

AWARD NUMBER: W81XWH-16-1-0098

TITLE: Antilyosphosphatidic Acid Antibodies in the Treatment of Post-TBI Neuropathic Pain

PRINCIPAL INVESTIGATOR: David C. Yeomans, PhD

CONTRACTING ORGANIZATION: Stanford University
Palo Alto, CA 94304

REPORT DATE: APRIL 2019

TYPE OF REPORT: Annual Report

PREPARED FOR: U.S. Army Medical Research and Materiel Command
Fort Detrick, Maryland 21702-5012

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REPORT DOCUMENTATION PAGE

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1. REPORT DATE (DD-MM-YYYY)	2. REPORT TYPE	3. DATES COVERED (From - To)
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	5b. GRANT NUMBER
	5c. PROGRAM ELEMENT NUMBER

6. AUTHOR(S) EMAIL: dcyemans@stanford.edu	5d. PROJECT NUMBER
	5e. TASK NUMBER
	5f. WORK UNIT NUMBER

7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES)	8. PERFORMING ORGANIZATION REPORT NUMBER
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9. SPONSORING/MONITORING AGENCY NAME(S) AND ADDRESS(ES) Fort Detrick, MD 21702-5012	10. SPONSOR/MONITOR'S ACRONYM(S)
	11. SPONSOR/MONITOR'S REPORT NUMBER(S)

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13. SUPPLEMENTARY NOTES

14. ABSTRACT

15. SUBJECT TERMS

16. SECURITY CLASSIFICATION OF:			17. LIMITATION OF ABSTRACT	18. NUMBER OF PAGES	19a. NAME OF RESPONSIBLE PERSON	
a. REPORT	b. ABSTRACT	c. THIS PAGE			19b. TELEPHONE NUMBER (Include area code)	

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INTRODUCTION

Lysophosphatidic acid (LPA) is a lipid inflammatory mediator that is released following nerve injury, including injury to the brain. LPA has been implicated in the development and maintenance of pain and other deleterious sequelae to brain injury. The broad, long-term goal of this project is to evaluate the therapeutic potential of a novel humanized anti-LPA antibody (Lpathomab) for attenuating post-traumatic brain injury (TBI) associated pain. Thus, we are studying the effect of Lpathomab, comparing the utility of nasal, intravenous, and subcutaneous administration, in preventing long-term pain sequelae due to fluid percussion-induced neurotrauma in rats.

This grant was originally awarded to LPath, Inc, a small biotech in San Diego, with Roger Sabbadini as principle investigator. Stanford University was the primary contractor, with David Yeomans, a Stanford Professor, as Co-PI. Unfortunately, during the reporting period, LPath went out of business, and Dr. Yeomans and Sabbadini requested transfer of the grant to Stanford, and PI-ship to Dr. Yeomans. This request was made in September of 2016 to transfer the grant as of November, 2016. However, Stanford did not receive a notice of allowance until March, 2018. Meanwhile, LPath went out of business as of January, 2017 – and all LPath email addresses became dead. Thus, if notices were sent to LPath for submitting an annual report, these were not received. In the mean time, Dr. Yeomans and Dr. Sabbadini have continued to work on the project and file quarterly reports with the idea that the funds would be transferred. The transfer notice is included in the appendix.

KEY WORDS

monoclonal antibody, lipid inflammatory mediator, analgesia, lysophosphatidic acid

ACCOMPLISHMENTS

Goals:

Traumatic brain injury (TBI) is the most common major injury suffered by warfighters in Iraq and Afghanistan, and frequency results in multiple symptoms of which chronic pain is the most common. Reviews have placed the prevalence of chronic pain following TBI at 88% in blast exposed soldiers with TBI in OIF/OEF and 65% in non-blast exposed soldiers with TBI from the same conflicts (Girona, 2009). Another general review of military veterans' medical charts places the prevalence of pain complaints at 50% in the general VA population (Clark, 2002) with 62.6% of TBI of patients reported taking narcotic analgesics (Ponsford, 2011). Lysophosphatidic acid (LPA), a bioactive lipid which is known to be increased in TBI and block of LPA effects using an anti-LPA antibody has been shown to provide neural protection after TBI and to block pain after peripheral nerve injury. The broad, long-term goal of this project is to evaluate the therapeutic potential of a novel anti-LPA antibody (Lpathomab) for attenuating post-TBI pain. Thus, we propose to study the effect of Lpathomab, using 3 different application procedures, in preventing long-term pain sequelae due to fluid percussion-induced neurotrauma.

Because of the availability of the specialized expertise for nasal administration in rats, we altered the order such that we accomplished and published the results of Specific Aim 4 of the original proposal began Specific Aim 1 of the revised SOW.

Task 2 (6-12 months): Analysis of LPA in cerebral spinal fluid (CSF). Concentration of LPA in CSF of sham and cFP TBI rats was evaluated by ELISA assay. Two groups of animals, including 12 cFP TBI rats and 12 sham cFP rats were anesthetized and surgically prepared for percussion injury. The 12 TBI rats underwent percussion injury wherein the surface of the cortex is exposed to a pressure wave, the remaining 12 rats underwent sham surgery wherein the cortex was exposed, but no pressure wave was introduced. Thirty minutes following TBI, and while still anesthetized, the atlanto-occipital membrane was surgically exposed and a needle introduced through the membrane for extraction of CSF which were frozen for off-line analysis by ELISA.

