NRC FORM 483 (7-1999)

U.S. NUCLEAR REGULATORY COMMISSION

REGISTRATION CERTIFICATE -- in vitro TESTING WITH BYPRODUCT MATERIAL UNDER **GENERAL LICENSE**

APPROVED BY OMB: NO. 3150-0038

EXPIRES: 07/31/2002

Estimated burden per response to comply with this mandatory collection request; 7 minutes. The validated registration serves as evidence to suppliers of byproduct material that the registrant is entitled to receive the byproduct material. Send comments regarding burden estimate to the Records Management Branch (T-6 F33), U.S. Nuclear Regulatory Commission. Management Branch (1-0 F33), U.S. Nuclear Regulatory Commission, Washington, DC 20555-0001, or by internet e-mail to bis1@nnc.gov, and to the Desk Officer, Office of Information and Regulatory Affairs, NEOB-10202, (3150-0038), Office of Management and Budget, Washington, DC 20503. If a means used to impose an information collection does not display a currently valid OMB control number, the NRC may not conduct or sponsor, and a rson is not required to respond to, the information collection

Section 31.11 of 10 CFR 31 establishes a general license authorizing physicians, clinical laboratories, hospitals, and veterinarians in the practice of veterinary medicine to possess certain small quantities of byproduct material for in vitro clinical or laboratory tests not involving the internal or external administration of the byproduct material or the radiation therefrom to human beings or animals. Possession of byproduct material under 10 CFR 31.11 is not authorized until the physician, clinical laboratory, hospital, or veterinarian in the practice of syproduct material under 10 CFR 31.11 is not authorized until the physician, clinical laboratory, hospital, or veterinarian in the practice of syproduct material under 10 CFR 31.11 is not authorized until the physician, clinical laboratory, hospital, or veterinarian in the practice of syproduct materials and second physician are second physician and second physician and second physician are second physician and second physician are second physician a

UMBER. NAME AND ADDRESS OF APPLICANT (See Instruction 3.8. below)	2. APPLICATION (Check one box only)
Eugene G. Martin, Ph.D. UMDNJ - Robert Wood Johnson Medical School	I hereby apply for a registration number pursuant to 10 CFR 31, Section 31.11, for use of byproduct materials for:
University Diagnostic Laboratories One Robert Wood Johnson Place - MEB 212	Myself, a duly licensed physician authorized to disperse drugs the practice of medicine.
New Brunswick, NJ 08901	The above-named clinical laboratory.
ELEPHONE NUMBER (Include Area Code):	The above named hospital.
(732) 235–8110	Veterinarian in the practice of veterinary medicine.
NSRUCTIONS	4. REGISTRATION
A. Submit this form in duplicate to:	REGISTRATION NUMBER:
Materials Safety Branch (T-8 F5) Division of Industrial and Medical Nuclear Safety	9205
Office of Nuclear Material Safety and Safeguards U.S. Nuclear Regulatory Commission Washington, DC 20555-0001	FOR THE U.S. NUCLEAR REGULATORY COMMISSION
(At NRC, a registration number will be assigned and a validated copy of NRC Form 483 will be returned.)	Traci Kime October 5, 2001
In the box above, print or type the name, address (including ZIP Code), and telephone number of the registrant physician, clinical laboratory, hospital, or veterinarian in the practice of veterinary medicine for whom or for which this registration form is filed.	(If this an initial registration, leave this space blank – number to be assigned by NRC. If this is a change of information from a previously registered general license, include your registration
place of use is different from address listed above, give complete address.	
	iv. Diagnostic Laboratores - MEB 215, New Bru
	TIFICATION NJ 0890
hereby certify that:	
A. All information in this registration certificate is true and comple	ete.
B. The registrant has appropriate radiation measuring instrume under the general license of 10 CFR 31.11. The tests will be and in the handling of the byproduct materials.	ents to carry out the tests for which byproduct material will be used performed only by personnel competent in the use of the instrument
certificate be reported to the Director of Nuclear Material Sa change.	hange in the information furnished by a registrant on this registration afety and Safeguards within 30 days from the effective date of such
form); and I understand that the registrant is required to c receives, acquires, possesses, uses, or transfers under the U.S. Nuclear Regulatory Commission.	of NRC regulations 10 CFR 31 (reprinted on the reverse side of this comply with those provisions as to all byproduct material which his general license for which this Registration Certificate is filed with the
PRINTED OR TYPED NAME AND TITLE OF APPLICANT	SIGNAPORE) DATE
Eugene G. Martin, Ph.D.	9/21/01

ANY DEPARTMENT OR AGENCY OF THE UNITED STATES AS TO ANY MATTER WITHIN ITS JURISDICTION. PRINTED ON RECYCLED PAPER

REGULATIONS REQUIRE THAT SUBMISSIONS TO THE NRC BE COMPLETE AND ACCURATE IN ALL MATERIAL RESPECTS. 18 U.S.C. 1001 MAKES IT A CRIMINAL OFFENSE TO MAKE A WILLFULLY FALSE STATEMENT OR REPRESENTATION TO

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