

This worksheet was originally created by a University of Michigan Research Administration Advisory Council (RAAC) Communications subcommittee working group, as a result of the implementation of the National Institutes of Health (NIH) SF424 (R&R) Application Packages – Version E, commonly referred to as "FORMS-E," and has subsequently been revised to accommodate the implementation of NIH SF424 (R&R) FORMS-F and FORMS-G.

The worksheet is consistent with the sequence and wording of the excerpted sections of FORMS-G contained herein, as of the "Last Updated" date in the document footer. Specifically, **this worksheet focuses** <u>only</u> on **those sections of FORMS-F related to** <u>human subjects and clinical trials</u>. It does not relate to other sections of the NIH SF424 (R&R) application package or guidance.

Information in this document is <u>only relevant to the University of Michigan (U-M)</u>. It should not be construed as having applicability anywhere outside the U-M. The worksheet serves to guide U-M personnel (specifically, investigators and research administrators) through the changes arising in FORMS-G as they relate to our institution.

FORMS-G must be used for all NIH submissions with <u>due dates on or after January 25, 2022</u>. See <u>NIH -</u> <u>How to Apply - Application Guide</u> for information related to NIH forms and applications.

The RAAC Communications subcommittee is comprised of volunteer membership. Although we will do our best to ensure this worksheet is up-to-date with the latest versions of NIH forms and instructions, we advise you e-mail the RAAC Communications subcommittee at <u>RAAC.Comm@umich.edu</u> to obtain the most current information about this worksheet.

Consult the table below for a list of recent changes to this worksheet. **For the most recent version of the worksheet, please visit the** <u>**ORSP Website**</u>.

VERSION HISTORY

Document Version	<u>Date</u>	Document Revisions
v.1	2/1/2022	(Original document – replaces NIH FORMS-F worksheet)



INSTRUCTIONS / REMINDERS:

- <u>Always consult</u> the Funding Opportunity Announcement (FOA) related to the application submission. The FOA may contain submission-specific information and requirements that are not addressed here.
- If your study qualifies as a clinical trial (*p. 4*), make sure you are <u>responding to an FOA that allows for</u> <u>clinical trial submissions</u>.
- If your study qualifies as a clinical trial (*p. 4*), make sure you <u>understand all of the requirements</u> that accompany designation as a clinical trial. For more information, see the <u>ORSP Clinical Trials website</u>.
- Create a <u>new copy</u> of this worksheet for each proposal submission.
- To make this worksheet concise, some of the text fields are <u>smaller than what their character limits will</u> <u>allow</u>. When transferring information from the worksheet to the SF424 (R&R) application package, be sure to <u>check all text fields in the worksheet</u>, as they may not print completely.
- When transferring information from this worksheet to the SF424 (R&R) application forms, be sure to review the SF424 (R&R) forms and clarify any questions with the Principal Investigator of the study.
- Answers appearing in checkboxes on this worksheet may <u>actually appear</u> as either checkboxes or drop-down menus in the SF424 (R&R) forms.
- This worksheet <u>does not address system validations within the actual SF424 (R&R) forms</u>. Be advised that system validations within the SF424 (R&R) sections referenced in this worksheet may be dependent upon other SF424 (R&R) sections <u>not contained in this worksheet</u>. Consult <u>eResearch</u> <u>Proposal Management support</u> for more information, including how to contact the ITS Help Desk.
- Use or adoption of this worksheet is entirely <u>optional and voluntary</u>. Please provide any feedback related to this worksheet to <u>RAAC.Comm@umich.edu</u>.



RESEARCH & RELATED Other Project Information

*1. Are Human Subjects Involved?

□ Yes	🗆 No	
If YES	If NO	
 answer <u>question 1.a.</u> and upload a <u>Study Record</u> (see below and p. 4) for each study. 	• skip to <u>PHS Human Subjects and</u> <u>Clinical Trials Information</u> <u>question</u> below.	

*1.a. If YES to Human Subjects (question 1 above)

Is the Project Exempt from Federal regulations?

□ Yes □ No

If yes	, check	the ap	propria	ate exe	mption	numbe	ər.	
-	□ 1			□ 4	5	□ 6	□7	□ 8
lf no	ic tha	DB rov	iow Po	ndina?				

If no, is the IRB review Pending? □ No □ Yes

IRB Approval Date (optional)

Human Subjects Assurance Number FWA 00004969

(This is the University of Michigan's Human Subjects Assurance Number.)

PHS Human Subjects and Clinical Trials Information

Use of Human Specimens and/or Data

*Does the proposed research in the application involve human specimens and/or data?

