required, since this action is categorically excluded under 10 CFR 51.22(c)(14).

Current NRC regulations and guidance are included on the NRC's website at [www.nrc.gov](http://www.nrc.gov); select **Nuclear Materials; Medical, Academic, and Industrial Uses of Nuclear Material**; then **Regulations, Guidance, and Communications**. You may also obtain these documents by contacting the Government Printing Office (GPO) toll-free at 1-866-512-1800. The GPO is open from 7:00 a.m. to 6:30 p.m. EST, Monday through Friday (except Federal holidays).

Thank you for your cooperation.

Sincerely,

***Original signed by Thomas K. Thompson***

Thomas K. Thompson  
Senior Health Physicist  
Commercial and R&D Branch  
Division of Nuclear Materials Safety

R. Parker

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Enclosure:  
Amendment No. 13

cc:  
Neil Hartman, Ph.D., Radiation Safety Officer

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**SUNSI Review Complete: TThompson**

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DATE	6/28/2011					

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