

U.S. Department of Health and Human Services Public Health Service Final Progress Report Instructions

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A. Final Progress Report Requirement and Submission Information

A final progress report is required for any grant that is terminated and any award that will not be extended through award of a new competitive segment. The report is due within 90 of the end of the project period. If a competitive renewal (Type 2) application has been submitted, whether funded or not, the progress report contained in that application may serve in lieu of a separate final progress report at the discretion of the funding Institute/Center (IC). Otherwise, a final progress report should be prepared in accordance with the requirements below and any specific requirements set forth in the terms and conditions of the award.

There is no form page for the final progress report. At the top of the first page provide the grant number, project title, name of grantee organization, project period (start and end dates), name of the PD/PI, and clearly indicate "Final Progress Report."

All grantees are strongly encouraged to submit the final progress report electronically through the eRA Commons at https://commons.era.nih.gov/commons/. See the eRA Commons User Guide, section 9.11 Closeout. Additional information on electronic submission of closeout documents is available at the NIH eRA Commons homepage or by contacting the eRA help desk at: http://ithelpdesk.nih.gov/eRA/ or Toll-free (866) 504-9552, Phone 301-402-7469, TTY 301-451-5939.

If not submitted electronically through the eRA Commons, the original final progress report should be submitted to the centralized mailing address at:

Division of Extramural Activities Support, OER National Institutes of Health 6705 Rockledge Drive, Room 2207, MSC 7987 Bethesda, MD 20892-7987 (for regular or US Postal Service Express mail)

Bethesda, MD 20817 (for other courier/express mail delivery only)

Phone Number: (301) 594-6584

If submitted via paper to the centralized mailing address, the report should contain the signature of a Signing Official/Authorized Organization Representative.

Additional information on submitting closeout documents to AHRQ, CDC, FDA and IHS can be obtained from their websites.

B. Instructions for All Final Progress Reports (exclusive of SBIR/STTR Phase II Final Progress Reports)

There is no form page for the final progress report. At the top of the first page provide the grant number, project title, name of grantee organization, project period (start and end dates), name of the PD/PI, and clearly indicate "Final Progress Report."

The final progress report should include a summary of progress made toward the achievement of the originally stated aims, a list of significant results (positive or negative), and a list of publications. Grantees should also report additional information required by the awarding IC in program-specific final progress report instructions. The final progress report also should address the following when applicable:

- 1. Report on the final enrollment data for study subjects based on sex/gender, race, and ethnicity (use the Inclusion Enrollment Report).
- 2. If appropriate, indicate whether children were involved in the study or how the study was relevant for conditions affecting children (see Public Policy Requirements and Objectives—Inclusion of Children as Subjects in Clinical Research).

- 3. Describe any data, research materials (such as cell lines, DNA probes, animal models), protocols, software, or other information resulting from the research that is available to be shared with other investigators and how it may be accessed. If the initial research plan addressed, or the terms of award require, a formal plan for sharing final research data, model organisms, Genome Wide Association Studies data, or other such project-specific data, provide a final statement on the implementation of that plan.
- 4. Publications that were authored or co-authored by the PD/PI and arose from the award must include the NIH Manuscript Submission reference number (e.g., NIHMS97531) or the PubMed Central (PMC) reference number (e.g., PMCID234567) for each article. If the PMCID is not yet available because the Journal submits articles directly to PMC on behalf of their authors, indicate "PMC Journal In Process." A list of these Journals is posted at: http://publicaccess.nih.gov/submit_process_journals.htm.
- 5. Any other specific requirements set forth in the terms and conditions of the award must also be addressed in the final progress report.

C. Instructions for SBIR/STTR Phase II Final Progress Reports

Final reports serve as an important source of material for staff of the IC in preparing annual agency reports, for planning purposes, for tracking program outcomes, and in communicating scientific accomplishments achieved through the SBIR/STTR program. There is no form page for a final SBIR/STTR report, but the format below is strongly recommended and is available as a fillable MS Word file at: http://grants.nih.gov/grants/funding/finalprogressreport_SBIR_PhaseII.doc. All 15 items, plus requested attachments, should be provided. If uploaded through the Commons all documents must be combined into a single pdf.