□ Yes	🗆 No
If YES	If NO

II 125	
provide an explanation of why the application does not involve human subjects (attachment)	

Study Record – add a study record for each proposed study involving human subjects (see p. 4)

Delayed Onset Study(ies)

Study Title

(Up to 600 characters)

Anticipated Clinical Trial? □ Yes □ No

Justification (attachment)



Study Record: PHS Human Subjects and Clinical Trials Information

NOTE: If the proposal has multiple studies, create a new study record with a unique title for each study.

<u>Section 1 – Basic Information</u>

*1.2. Is this study exempt from Federal Regulations?

If YES	If NO
 check the appropriate <u>exemption number</u> below. 	 skip to <u>question 1.4</u> below.

1.3. Exemption Number □1 □2 □3 □4 □5 □6 □7 □8

- *1.4. Clinical Trial Questionnaire
 - 1.4.a. Does the study involve human participants?
 - 1.4.b. Are the participants prospectively assigned to an intervention?
 - 1.4.c. Is the study designed to evaluate the effect of the intervention on the participants?
 - 1.4.d. Is the effect that will be evaluated a healthrelated biomedical or behavioral outcome?

After responding to the Clinical Trial Questionnaire, refer to the table below to determine the required application sections.

Form Section	If you answered "Yes" to <u>all</u> the questions (see Clinical Trial Questionnaire)	If you answered "No" to <u>any</u> of the questions (see Clinical Trial Questionnaire)	
<u>Section 2 – Study</u> Population Characteristics	Required	Required	
<u>Section 3 – Protection and</u> <u>Monitoring Plans</u>	Required	Required	
<u>Section 4 – Protocol</u> <u>Synopsis</u>	Required	Do not complete	
<u>Section 5 – Other Clinical</u> <u>Trial-related Attachments</u>	Required if specified in the FOA	Do not complete	

□ Yes

□ Yes

□ Yes

□ Yes

□ No

□ No

1.5 Provide the ClinicalTrials.gov Identifier (e.g., NCT87654321) for this trial, if applicable.

(Newly proposed studies do not need to be entered in ClinicalTrials.gov at the time of application.)



Section 2 – Study Population Characteristics NOTE: This section is required <u>unless</u> you selected only
Exemption 4 and no other exemptions on the "1.3 Exemption Number" question

2.1	Conditions or Focus of Study			
2.2	Eligibility Criteria			
2.3	Age Limits			
	Minimum age (enter number) (check one) □ Years □ Months □ Weeks □ Days □ Hours □ Minutes □ N/A (no limit)			
	 Maximum age (enter number) (check one) □ Years □ Months □ Weeks □ Days □ Hours □ Minutes □ N/A (no limit) 2.3.a Inclusion of Individuals Across the Lifespan (attachment) 			
2.4	Inclusion of Women and Minorities (attachment)			
2.5	Recruitment and Retention Plan (attachment)			
2.6	Recruitment Status			
	Image: Not yet recruitingImage: Completed(check one)RecruitingImage: SuspendedImage: DescriptionImage: Terminated (Halted PrematurelyImage: DescriptionImage: SuspendedImage: DescriptionImage: Terminated (Halted PrematurelyImage: DescriptionImage: SuspendedImage: DescriptionImage: Terminated (Halted PrematurelyImage: DescriptionImage: Terminated (Halted Prematurely)Image: Description			
2.7	Study Timeline (attachment)			
2.8	Enrollment of First Participant Date (check one) Anticipated Actual			
2.9	Inclusion Enrollment Report(s) (attachment(s)) (see sample, p. 9)			
Section 3	8 – Protection and Monitoring Plans			
3.1	Protection of Human Subjects (attachment)			
3.2	Is this a multi-site study that will use the same protocol to conduct non-exempt hum			

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?

("N/A" is only a valid option if you answered "Yes" to qu	estion 1.2 above or you are a training g	grant applicant.)

If YES	If NO	
• describe the single IRB plan (attachment)	• skip to <u>question 3.3</u> below.	

- 3.3 Data and Safety Monitoring Plan (attachment) (<u>Required for clinical trial; optional for human subjects.</u>)
- **3.4 Will a Data Safety and Monitoring Board by appointed for this study?**
 U Yes
 No (Answer is required for clinical trial; answer is optional for human subjects.)
- 3.5 **Overall Structure of the Study Team** (attachment) (Optional refer to your specific FOA)



<u>Section 4 – Protocol Synopsis</u> *Refer to the Clinical Trial Questionnaire on p. 4. If you answered "No" to any of the questions, this section is <u>not required</u>.

