

DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service Centers for Disease Control and Prevention (CDC)

Memorandum

Date December 4, 2020

From Kathy Masterson

IRB Administrator, NIOSH Institutional Review Board

Subject IRB Approval of Continuation of NIOSH Protocol 13-HELD-03XP, "Factors Influencing the

Transmission of Influenza" (Expedited)

To William G. Lindsley, PhD

Project Officer, HELD, NIOSH

The NIOSH IRB has reviewed and approved your request to continue protocol 13-HELD-03XP for the maximum allowable period of one year and it will expire on November 17, 2021. The protocol was reviewed in accordance with the expedited review process outlined in 45 CFR 46.110(b)(1), Categories (3), (4) and (7).

The IRB determined the study poses no more than minimal risk to subjects.

If other institutions involved in this protocol are being awarded NIOSH funds through the CDC Procurement and Grants Office (PGO), you are required to send a copy of this IRB approval to the CDC PGO award specialist handling the award. You are also required to verify with the award specialist the awardee has provided PGO with the required documentation and has approval to begin or continue research involving human subjects as described in this protocol.

As a reminder, the IRB must review and approve all human subjects research protocols at intervals appropriate to the degree of risk, but not less than once per year. There is no grace period beyond one year from the last IRB approval date. It is ultimately your responsibility to submit your research protocol for continuation review and approval by the IRB along with available IRB approvals from all collaborators. Please keep this approval in your protocol file as proof of IRB approval and as a reminder of the expiration date. To avoid lapses in approval of your research and the possible suspension of subject enrollment and/or termination of the protocol, please submit your continuation request along with all completed supporting documentation at least six weeks before the protocol's expiration date of November 17, 2021.

Any problems of a serious nature must be brought to the immediate attention of the NIOSH IRB, and any proposed changes to the protocol should be submitted as an amendment to the protocol for NIOSH IRB approval <u>before</u> they are implemented.

If you have any questions, please contact the CDC Human Research Protection Program (513) 533-8591 or e-mail: cin-hsrb@cdc.gov.

Centers for Disease Control and Prevention

NIOSH Institutional Review Board



Request for Subsequent Action of IRB-Approved Protocol

Use this form to submit a protocol for continuing review or amendment by a CDC IRB or a non-CDC IRB. [See 45 CFR 46.109(e).] See *HRPO Guide: Non-Exempt Review Cycle* for further details on how to complete this form.

CDC protocol ID: 13-F	IELD-03XP actors Influencing the Transn	Protocol version of Influenza		Version date <u>10/05/2</u> 018	
Continuing Review Review of changes	* ***	*Requesting transition to the 2018 Common Rule (*Optional) *Note: This may require changes to the study, including informed consent documents, to comply with the 2018 Common Rule			
2 Key CDC p	ersonnel				
No change in key (CDC personnel. If no changes	s, please list only th	e primary contact a	nd principal investigator.	
	Name and degrees (FirstName LastName, Degrees)	User ID	CITI Course Expiration Date	CDC CIO/division	
Primary contact (required)	William G. Lindsley, PhD	wdl7	04/01/2022	NIOSH/HELD	
Principal investigator (required)	William G. Lindsley, PhD	wdl7	04/01/2022	NIOSH/HELD	
Co-Investigator					
Co-Investigator					
Co-Investigator					
in current training can	nuired CITI training is expired result in removal from the states stigators. Include name and continue and co	udy or suspension o	of the study until red	quirements are met.	
y ⊠n CDC employed y □n CDC employed y □n CDC employed	• •	e identifiable (inclu e anonymous or unl chnical assistance o	ding coded) private inked data or biosp r oversight.	data or biospecimens.	

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4 Study Subjects	
Have any subjects been enrolled in the last 12 months?	⊠ no
Total number of subjects needed for study: 240 Total	I number of subjects enrolled to date: 118
Comments on sample size:	·
Additional subjects total will allow us to recruit about another 60 s	subjects with influenza and 60 healthy subjects
4.1 Contact status	
"Contact" means intervention or interaction with subjections consent, enrollment, and collection of data and biospection following.	0 0
 Study is not designed to involve research-related contact v records); study activities involve only access to or analysis Study is designed to involve contact with participants. Che 	s of data or biospecimens and writing reports.
Contact with subjects has not yet begun.	sek one of the following.
Contact with subjects has begun and continues; the notification of results.	his may include follow-up for debriefing or
Contact with subjects is completed; study activiti	es involve only data analysis or report writing.
4.2 Consent status	
	organization Charles and Alexander
"Consent" includes adult consent, child assent, and parental p	
The IRB previously waived all requirements both to obtain	•
Although not waived, there is no further need to obtain or o	
*Subjects will be asked to provide consent (with or withou	
* If you check the third box, please include all current consent (e.g., scripts, documents) from each study site with this submis	
5 Study status—overall conduct (This section [Comment 5.1] Summary of research activities to date. Briefly sum Include the number of potential subjects who declined enrollment and this study involves a registrable clinical trial, summarize registration	marize study progress and interim findings. and the number who withdrew from the study. If
In our most recent studies, we collected the aerosol particles product mask for 20 minutes. About a third of potential subjects declined to after enrollment. We intend to continue these studies with a similar mass spectroscopy and PCR rather than by culturing the virus.	enroll in the study. No participants withdrew
[Comment 5.2] Summary of study changes reviewed and approved s changes submitted with or before approval of this continuation but to None	

Request for subsequent action of IRB Approved Protocol

[Comment 5.3] Summary of any recent literature or other information relevant to the research study (not limited to information with CDC co-authorship).

