KESSLER FOUNDATION

INSTITUTIONAL REVIEW BOARD

APPLICATION TO UNDERTAKE RESEARCH INVOLVING HUMAN PARTICIPANTS

IRB # (for use by administrator) —						
Submission Date: Proposed Start Date of Project: Target Completion Date of Project (i.e. publication submission):						
Title of Proposed Project:						
Description of Project:						
Principal Investigator (full name, degree) and Contact Information for PI (Mailing Address, Telephone, Email): Percentage of time to be devoted to project:% Signature of principal investigator (required)						
Co-Investigators and Full name, Degree	Study Coordina Co-Investigator	ators Department	Phone no.,	Email address	Signature (required)	
r un name, Degree	(Co-I) or Study Coordinator (SC)	or Institution	ext. (include area code)	Linaii address	Signature (required)	

Application To Undertake Research Involving Human Participants (Initial IRB Application)

I. Proj	ect Description (check all that applies):
	Is part of a grant proposal that will be/has already been submitted to a funding agency? Name of Funding Agency: Grant Application Deadline Date: Amount of Funding Requested: Time Period of Funding: Grant no.: I certify that the research protocols submitted to the IRB and to the funding agency identified above are identical.
	If the protocols submitted to the IRB and the funding agency are different, please explain.
	Principal Investigator Signature
	Is a dissertation proposal and has been approved by the dissertation committee Is a collaboration with another institution (IRB approvals for all collaborating institutions will be required)
	Name of Institution:
	☐ IRB approval copy attached, or provide explanation
	Pilot project Clinical trial
Ш	Pharmaceutical sponsor name:
	Sponsor protocol no.:
	Form 1572 copy attached (required for clinical research studies involving drugs or devices regulated by the FDA, investigator's agreement to perform the study according to applicable federal regulations)
	☐ IND copy attached (Investigational New Drug filing with FDA)
	None of the above (provide description of project)
and is contin	FEES – a fee of \$2500 may be applied to all protocols reviewed by the Kessler Foundation IRB due once the protocol has been approved and the contract or grant has been finalized; nuing review fee of \$750 will be applied annual. An exemption may be applied for through the IRB nistrator.
	Grant proposal for which an internal transfer of funds will be authorized (att. appropriate invoice memo)
	Industry sponsored study (att. appropriate invoice memo)
	Exemption - IRB fee does not apply, e.g. Federal grant, IRB fees are part of indirect costs
	Explanation:

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III. Ty	pe of Review Requested (check only one	e box):			
	EXEMPTION FROM FULL IRB REVIEW 45 CFR 46, SECTION 46.101(b)* PARAGRAPH				
	EXPEDITED REVIEW 45 CFR 46, SECTION 46.110* PARAGRAPH				
	FULL IRB REVIEW				
*See /	APPENDIX B for citation from the Code of Feder	al Regulations			
IV. Fa	cility at which the Research is to be cond	ducted (<i>check all th</i>	nat apply):		
□ We	st Orange				
□Che	_	cility)			
_	t Hanover search Population (check all that apply)				
☐ Am	inutee	Cerebrovascular A	Accident-Stroke (CVA)		
	ronic Fatigue Syndrome (CFS)	☐ Huntington's Disea	• • •		
_	Itiple Sclerosis (MS)	Orthopedic (hip, ki	` ,		
	in Management	☐ Spinal Cord Injury	,		
	numatic Brain Injury (TBI)	☐ Healthy Volunteers	• •		
_	ner (indicate research population treatment category)				
VI. H	uman subjects to be involved in the prop	osed research (<i>che</i>			
□ м	inors*		Pregnant women		
	ognitively impaired		Genetic material		
	☐ Intellectually impaired – impaired decision making ☐ Specific cognitive deficits – intact decision-making, but some deficits on certain cognitive test ☐ Non-English speaking				
*Minor detern "Unen duty in	soners rs - Persons who have not attained the legal age nined under the applicable law of the jurisdiction nancipated minor" means a person under the age n one of the military services of the United States ant to N.J.S.A. 3B:12-25 because of a finding of	in which the research e of 18 years who is ur s of America, or someo	will be conducted [45 CFR nmarried and is not current	46.402(a)]. ly serving active	
VII. R	ecruitment process: outline process for re	ecruitment			
☐ Ad	vertisements, brochures, flyers, website, letters ((ATTACHED)			
	tabases, hospital or clinic records (logbooks, sch RM is required	nedules) – Notice of Pri	ivacy Practices (NOPP) Su	ubject Certification	
□Wo	ord of mouth				
☐ Oth	ner	_			

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VIII. Study procedures (<i>check all that apply</i>): ☐ Invasive procedures	☐ MRI
Exposure to radiation	Investigational drug or device*
☐ None of the above	Questionnaire with sensitive information**
products; 2) about the subject's sexual activities and of employability, or reputation within the community; or 4 must review and approve in advance any questionnain IRB-approved study. Note: Sensitive information about protocol, when such information has previously been a criteria. IX. Conflict of Interest Statement (refer to policy)	about personal use of alcohol, illegal drugs or other addictive prientation; 3) that could damage an individual's financial standing, (1) that could lead to social stigmatization or discrimination. The IRE re that collects sensitive information from subjects enrolled in an out a subject may be recorded as part of subject recruitment into a approved by the IRB as part of the protocol's inclusion/exclusion icy #5016) ersonal financial or other interest or advisory relationship to the
If yes, please explain	
X. Consent Forms	
Provide the number of participants	

XI. Certification of Study Team Members:

Provide the number of consent forms attached

Starting January 2008, the Kessler Foundation's IRB has required that all participants in IRB-approved studies obtain certification by the Collaborative Institutional Training Initiative (CITI) by passing the CITI Course in the Protection of Human Research Subjects. Researchers should contact the IRB office for instructions on how to access the CITI web-based course. CITI certification is provided for a three year period; investigators will be reminded by CITI 90 days before their anniversary date and will be required to renew their certification at that time. For general information on the CITI program see: www.citiprogram.org

Training certifications for study team members – ATTACHED

XII. HIPAA

The Federal Health Insurance Portability and Accountability Act (HIPAA) of 1996 requires authorization to be obtained from subjects prior to their participation in research. At Kessler Foundation an application (Authorization to Use and Disclose Protected Health Information for Research Purposes) needs to be reviewed an approval provided by the Privacy Officer.

Application "HIPAA Waiver of Authorization" - ATTACHED

PROJECT APPROVAL SIGNATURE FORM

NAME OF PRINCIPAL INVESTIGATORS:				
PROJECT TITLE:				
BRIEF DESCRIPTION:				
PROJE	CT APPROVALS			
NAME (printed)	SIGNATURE	DATE		
** PRINCIPAL INVESTIGATOR				
**LABORATORY DIRECTOR(S) (if applicable)				
		<u>.</u>		
John DeLuca, PhD SENIOR VICE PRESIDENT OF RESEARCH (or designee)				
Bruce Gans, M.D. ***CHIEF MEDICAL OFFICER, KIR (or designee)				

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^{*} SIGNATURE REQUIRED for Exempt studies (Chair-IRB)

^{*}SIGNATURE REQUIRED for all <u>non-funded</u> studies (Chair-RRC)

^{**} SIGNATURES REQUIRED for all studies, PRIOR TO submission of the application to the IRB Office

^{***}SIGNATURE REQUIRED for all new Pls from KIR, PRIOR TO submission of the application to the IRB Office