NIH R01 Submission

Multidisciplinary Studies (R01 Clinical Trial Optional)

<u>Application Components</u> (Refer to the FOA for specific instructions)

Section of Application	Page Limits (If different from FOA, FOA supersedes)	Action
SF-424 Application/R&R Cover	. ,	SHP - OR
a. Project Title	Limited to 200 characters (including spaces and punctuation) Examining the effect of High-intensity Exercise to Attenuate Cognitive function Limitations and Train exercise Habits in older people living with HIV (HEALTH-Cog)	PI
Other Project Information	F 1 2 1 (1 1 2 1 1 1 1 1 1 1 1 1 1 1 1 1	
a. Project Summary/Abstract	30 lines including title	PI
b. Project Narrative	3 sentences; relevance to public health	PI
c. Bibliography & References Cited (in Research Plan and Human Subjects/Clinical Trials Information)	No page limit, concise	PI
d. Facilities & Other Resources	No page limit, concise	PI
e. Equipment (if applicable)	No page limit, concise	Pl
Sr./Key Person Profile		
a. Biographical Sketches	5 pages per Biosketch	Pl
R&R Budget / Budget Justification		
a. Budget / Budget Justification	No page limit, concise	PI / SHP-GAO
Research Plan		
a. Specific Aims	1 page limit	Pl
b. Research Strategy	12 page limit	Pl
c. Consortium/Contractual Arrangements (if applicable)	No page limit, concise	Pl
d. Letters of Support (if applicable)	No page limit, concise	Pl
e. Resource Sharing Plan(s)	No page limit, concise	Pl
f. Other Plans	No page limit, concise	
g. Appendix (if applicable)	Maximum of 10 pdfs allowed	Pl
Human Subjects and Clinical Trials	Forms H	
a. Use of Human Specimens and/or Data	Does any of the proposed research in the application involve human specimens and/or data? (Yes or No) Add an attachment that provides an explanation for any use of human specimen and/or data NOT considered to be human subjects research.	PI
b. Are Human Subjects involved? c. If Yes, Is the project exempt from federal regulations?	Yes or No Yes or No Select an Exemption Number: 1-8	PI
d. Exemption Number: e. If No to Human Subjects:	SKIP THE REST OF THE PHS HUMAN SUBJECTS AND CLINICAL TRIALS INFORMATION FORM	PI
f. Study Record(s)	Attach human subject study records using unique filenames.	PI

1.1. Study Title	Each study title must be unique; 600 character maximum	Pl
1.2. Is the Study Exempt from Federal Regulations	Yes or No	PI
1.3. Exemption Number	Select an Exemption Number: 1-8	Pl
1.4. Clinical Trial Questionnaire	Does this study meet the definition of a Clinical Trial? (Answer 4 questions)	PI
1.5. Provide Clinical Trials.gov Identifier for this trial, if applicable	If Applicable	Pl
2.1. Conditions or Focus of Study	Identify the name(s) of the disease(s) or condition(s) you are studying, or the focus of the study	PI
2.2. Eligibility Criteria	Up to 15,000 characters	Pl
2.3. Age Limits	Enter the minimum and maximum ages (or No Age Limit)	PI
2.3.a. Inclusion of Individuals Across the Lifespan	Add attachment along with an Inclusion Enrollment Report (IER) No page limit, concise	Pl
2.4. Inclusion of Women and Minorities	Add attachment along with an Inclusion Enrollment Report (IER) No page limit, concise	PI
2.5. Recruitment and Retention Plan	No page limit, concise	Pl
2.6. Recruitment Status: Choose from (not yet recruiting, recruiting, enrolling by invitation, active-not recruiting, completed, suspended, terminated, withdrawn)	Select from dropdown menu	PI
2.7. Study Timeline or Description	No page limit, concise	Pl
2.8. Enrollment of First Participant	Enter date and indicate Anticipated or Actual	Pl
2.9. Inclusion Enrollment Report(s)	Required for EACH STUDY – Add New Inclusion Enrollment Report - maximum of 20 reports	PI
3.1. Protection of Human Subjects	No page limit, concise	Pl
3.2. Is this a multi-site study that will use the same protocol to conduct non-exempt human subjects research at more than one domestic site?	Yes, No, or N/A If yes, describe the single IRB plan	PI
3.3. Data and Safety Monitoring Plan	No page limit, concise	Pl
3.4. Will a Data and Safety Monitoring Board be appointed for this study?	Yes or No	PI
3.5. Overall Structure of the Study Team	No page limit, concise	PI
4.1. Study Design		
4.1.a. Detailed Description	Up to 32,000 characters	Pl
4.1.b. Primary Purpose	Select from dropdown menu	Pl
4.1.c. Interventions	Add new intervention	Pl
4.1.d. Study Phase	Select from dropdown menu	Pl
Is this an NIH-defined Phase III Clinical Trial?	Yes or No	Pl
4.1.e. Intervention Model	Select from dropdown menu	Pl
4.1.f. Masking	Yes or No Choose One: Participant, Care Provider, Investigator, Outcomes Assessor	Pl

4.1.g. Allocation	Select from dropdown menu	PI
4.2. Outcome Measures	Add New Outcome	PI
4.3. Statistical Design and Power	Add attachment	Pl
4.4. Study Participation Duration	Fill in blank	Pl

4.5. Will the study use an FDA-regulated intervention?	Yes or No	PI
4.5.a. If yes, describe the availability of Investigational	Add attachment	PI
Product (IP) and Investigational New Drug		
(IND)/Investigational Device Exemption (IDE) status		
4.6 Is this an applicable clinical trial under FDAAA	Yes or No	PI
4.7 Dissemination Plan	Add attachment	PI
Project/Performance Site Location(s)		SHP-
		OR
PHS Assignment Request Form	Optional	PI

*Send <u>FINAL</u> items to Jill Meredith (<u>jillmeredith@uab.edu</u>) or Patrick Singer (<u>psinger@uab.edu</u>) immediately upon completion.