Our work and work by other groups has shown that patients with influenza expel potentially infectious virus in airborne particles as they cough and breathe. Our focus is now shifting to analyzing the content of the airborne particles and understanding how this relates to the production of potentially infectious aerosols and the body's response to the influenza infection.

[Comment 5.4] Summary of all adverse events to date. In particular, address adverse events that were serious, unexpected (or more frequent or severe than expected), or at least possibly related to the research.

No adverse events have occurred.

[Comment 5.5] Summary of (a) incidents that are not adverse events and (b) other substantial concerns since last continuation.

No incidents or substantial concerns have arisen.

[Comment 5.6] List and include copies of progress or monitoring reports on safety or compliance (e.g., site monitor, safety review, DSM report, multi-center trial report, but not reports to PGO).

N/A

[Comment 5.7] Summary of remaining research activities, emphasizing future contact with subjects, use of identifiable private data and biospecimens, and preparation of primary reports.

Subjects will be enrolled in the study and tested using our new aerosol collection system. Additional studies will be prepared when we have significant results.

6 Regulation and policy

6.1 Vulnerable populations

Check one of the following:

- Change in vulnerable populations (added or dropped).
- No Change

6.2 Free and informed consent

Check one of the following:

- Change in consent process, forms, or approved waivers.
- **⋈** No Change

6.3 Other regulation and policy considerations

Check one of the following:

- Change in other regulation and policy considerations.
 - Exception to PHS policy regarding notification of HIV test results
 - Human genetic testing
 - Inclusion of a registrable clinical trial or change in registration status
 - Plans for long-term storage of identifiable biospecimens
 - Involvement of drug, biologic, or device, including Investigational New Drug or Investigational Device Exemption status (See *HRPO Worksheet to Determine FDA Regulatory Coverage* for guidance on whether or not FDA regulations apply.)
- **⋈** No Change

6.4 Confidentiality protections

Check one of the following:

- **Change** in confidentiality protections (e.g., granted, applied for, denied).
- **⋈** No Change

7 Summary of proposed changes

Describe and justify proposed modifications to the protocol. Include page numbers in reference to clean version and tracked version. Continue summary below in additional comments or in supplemental document if necessary.

8 Suggested Mode of IRB review on CDC's behalf

Location of IR	RB (check one):
CDC IRB	
Non-CDC	IRB through IRB authorization agreement [submit form 0.1371 if this is a new request]
Institution	or organization providing IRB review:
IRB regist	ration number:
_	ide assurance number:
Suggested lev	rel of risk to subjects (check one):
Proposed o	changes to protocol are minor
✓ Minimal	
Greater that	an minimal
_	
Suggested lev	rel of IRB review (check one):
	-board review is suggested
_	nvened review:
_	review is suggested, under the following categories (check all that apply):
1	Study of drugs not requiring Investigational New Drug exemption from FDA
1 b	Study of medical devices not requiring Investigational Device Exemption from FDA
2a	Collection of blood from healthy, non-pregnant adults; below volume limit, minimally invasive
2b	Collection of blood from other adults and children; below volume limit, minimally invasive
X 3	Prospective noninvasive collection of biological specimens for research purposes
× 4	Collection of data through routine, noninvasive procedures, involving no general anesthesia,
_	sedation, x-rays, or microwaves
<u> </u>	Research that uses materials collected solely for non-research purposes
<u> </u>	Collection of data from voice, video, digital, or image recordings made for research purposes
X 7	Research that uses interview, program evaluation, human factors, or quality assurance methods
~	
	ing review of research previously approved by the convened IRB where
■ 8a	The research is permanently closed to the enrollment of new subjects; all subjects have completed all research-related interventions; and the research remains active only for long-term
	follow-up of subjects
■ 8b	No subjects have been enrolled and no additional risks have been identified
■ 8c	The remaining research activities are limited to data analysis
1 9	Continuing review of research, not under IND/IDE, where categories 2 through 8
	do not apply but the IRB has determined and documented at a convened meeting
	that the research involves no greater than minimal risk and no additional risks
	have been identified

9 CDC's research partners

Research partners include *all* direct and indirect recipients of CDC funding (e.g., grants, cooperative agreements, contracts, subcontracts, purchase orders) other CDC support (e.g., identifiable private information, supplies, products, drugs, or other tangible support and collaborators who do not receive such support. Include current information on partners added or dropped since the last review using form 0.1370. Check one of the following.