Results for Task 2: CSF levels of LPA following TBI

After mild TBI, rats demonstrated significantly ($p < 0.05$, Student's t test) higher levels of LPA when compared to rats that had undergone sham procedures. Average CSF concentration of LPA 30 minutes after TBI was 4.5 (+/- 0.3 SEM) micromolar vs 1.1 (+/- 0.2 SEM) micromolar for sham rats. These values are consistent with values we previously found for mice that had undergone nerve injury where we observed a 2.8X increase and with where we observed a 5.4 X increase at 24 hours after TBI in humans (Crack et al., 2014).

Specific Aim 1: I.V Administration of Lpathomab anti-LPA antibody

The experiment has four levels (dosages) of Lpathomab (2.5, 25, 50, and 100 mg/kg) and several types of controls. Sham surgery served as an injury control and injection of a matched control antibody served as an Lpathomab treatment control.

Task 4: I.V. Lpathomab Administration and Neurobehavioral assessment

Neurologic function by standard assessment and pain sensitivity by mechanical stimulation and thermal (laser) stimulation were assessed at various times after cFP TBI or sham injury and after I.V. application of Lpathomab or a control antibody. Thus, a dose-response relationship was determined in terms of analgesic efficacy.

Thirty minutes after percussion injury (for TBI rats), or suture closing (for sham rats), animals were briefly anesthetized with inhaled isoflurane. Anesthetized rats were then administered an intravenous (tail vein) injection of Lpathomab, an equivalent amount of matched antibody, or a control solution. Lpathomab dosage levels were based on values found to reduce pain symptoms following neuropathic injury in rats.