4.1 Study Design:

4.1.a. **Detailed Description**

(Up to 32,000 characters; typically needs only 5,000 characters)

4

4.1.b.	Primar	y Purpose	
(check one)		 □ Treatment □ Prevention □ Diagnostics □ Supportive Care □ Other 	 Screening Health Services Research Basic Science Device Feasibility
4.1.c.	Interve (up to 20	entions Interventions allowed.)	
(check	one)	Intervention Type: Drug (including placebo) Device (including sham) Biological/Vaccine Procedure/Surgery Radiation Behavioral (e.g., Psychotherapy, Lifestyle Counseling)	 Genetic (including gene transfer, stem cell, and recombinant DNA) Dietary Supplement (e.g., vitamins, minerals) Combination Product Diagnostic Test Other
4.1.d . (c	(check one) □ Early Phase 1 (or Phase □ Phase 1 □ Phase 1/2 □ Phase 2 (If selecting "Other", response is		[•] Other" and provide explanation.) □ Phase 2/3 □ Phase 3 □ Phase 4 □ N/A
4.1.e.		ention Model	

4

□ Single Group □ Parallel (check one)

□ Factorial □ Sequential

□ Other _

□ Cross-Over

(If selecting "Other", response is limited to 255 characters.)



Section 4 – Protocol Synopsis (cont.)

	4.1.f. N	lasking		
		□ Yes	🗆 No	
		If YES	If NO	
	 check all that apply below. 		• skip to <u>question 4.1.g</u> below.	
		icnerk all that annivi	rticipant □ Investigato re Provider □ Outcomes	
	4.1.g. A	llocation		
	(*	check one) □ N/A	\Box Non-randomized \Box F	Randomized
4.2.	Outcome N (At least one of	leasures Outcome Measure required, unless n	oted in the opportunity. Up to 50 Out	come Measures allowed.)
	Na	me		
	(Up	to 255 characters.)		
	Ty (ch	pe eck one) □ Primary □ Se	condary 🛛 Other	
	Tin	ne Frame		
		to 255 characters.)		
		ef Description		
	• •	to 999 characters.) <u>outcome measure entries avai</u>	ilable on p. 12. if needed)	
4.2				
4.3.	Statistical	Design and Power (attachmer	IT)	
4.4.	Subject Pa (Up to 255 ch	rticipation Duration		
4.5.	Will the stu	udy use an FDA-regulated in		
			□ No	
	4.5. <u>a</u> .			
		If YES	If NO	
	Prod (IND	ribe the availability of Investiga uct (IP) and Investigational Ne)/ Investigational New Drug nption (IDE) status <i>(attachment)</i>	• skip to <u>question 4.</u>	<u>6</u> below.
4.6.	ls this an a	upplicable clinical trial under	FDAAA? 🗆 Yes 🗆 No	
4.7		tion Plan (attachment)		



Section 5 – Other Clinical Trial Attachments (attachments, if required by FOA)

*Refer to the Clinical Trial Questionnaire on p. 4. If you answered "No" to any of the questions, this section is not required.

REQUIREMENTS:

RESEARCH & RELATED Other Project Information

Yes	1	•	Attach a Study Record (p. 4) for each study
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PHS Human Subjects and Clinical Trials Information

Yes	Human Specimens and/or Data	• Attach an explanation of why the application does not involve human subjects (<i>p. 3</i>)
Yes	Delayed Onset Study(ies)	 Complete Delayed Onset Study questions (see p. 3) Attach Justification (see p. 3)

Study Record: PHS Human Subjects and Clinical Trials Information

Yes to <u>all</u>	1.4.a. through 1.4.d.	 Complete Section 2 (p. 4) Complete Section 3 (p. 5) Complete Section 4 (p. 6) Complete Section 5 (p. 8), if required by FOA
No to <u>any</u>	1.4.a. through 1.4.d.	 Complete Section 2 (p. 4) Complete Section 3 (p. 5)
Yes	3.2	Attach Single IRB Plan (p. 5)
Yes	4.6	Attach Description (p. 7)

ATTACHMENT CHECKLIST: (Not all of the attachments listed below may be required. Consult the table above.)