- 1. Provide the grant number, project title, name of grantee organization, project period (start and end dates), and name of the PD/PI.
- 2. If the company has undergone a recent name change provide the new name.
- 3. Provide a summary of the specific aims and impact on public health of the Phase II grant. (limit 1,300 characters)
- 4. Provide a succinct account of published and unpublished results, indicating progress toward achievement of the originally stated aims.
- 5. List patents (U.S. and international), copyrights, trademarks, and invention reports, if any, that resulted from the award.

	# Filed (Enter Numeric Value)	# Approved (Enter Numeric Value)	Patent Numbers (separated by commas)
Patents			
Copyrights			
Trademarks			
Invention Reports			

Describe other printed materials or demonstration of IP protection, if any, that resulted from the award. (limit 500 characters)

6. Check all boxes below that best describe the technology developed from this SBIR/STTR.

	Small Molecules: The development or reformulation of drugs as chemical substances used in the treatment, cure, prevention, or diagnosis (<i>in vivo</i> , imaging agents, etc) of disease or used to otherwise enhance physical or mental well-being; includes so-called "naturopathic" or naturally-derived substances in alternative care regimes.
	Biologics: A medicinal product created by biologic processes, such as a vaccine, blood or blood component, allergenic, somatic cell, gene therapy, tissue, recombinant therapeutic protein, or living cells.
	Companion Product: A diagnostic, therapeutic, or device that must be used in combination with another diagnostic, therapeutic, or device type (e.g. companion diagnostic for a specific therapy; a small molecule that activates expression from a gene therapy vector; a device and imaging agent that work together). This does not include "drug cocktails." The Phase II project may include only one aspect of the companion product.
	Medical Devices: The development and/or use of instruments or machines, used in the diagnosis of disease or in the cure, mitigation, treatment, or prevention of disease or conditions associated with the deterioration of physiological function (e.g., prostheses); this would also include medical imaging devices and the use of innovative materials to construct new devices.
	Research Tools: The development of new or improved tools, devices, and sensors to enhance laboratory or field studies on humans, animals, or any model system. This includes tools to broaden the research knowledge base and for biomonitoring.
	Biotechnology: The use of microorganisms, such as bacteria or yeasts, to perform specific industrial or manufacturing processes.
	<i>In Vitro</i> and <i>Ex Vivo</i> Diagnostics: The use of tools (software, hardware or combinations) to identify or screen for medical conditions and determine whether specified diseases or disease processes are present in living organisms. Includes the use of these tools for non-clinical screenings and to provide insights in the work of clinicians, providers, manufacturers of equipment, and companies involved in therapies associated with disease.
	Healthcare IT: Approaches and tools derived from information technology that allow for the management of research, educational and medical information. Includes software, media, educational tools, and digital health.
	Other, please specify. (limit 500 characters)
	cribe the technology's intended commercial application, potential market size, and who will use it. (limit characters)
7.	Check the box that best describes the current R&D status of the product.
	Non-clinical technology in prototype development/testing stage
	Non-clinical technology in full development/testing stage
	Pre-clinical development
	Clinical development
	Commercially available
	Discontinued
	Other (limit 500 characters)
Des	cribe the current status of this product and explain reasons if discontinued. (limit 500 characters)
8.	Check the boxes that best describes the regulatory approval status for your product, process, or service.
(Cho	eck all that apply)
	Not applicable (no regulatory approval needed)