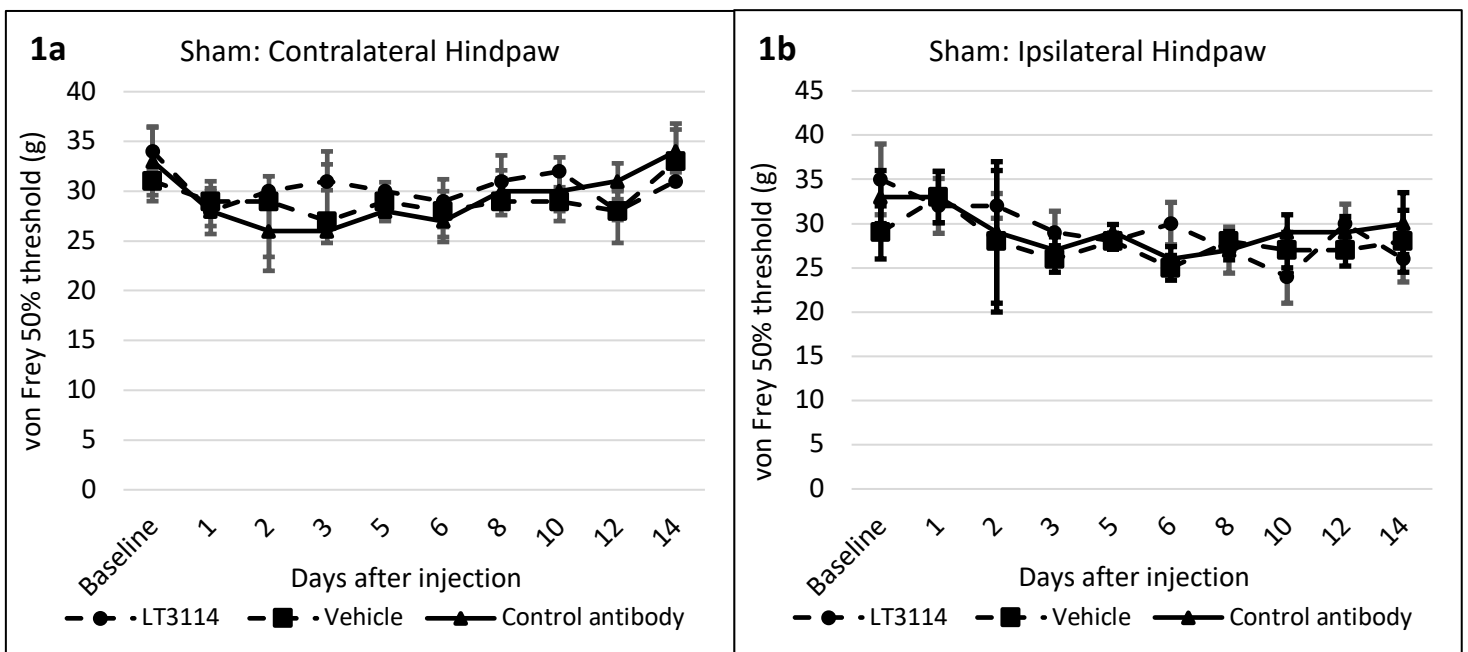
A. Neurological severity score

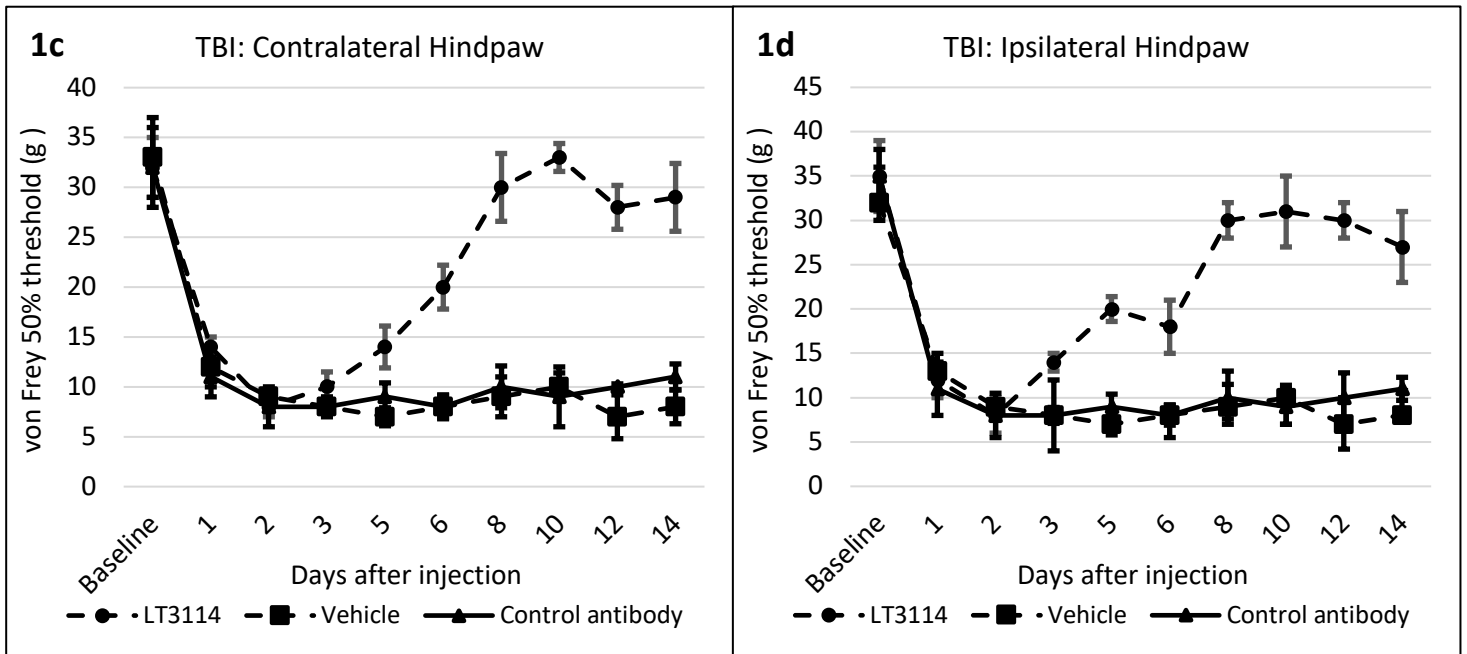
Neurological status after TBI was assessed using a standard neurological severity score described using a 12-level test that assesses, e.g., righting reflex, hemiplegia, and loss of pinna reflex and has been shown to correlate closely with the applied pressure. After TBI, all rats demonstrated no or only mild signs of neurological impairment. Thus, no rats had to be excluded because of preset parameters (elevated neurological severity score [>0]) after 1 week or with delayed emergence (>4 minutes after TBI), which would have been indicative of higher grade TBI.