- No research partners have been added since the last review.
- Research partners have been added and are listed on form 0.1370, which accompanies this form.
- One or more research partners no longer collaborate for this study, and are listed as follows:

10 Material submitted with this form

Check all that apply. Describe additional material in the comments section. Required items are indicated. Optional items may be requested by HRPO or the IRB.

Clean	Tracked	
X		Complete protocol
\times		Consent, assent, and permission documents or scripts
×		Other information for recruits or participants (e.g., ads, brochures, flyers, scripts)
X		Data collection instruments (e.g., questionnaires, interview scripts, record abstraction tools)
		Certification of IRB approval or exemption for supported/nonexempt partner
		Progress and monitoring reports (recommended when available)

11 Additional comments

Additional partners are listed on ancillary 1370 form

Partner 1 West Virginia University	Partner 2		
Institution name:	Institution name:		
Institution location: Morgantown, WV 26508	Institution location:		
Individual name (IIA only):	Individual name (IIA only):		
Reporting status: Previously Reported	Reporting status [Enter Status Here]		
Regulatory coverage Engaged/Non-Exempt	Regulatory coverage [Enter Status Here]		
Financial support No Financial Support	Financial support [Enter Status Here]		
Support award number:	Support award number:		
Support end date:	Support end date:		
Nonfinancial support: No Financial Support	Nonfinancial support [Enter Status Here]		
FWA number: <u>00005</u> 078	FWA number:		
HS Training (IIA only): Other Human Subject Training-Yes	HS Training (IIA only): [Enter Status Here]		
IRB review status: Relying on CDC IRB	IRB review status [Enter Status Here]		
IRB approval expiration date: 11/17/2020	IRB approval expiration date:		
Comments: Agreement 1372A in place KM	Comments:		
Partner 3	Partner 4		
Institution name:	Institution name:		
Institution location:	Institution location:		
Individual name (IIA only):	Individual name (IIA only):		
Reporting status [Enter Status Here]	Reporting status [Enter Status Here]		
Regulatory coverage: [Enter Status Here]	Regulatory coverage: [Enter Status Here]		
Financial support: [Enter Status Here]	Financial support [Enter Status Here]		
Support award number:	Support award number:		
Support end date:	Support end date:		
Nonfinancial support: [Enter Status Here]	Nonfinancial support: [Enter Status Here]		
FWA number:	FWA number:		
HS Training (IIA only): [Enter Status Here]	HS Training (IIA only): [Enter Status Here]		
IRB review status [Enter Status Here]	IRB review status [Enter Status Here]		
IRB approval expiration date:	IRB approval expiration date:		
Comments:	Comments:		
Partner 5	Partner 6		
Institution name:	Institution name:		
Institution location:	Institution location:		
Individual name (IIA only):	Individual name (IIA only):		
Reporting status: [Enter Status Here]	Reporting status [Enter Status Here]		
Regulatory coverage: [Enter Status Here]	Regulatory coverage: [Enter Status Here]		
Financial support: [Enter Status Here]	Financial support: [Enter Status Here]		
Support award number:	Support award number:		
Support end date:	Support end date:		
Nonfinancial support: [Enter Status Here]	Nonfinancial support: [Enter Status Here]		
FWA number:	FWA number:		
HS Training (IIA only): [Enter Status Here]	HS Training (IIA only): [Enter Status Here]		
IRB review status: [Enter Status Here]	IRB review status [Enter Status Here]		
IRB approval expiration date:	IRB approval expiration date:		
Comments:	Comments:		

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12 Signatures

As principal investigator, I hereby accept responsibility for conducting this CDC-sponsored research project in an ethical manner, consistent with the policies and procedures contained in CDC's *Procedures for Protection of Human Research Participants*, and to abide by the principles outlined in federal policies for the protection of human subjects at 45 CFR part 46, 21 CFR part 50, and 21 CFR part 56.

	Date	Kemarks
Signature		
Principal CDC Investigator:		
William G. Lindsley - Digitally signed by William G. Lindsley -S Date: 2020.11.16 14:21:50 -05'00'		

As a supervisor of the principal investigator, I hereby accept responsibility for ensuring that this CDC-sponsored research project is conducted in an ethical manner, consistent with the policies and procedures contained in CDC's *Procedures for Protection of Human Research Participants*, and to abide by the principles outlined in federal policies for the protection of human subjects at 45 CFR part 46, 21 CFR part 50, and 21 CFR part 56.

	Date	Remarks
Signature		
Team Lead:		Check if PI is Team Lead:
John D. Noti -S Digitally signed by John D. Noti -S Date: 2020.11.16 14:08:00 -05'00'		
Branch Official (e.g., Chief or Senior Scientist):		Check if PI is Branch Official:
John D. Noti -S Digitally signed by John D. Noti -S Date: 2020.11.16 14:08:25 -05'00'		
Division Official (e.g., Director or ADS):	4.4.4.0.100.00	Check if PI is Division Official:
Paul D. Siegel -S Digitally signed by Paul D. Siegel -S Date: 2020.11.16 14:48:11 -05'00'	11/16/2020	