FDA	<u>approval</u> :						
PMA	A	☐ Not yet subr	nitted	Submitted	☐ Approved	Rejected	
510(k) Not yet subn		nitted	Submitted	Approved	Rejected		
IDE Not yet subn			nitted	Submitted	Approved	Rejected	
BLA	L	☐ Not yet subr	nitted	Submitted	Approved	Rejected	
IND		☐ Not yet subr	nitted	Submitted	Approved	Rejected	
NDA	A	☐ Not yet subr	nitted	Submitted	Approved	Rejected	
FDA	Facility Registrations	☐ Not yet subr	nitted	Submitted	Approved	Rejected	
EU/	<u>UK approval</u> :						
CE I	Mark	☐ Not yet subr	nitted	Submitted	☐ Approved	Rejected	
	Other regulatory submission including any foreign subm	* *		•	ubmitted regulator	y applications,	
9.	Check the boxes that best of	describe the reim	bursemer	nt approval status of	the product, proce	ess, or service.	
(Che	eck all that apply)						
	Not applicable						
CMS	S Reimbursement	☐ Not yet subr	nitted	Submitted	☐ Approved	Rejected	
Priva	ate Payer Reimbursement	☐ Not yet subr	nitted	Submitted	Approved	Rejected	
10.	Check the boxes that best of	describe the statu	s of clini	cal trials for your pro	oduct, process, or	service.	
(Che	eck all that apply)						
	Not applicable						
Phas	e I clinical trial		Ongo	oing	Completed	l	
Phas	e II clinical trial		Ongo	oing	Completed	l	
Phas	e III clinical trial		Ongo	oing	Completed	l	
Premarket approval (PMA) device trial			Ongo	oing	Completed	l	
Phas	e IV Postmarketing study		Ongo	oing	Completed	l	
Outs	ide of the United States (O	US)	Ongo	oing	Completed	1	
11. Describe company outcomes occurring, at least in part, as a result of this award.							
(Che	eck all that apply)						
	Follow on funding		Total c	umulative dollar amo	ount \$		
(check all that apply and enter amount invested)							
	Venture Capital (VC)		Total co	umulative dollar amo	ount		
	Angel		Total co	umulative dollar amo	ount		
	state/Local		Total co	umulative dollar amo	ount		
Strategic partnership				Total cumulative dollar amount			
☐ Federal				umulative dollar amo	ount		
☐ Internal SBC Funds			Total cumulative dollar amount				

Other (Foundations, bar	nk loans, etc)	Total cumu	lative dollar amount	
Out-licensing agreemen	ts/sale of IP	Number		
			lative dollar amount	
			greement	
☐ In-licensing agreements		Number		
_			lative dollar amount	
		Nature of a	greement	
☐ Strategic partnership/s t	hat do not include fu	ınding		
		Name(s)		
☐ Spin-off companies				
☐ Public offering				
		Year		
		Value		
☐ Merger or acquisition of	f Awardee	Name of ac	equirer	
		Year		
		Total value		
			tcomes attributable to the award, incl I nature of significant partnerships, it	
12. Describe the sales or refunds).	evenues, if any, whi	ch resulted fro	om this SBIR/STTR award (not inclu	ding award
No sales or revenue to	date.			
Please provide projected da	te of first sale/comn	nercial service	launch in MM/DD/YYYY:	
Sales or service to:				
(check all that apply and en	ter the total cumulat	ive dollar am	ount to date)	
Federal				
Private sector				
Other				
List the generic and/or comleast in part, from this awar			rocess(es), or service(s), if any, that is er of products sold.	esulted, at
* If the SBIR/STTR-support revenues of both the compo	•		rger commercial product, please list	the sales
Product or Service	Revenues Generate	ed	Number Sold (if applicable)	
	l			

Product or Service	Revenues Generated	Number Sold (if applicable)

13.	List titles and complete references to publications, and manuscripts accepted for publication, if any, that
	resulted from the Phase II award. When citing articles that fall under the Public Access Policy, provide the
	NIH Manuscript Submission reference number (e.g., NIHMS97531) or the PubMed Central (PMC)
	reference number (e.g., PMCID234567) for each article. If the PMCID is not yet available because the
	Journal submits articles directly to PMC on behalf of their authors, indicate "PMC Journal - In Process." A
	list of these Journals is posted at: http://publicaccess.nih.gov/submit_process_journals.htm .

14.	Provide the current number of employees (total full time equivalents or FTEs):
	Provide the number of FTEs directly supported by this award:
	Provide an estimate of the total number of FTEs attributable to all previous and current SBIR/STTR funding received:

15. Attach the Inclusion Enrollment Report from the competing application instructions, with the final enrollment data for clinical research.