B. Mechanical Allodynia

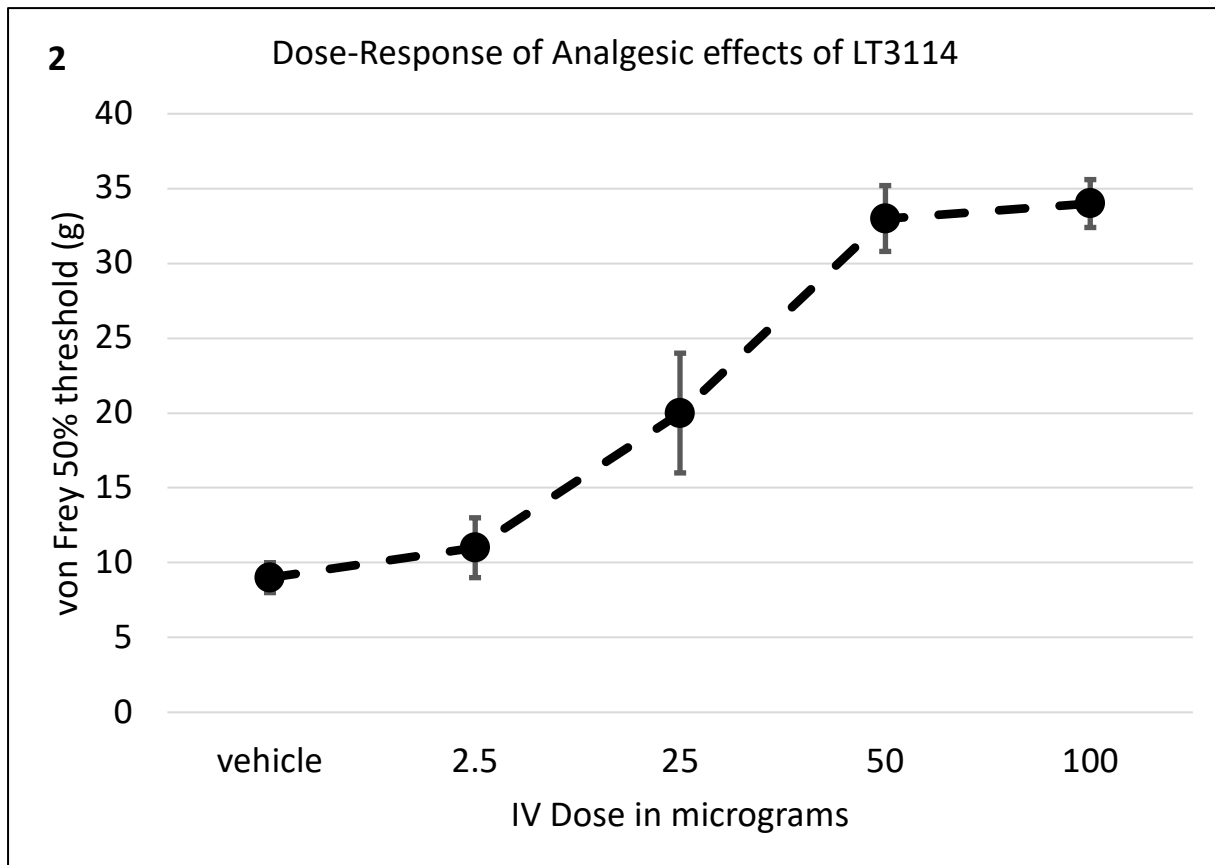
Before the first testing session, animals (N = 6 per group) were habituated for at least 2 hours on a metal mesh (0.5 x 0.5 cm) inside a plastic chamber. One day before TBI, rats were tested for mechanical sensitivity using von Frey filaments (Bioseb, Chaville, France) applied to the plantar surfaces of the hind paws. Fifty percent mechanical withdrawal thresholds to the application of a von Frey probe to the foot was calculated by using the up-down method. To assess mechanical allodynia, an ascending series of von Frey hairs of logarithmically incremental force (0.32-16.31 g) was applied to the midregion of the plantar surface of the hind paw. Each von Frey hair was applied to the test area for 2 to 3 seconds, with a 1- to 2-minute interval between stimuli. Testing was performed on post-TBI day 1, 2, 3, 5, 6, 8, 10, 12, and 14.

Figures 1a and b shows that there was no change in mechanical withdrawal thresholds for either hindpaw following sham TBI. In addition, no effect was seen on thresholds after IV administration of 100 μ g of LT3114, control antibody, or vehicle. On the other hand, rats exposed to true TBI showed robust, bilateral mechanical allodynia (Figures 1 c and d) as demonstrated by a significant ($p < 0.05$) decrease in withdrawal thresholds. However, unlike rats that received injections of control antibody or vehicle, rats that received an injection of 100 μ g of LT3114 recovered to their pre-TBI levels of mechanical sensitivity over 8 to 10 days following injection. This recovery of normal sensitivity was maintained at least through 14 days.





These anti-allodynic effects were dose-dependent, as is made clear by Figure 2, which represents the average (+/- SEM) von Frey thresholds at 10 days following LT3114 injection.



IMPACT

What was the impact on development of the principal discipline(s) of the project?

The results of our work demonstrated that LPA levels are elevated in CSF following TBI – indicating the presence of the target molecule in the target tissue. In addition, these experiments demonstrated that IV application of an antibody can result in decreased pain sensitivity following TBI. Thus, these results give additional confidence that IV or intranasal application of anti-LPA antibody would be useful for the treatment of post-TBI pain.

What was the impact on other disciplines?

These results can impact other disciplines in that they show that Intranasal and intravenous application of an antibody can have therapeutic effects.

What was the impact on technology transfer?

Nothing to report at this time.

What was the impact on society beyond science and technology?

Nothing to report at this time.

CHANGES/PROBLEMS

Changes in approach and reasons for change

We have changed the order of the tasks secondary to problems that arose during the previous reporting period.

Actual or anticipated problems or delays and actions or plans to resolve them

We have run into a serious problem with the reliability of the rat TBI model. We believe that we have fixed this problem, but it is still possible that additional delays will occur because of this.

Changes that had a significant impact on expenditures

We have delayed initiating the next round of experiments while we determine the cause of the problem with reliability. This has caused a delay during which we have used both salary and animals while we try to figure out the issue.

Significant changes in the use of care of human subjects, vertebrate animals, biohazards, and/or select agents

None to report at this time.

PRODUCTS

Publications, conference papers, and presentations

We have published our work in the leading pain journal, *Pain*:

Eisenried A, Meidahl ACN, Klukinov M, Tzabazis AZ, Sabbadini RA, Clark JD, Yeomans DC.