Section	Question	Description	-		Page
Oection	Question	Description			i age

PHS Human Subjects and Clinical Trials Information

N/A	Human Specimens and/or Data	Explanation of why the application does not involve human subjects	3
N/A	Study Record(s)	Study Record: PHS Human Subjects and Clinical Trials Information for each study	3
N/A	Delayed Onset Study(ies)	Justification explaining why human subjects study information is not available at the time of application	3

Study Record: PHS Human Subjects and Clinical Trials Information

2	2.3.a	Inclusion of Individuals Across the Lifespan	5
2	2.4	Inclusion of Women, Minorities, and Children	5
2	2.5	Recruitment and Retention Plan	5
2	2.7	Study Timeline	5
2	N/A	Inclusion Enrollment Report(s)	5, 9
3	3.1	Protection of Human Subjects	5
3	3.2	Single IRB Plan	5
3	3.3	Data and Safety Monitoring Plan	5
3	3.5	Overall Structure of the Study Team	5
4	4.4	Statistical Design and Power	7
4	4.6	FDA-regulated Intervention	7
4	4.7	Dissemination Plan	7
5	see FOA	See FOA for details	8



Inclusion Enrollment Report (Sample)

*1. Inclusion Enrollment Report Title _____

- 3. Enrollment Location Type Domestic Foreign
- 4. Enrollment Country(ies)
- 6. Comments (Up to 500 characters.)

Planned

			Ethnic Categories		
Racial Categories	Not Hispani	c or Latino	Hispanic	or Latino	Total
	Female	Male	Female	Male	
American Indian/ Alaska Native					
Asian					
Native Hawaiian or Other Pacific Islander					
Black or African American					
White					
More than One Race					
Total					



Inclusion Enrollment Report (Sample) (cont.)

Cumulative (Actual)

					Ethnic C	ategories				
Racial	Not Hispanic or Latino			Hispanic or Latino			Unknown/Not Reported Ethnicity			Total
Categories	Female	Male	Unknown/ Not Reported	Female	Male	Unknown/ Not Reported	Female	Male	Unknown/ Not Reported	
American Indian/ Alaska Native										
Asian										
Native Hawaiian or Other Pacific Islander										
Black or African American										
White										
More than One Race										
Unknown or Not Reported										
Total										

Report ____ of ____



Additional Interventions (if needed)

4.2.c.	Interventions	s (cont.)	
	(check one)	 Intervention Type: □ Drug (including placebo) □ Device (including sham) □ Biological/Vaccine □ Procedure/Surgery □ Radiation □ Behavioral (e.g., Psychotherapy, Lifestyle Counseling) 	 Genetic (including gene transfer, stem cell, and recombinant DNA) Dietary Supplement (e.g., vitamins, minerals) Combination Product Diagnostic Test Other
		Name (Up to 200 characters.)	
		Description	
		(Up to 1,000 characters.)	
4.2.c.	Interventions		
		Intervention Type: □ Drug (including placebo) □ Device (including sham) □ Biological/Vaccine	 Genetic (including gene transfer, stem cell, and recombinant DNA) Dietary Supplement (e.g.,
	(check one)	 Procedure/Surgery Radiation Behavioral (e.g., Psychotherapy, Lifestyle Counseling) 	vitamins, minerals) Combination Product Diagnostic Test Other
		Name (Up to 200 characters.)	
		Description	
		(Up to 1,000 characters.)	
4.2.c.	Interventions	s (cont.)	
		Intervention Type: □ Drug (including placebo) □ Device (including sham) □ Biological/Vaccine	 Genetic (including gene transfer, stem cell, and recombinant DNA) Dietary Supplement (e.g.,
	(check one)	 Procedure/Surgery Radiation Behavioral (e.g., Psychotherapy, Lifestyle Counseling) 	vitamins, minerals) Combination Product Diagnostic Test Other
		Name	
		Description	

(Up to 1,000 characters.)



Additional Outcome Measures (if needed)

4.3. Outcome Measures

(At least one Outcome Measure required, unless noted in the opportunity. Up to 50 Outcome Measures allowed.)

Name _____

1	(U	р	to	255	characters.)
---	----	---	----	-----	--------------

Туре

(check one)	Primary	Secondary	Other
(,	,	-

Time Frame

(Up to 255 characters.)

Brief Description

(Up to 999 characters.)

4.3. Outcome Measures

(At least one Outcome Measure required, unless noted in the opportunity. Up to 50 Outcome Measures allowed.)

Name (Up to 255 char	acters.)			
Type (check one)	□ Primary	□ Secondary	□ Other	
Time Frame (Up to 255 char				
Brief Descri (Up to 999 char				

4.3. Outcome Measures

(At least one Outcome Measure required, unless noted in the opportunity. Up to 50 Outcome Measures allowed.)

Name (Up to 255 char	racters.)		
Type (check one)	□ Primary	□ Secondary	□ Other
Time Frame (Up to 255 char			
Brief Descri (Up to 999 char			