Nervous system delivery of antilyso-phosphatidic acid antibody by nasal application attenuates mechanical allodynia after traumatic brain injury in rats. *Pain*. 2017 Nov;158(11):2181-21.

Website(s) and other Internet site(s)

None to report at this time.

Technologies or techniques

None to report at this time.

Inventions, patent applications, and/or licenses

None to report at this time.

Other Products

None to report at this time.

PARTICIPANTS AND OTHER COLLABORATING ORGANIZATIONS

Name	Roger Sabbadini, PhD
Project Role	Principal Investigator
Researcher Identifier	
Nearest person month worked	3.0
Contribution to Project	Provides overall direction to the project
Funding Support	LPath Inc.

Name	David C. Yeomans, PhD
Project Role	Co-Principal Investigator
Researcher Identifier	0000-0002-9389-8539
Nearest person month worked	3
Contribution to Project	Provides overall direction to the Stanford component of the research
Funding Support	NIH, DOD

Name	Michael Nemenov, PhD
Project Role	Collaborator
Researcher Identifier	
Nearest person month worked	2.4
Contribution to Project	Provided expertise around assessment of effects of TBI on thermal pain sensitivity
Funding Support	NIH

Name	Lena Weber, MD
Project Role	Postdoctoral Fellow
Researcher Identifier	
Nearest person month worked	12
Contribution to Project	Performed animal surgery and testing procedures
Funding Support	Stanford University

Name	Mikhail Klukinov, MD
Project Role	Senior Research Scientist
Researcher Identifier	
Nearest person month worked	7.08
Contribution to Project	Helped with behavioral testing of animals, ran biochemical analyses
Funding Support	Stanford University

No change in active support of key personnel
 No other organizations were involved as partners

SPECIAL REPORTING REQUIREMENTS

An updated quad chart is included as Appendix 1

APPENDICES

Appendix 1: updated quad chart

Appendix 2: Approved transfer - amendment of solicitation/modification of contract

Anti-Lysophosphatidic Acid Antibodies in the Treatment of Post-TBI Neuropathic Pain

MR141271 Task Title: Task 6: Test effects of Intranasal application of LPA antibody in TBI rats

W81XWH-16-1-0098

PI: David C. Yeomans

Org: Stanford University Award Amount: \$1,446,655.00

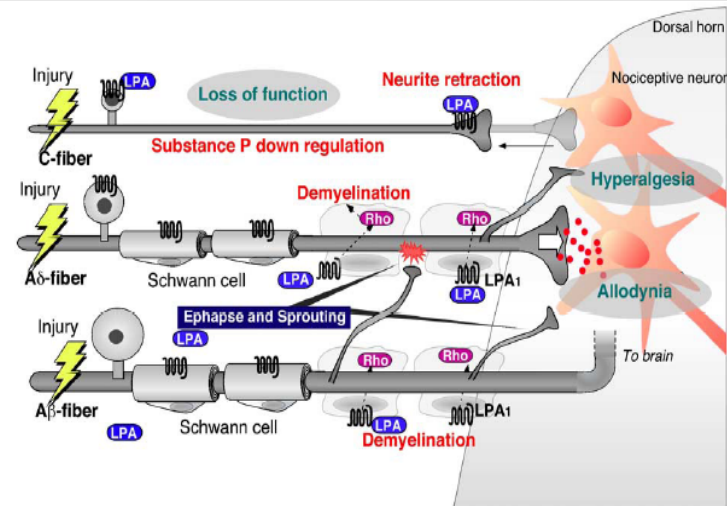


Study/Product Aims

- **Specific Aim 1:** Investigate the efficacy of intravenous (I.V.) injection of humanized anti-LPA mAb (LT3114, Lpathomab-h) in the central fluid percussion (cFP) injury model of TBI in the rat by measuring several neuropathic pain behaviors following cFP TBI.
- **Specific Aim 2:** To determine the optimal mode of administration by comparing effects of intravenous (I.V.) administration, garnered from Specific Aim 1, to the effects of subcutaneous (S.C.) or intranasal (I.N.) dosing in preventing or ameliorating post-injury pain as well as cognitive deficits.

Approach

Evaluate the efficacy of anti-LPA antibody for protection against post-TBI neuropathic pain: Rats will subjected to cFP injury followed by anti-LPA antibody treatment and we will correlate with mechanical allodynia and thermal hyperalgesia as well as other neurobehavioral measures



Interfering with LPA signaling will prevent white matter damage and improve pain responses. H. Ueda / Pharmacology & Therapeutics 109 (2006) 57–77

Timeline and Cost

Activities by CY	2016	2017	2018	2019
Investigate the efficacy of Lpathomab in cFP model of post-TBI neuropathic pain		[Bar chart showing activity from 2017 to 2019]		
Determine best route of administration (sc, iv, intranasal) and therapeutic time window		[Bar chart showing activity from 2017 to 2019]		
Submit FDA and IRB approval for TBI pain as an extension of Lpath's anticipated neuropathic pain IND				[Bar chart showing activity in 2019]
Estimated Total Budget (\$K, direct + indirect)	212	428	513	294

Updated: 04/29/2019

CY16 Milestones accomplished

- ✓ Put animal protocols in place
- ✓ Train personnel and conduct pilot in vivo studies for model development

CY17 Goals

- ✓ Analyze of LPA in cerebral spinal fluid (CSF) after injury
- ✓ Investigate I.N. Lpathomab Administration and Neurobehavioral assessment
- ✓ Investigate I.V. Lpathomab Administration and Neurobehavioral assessment

CY18 Goals

- Investigate S. C. Lpathomab Administration and Neurobehavioral assessment


CY19 Goals

- ✓ Optimize I. N Lpathomab Administration and Neurobehavioral assessment

- Determination of optimal administration method

Budget Expenditure to Date

Actual Expenditure: \$1,155,760.01

AMENDMENT OF SOLICITATION/MODIFICATION OF CONTRACT			1. CONTRACT ID CODE S	PAGE OF PAGES 1 16
2. AMENDMENT/MODIFICATION NO. P00001	3. EFFECTIVE DATE 08-Mar-2018	4. REQUISITION/PURCHASE REQ. NO. 0010768265-0001		5. PROJECT NO.(If applicable)
6. ISSUED BY USA MED RESEARCH ACQ ACTIVITY 820 CHANDLER ST FORT DETRICK MD 21702-5014	CODE W81XWH	7. ADMINISTERED BY (If other than item 6) See Item 6		
8. NAME AND ADDRESS OF CONTRACTOR (No., Street, County, State and Zip Code) LELAND STANFORD JUNIOR UNIVERSITY, THE STANFORD UNIVERSITY 450 SERRA MALL STANFORD CA 94305-2004		9A. AMENDMENT OF SOLICITATION NO.		
		9B. DATED (SEE ITEM 11)		
		X 10A. MOD. OF CONTRACT/ORDER NO. W81XWH-16-1-0098		
		X 10B. DATED (SEE ITEM 13) 15-Apr-2016		
CODE 1KN27	FACILITY CODE			
11. THIS ITEM ONLY APPLIES TO AMENDMENTS OF SOLICITATIONS				
<input type="checkbox"/> The above numbered solicitation is amended as set forth in Item 14. The hour and date specified for receipt of Offer <input type="checkbox"/> is extended, <input type="checkbox"/> is not extended. Offer must acknowledge receipt of this amendment prior to the hour and date specified in the solicitation or as amended by one of the following methods: (a) By completing Items 8 and 15, and returning _____ copies of the amendment; (b) By acknowledging receipt of this amendment on each copy of the offer submitted; or (c) By separate letter or telegram which includes a reference to the solicitation and amendment numbers. FAILURE OF YOUR ACKNOWLEDGMENT TO BE RECEIVED AT THE PLACE DESIGNATED FOR THE RECEIPT OF OFFERS PRIOR TO THE HOUR AND DATE SPECIFIED MAY RESULT IN REJECTION OF YOUR OFFER. If by virtue of this amendment you desire to change an offer already submitted, such change may be made by telegram or letter, provided each telegram or letter makes reference to the solicitation and this amendment, and is received prior to the opening hour and date specified.				
12. ACCOUNTING AND APPROPRIATION DATA (If required)				
13. THIS ITEM APPLIES ONLY TO MODIFICATIONS OF CONTRACTS/ORDERS. IT MODIFIES THE CONTRACT/ORDER NO. AS DESCRIBED IN ITEM 14.				
A. THIS CHANGE ORDER IS ISSUED PURSUANT TO: (Specify authority) THE CHANGES SET FORTH IN ITEM 14 ARE MADE IN THE CONTRACT ORDER NO. IN ITEM 10A.				
B. THE ABOVE NUMBERED CONTRACT/ORDER IS MODIFIED TO REFLECT THE ADMINISTRATIVE CHANGES (such as changes in paying office, appropriation date, etc.) SET FORTH IN ITEM 14, PURSUANT TO THE AUTHORITY OF FAR 43.103(B).				
C. THIS SUPPLEMENTAL AGREEMENT IS ENTERED INTO PURSUANT TO AUTHORITY OF:				
X D. OTHER (Specify type of modification and authority) Award Transfer, USAMRAA Terms and Conditions				
E. IMPORTANT: Contractor <input checked="" type="checkbox"/> is not, <input type="checkbox"/> is required to sign this document and return _____ copies to the issuing office.				
14. DESCRIPTION OF AMENDMENT/MODIFICATION (Organized by UCF section headings, including solicitation/contract subject matter where feasible.) Modification Control Number: cmeinber181789 Effective this date; this award is hereby transferred: FROM Lpath, Inc. 4025 Sorrento Valley BLVD San Diego, CA 92121-1404 TO: The Leland Stanford Junior University 3172 Porter Dr. Palo Alto, CA 94304-1212				
Except as provided herein, all terms and conditions of the document referenced in Item 9A or 10A, as heretofore changed, remains unchanged and in full force and effect.				
15A. NAME AND TITLE OF SIGNER (Type or print)		16A. NAME AND TITLE OF CONTRACTING OFFICER (Type or print) JASON KUHN / CONTRACTING OFFICER TEL: 301-619-1861 EMAIL: jason.d.kuhns.civ@mail.mil		
15B. CONTRACTOR/OFFEROR	15C. DATE SIGNED	16B. UNITED STATES OF AMERICA BY 	16C. DATE SIGNED 07-Mar-2018	
(Signature of person authorized to sign)		(Signature of Contracting Officer)		

SECTION SF 30 BLOCK 14 CONTINUATION PAGE

SUMMARY OF CHANGES

SECTION 00010 - SOLICITATION CONTRACT FORM

DELIVERIES AND PERFORMANCE

The following Delivery Schedule item for CLIN 0001 has been changed from:

DELIVERY DATE	QUANTITY	SHIP TO ADDRESS	DODAAC / CAGE
POP 15-APR-2016 TO 14-APR-2018	N/A	W03J USA MED RESEARCH MAT CMD W03J USA MED RESEARCH MAT CMD 1077 PATCHEL STREET FORT DETRICK MD 21702-5024 301-619-7416 FOB: Destination	W91ZSQ

To:

DELIVERY DATE	QUANTITY	SHIP TO ADDRESS	DODAAC / CAGE
POP 15-APR-2016 TO 31-OCT-2019	N/A	W03J USA MED RESEARCH MAT CMD W03J USA MED RESEARCH MAT CMD 1077 PATCHEL STREET FORT DETRICK MD 21702-5024 301-619-7416 FOB: Destination	W91ZSQ

The following have been added by full text:

AWARD TRANSFER DETAILS**Award Transfer Details**

PI: David Yeomans

Project Title: Antilyso-phosphatidic Acid Antibodies in the Treatment of Post-TBI Neuropathic Pain

1. Lpath, Inc. submitted a relinquishment statement dated 10 October 2016, incorporated herein by reference, relinquishing this award as of 10 November 2016. Lpath, Inc. hereby relinquishes all future claims under this award.

2. There were no refund checks submitted by Lpath, Inc.

3. The total amount of this award for the full period of performance is \$1,446,655

Total amount awarded to Lpath, Inc.: \$269,779.26

Total amount awarded to Stanford University: \$1,176,875.74

4. The entire period of performance for this award is: 15 April 2016 – 31 October 2019

15 April 2016 – 07 March 2018: Lpath, Inc.

08 March 2018 – 31 October 2019: Stanford University

This award incorporates an approximate 19-month extension through 31 October 2019 with no additional funds.

5. The Final Technical Report to be submitted by Stanford University shall encompass the entire period of performance 15 April 2016 – 31 October 2019.

6. Payment Terms: Cost Reimbursement, Net 30 days.

7. The revised budget dated 16 November 2017 and the revised SOW, dated 16 February 2017, Submitted by Stanford University is incorporated, by reference, into this award.

SECTION 00800 - SPECIAL CONTRACT REQUIREMENTS

The following have been added by full text:

U.S. ARMY MEDICAL RESEARCH ACQUISITION ACTIVITY AWARD SPECIFIC RESEARCH TERMS AND CONDITIONS WITH INSTITUTIONS OF HIGHER EDUCATION, HOSPITALS, AND NON-PROFIT ORGANIZATIONS

DIVISION I – AWARD COVER PAGES

A. Award Information

1. **Department of Defense Awarding Office:** USAMRAA
2. **Award number/Project title:** W81XWH-16-1-0098/**Antilyosphosphatidic Acid Antibodies in the Treatment of Post-TBI Neuropathic Pain**
3. **Type of Award:** Grant
4. **Type of Award Action:** Modification – Grant Transfer
5. **i. Brief description of project or program:**

The DOD has solicited neurosensory research proposals to “support both applied (preclinical) research and clinical studies assessing traumatic brain injury (TBI) within specific Focus Areas of pain management, hearing loss/dysfunction, balance disorders, tinnitus, vision, or physical rehabilitation.” The proposed project responds to this solicitation and concerns the development of a new approach to the treatment of acute and chronic pain that is a frequent sequel to head injury. The proposed project responds to this solicitation and concerns the development of a new approach to the treatment of acute and chronic pain that is a frequent sequel to head injury. The prevalence of concussions in warfighters returning from Iraq or Afghanistan has been estimated at approximately 19.6% and the prevalence of posttraumatic craniofacial pain were present in 37% soldiers with 27% diagnosed with chronic daily headache and central TBI neuropathic pain. Chronic pain can contribute to psychological dysfunctions accompanying post-traumatic stress disorder. Current therapies, such as non-steroidal anti-inflammatory drugs (NSAID), opiates, anti-epileptics, and triptans can help some patients, but are far from universally effective and have, in some cases, substantial side-effect and abuse potential. Thus there is an acute need for improved strategies to prevent the development or treat existing chronic pain in civilians, war-fighters and veterans suffering from head trauma. The literature and our preliminary evidence clearly show that lysophosphatidic acid (LPA) plays a prominent role in nerve injury-induced pain in experimental animal models. In the CNS, LPA is synthesized by astrocytes, choroid plexus epithelial cells and inflammatory cells and is released upon cell activation. Its concentration within the brain increases during inflammation, clotting and neurotrauma, at which time it likely potentiates its known roles in astrocyte proliferation, neuronal death, axonal injury and

microglial activation. Our preliminary and published data indicate a specific upregulation of LPA receptors following injury to the adult mouse central nervous system (CNS), where LPA has been shown to induce neuronal apoptosis and to inhibit neural stem/progenitor cell differentiation along a neuronal lineage. In addition, LPA receptors are similarly increased after human brain injury. Moreover, LPA levels are upregulated in the CSF of TBI patients, suggesting.

ii. Funding Overview

	Federal funds	Cost Sharing	Total amount
a. Obligated or deobligated this action	\$0		\$0
b. Cumulative obligations to date, including this and previous actions	\$1,446,655		\$1,446,655
c. Planned project costs in the currently approved budget through the end of the period of performance, to include any future incremental funding obligations	\$1,176,875.74		\$1,176,875.74
d. Total value, which includes any unexercised options for which amounts were established in the award	\$1,446,655		\$1,446,655

6. **Obligation/Effective Date:** 15 April 2016
7. **Period of performance:** 15 April 2016 – 31 October 2019
8. **Authorities:** This award is made under the authority of 10 U.S.C. 2358.
9. **Catalog of Federal Domestic Assistance Number:** 12.420-Military Medical Research and Development
10. **Project Performance Information:**
 - i. This award is for research and development. Construction activities under this award are not authorized. (Reference Department of the Army Pamphlet 420-11, dated 18 March 2010, for the definition of construction activities.)
 - ii. Statement of Work and Budget: The revised Statement of Work (SOW) dated 16 February 2017 and the revised budget dated 16 November 2017 for your application submitted in response to the Fiscal Year 2014 DoD Clinical and Rehabilitative Medicine Research Program/Joint Program Committee 8, Neurosensory and Rehabilitation Research Award Program Announcement (Funding Opportunity Announcement Number W81XWH-14-CRMRP-NSRRA, which closed 11 February 2015) are incorporated herein by reference. You may rebudget allowable costs in accordance with applicable cost principles and in accordance with the prior approval requirements as stated in this award. Additional terms and conditions applicable to this award are in Division II and Division III.
 - iii. The following terms and conditions are incorporated herein by reference:
 - a. Division III - USAMRAA Addendum to the DoD R&D General Terms and Conditions available at <http://www.usamraa.army.mil/Pages/Resources.aspx>.
 - b. The DoD R&D General Terms and Conditions (September 2017), available at <http://www.onr.navy.mil/Contracts-Grants/submit-proposal/grants-proposal/grants-terms-conditions.aspx>.
 - iv. These USAMRAA Award Specific Research Terms and Conditions are in addition to the terms and conditions incorporated above. Any inconsistencies in the requirements of this award will be resolved in the following order:
 - a. Federal statutes
 - b. Federal regulations
 - c. 2 CFR part 200 with amendments, as modified and supplemented by DoD's interim implementation found in 2 CFR part 1103
 - d. Division II - USAMRAA Award Specific Research Terms and Conditions
 - e. Division III – USAMRAA Addendum to the DoD R&D General Terms and Conditions

f. DoD R&D General Terms and Conditions (September 2017)

v. **Grants Administration Office**

Grants Specialist: Christopher Meinberg

Phone: 301-619-2657

Email: christopher.l.meinberg.civ@mail.mil

Assistance Agreement Branch Email: usarmy.detrick.medcom-usamraa.mbx.aal@mail.mil

vi. **Grants Officer's Representative**

Congressionally Directed Medical Research Program Office

Phone: 301-619-7071

Email: usarmy.detrick.medcom-cdmrp.mbx.cdmrp-reporting@mail.mil

B. Recipient Information

1. **Unique Entity Identifier:** 009214214
2. **Recipient Business Name and Address:** The Leland Stanford Junior University
3. **Name and Title of Authorized Representative:** Natalie Muzzio
 - a. Phone: 650-724-0907
 - b. Email: muzzio@stanford.edu
4. **Principal Investigator and Organization:** David Yeomans
 - a. Phone: 650-725-5864
 - b. Email: dyeomans@stanford.edu
5. **Recipient's Indirect Cost Rate at the Start of the Performance Period:**
57%, Predetermined, MTDC, 02 August 2016
Negotiation Agency: Department of the Navy

83%, Predetermined, Animal Care, 02 August 2016
Negotiation Agency: Department of the Navy

C. Additional Information:

1. **Award Modification:** The only method by which the award may be modified is by a formal, written modification signed by the USAMRAA Grants Officer. No other communications, whether oral or in writing, are valid to change the terms and conditions of this award. Awards will not be modified to provide additional funds for such purposes as reimbursement for unrecovered indirect costs resulting from the establishment of final negotiated rates or for increases in salaries, fringe benefits, changes in exchange rates, or other costs.
2. **Expiration of Funds:** Funds obligated on this award are available for use for a limited period based on the fiscal year (FY) of the funds. That time is considered when establishing your period of performance. **This award is funded with FY15 funds in the amount of \$1,446,655 which will expire for use on September 30, 2021.** You must monitor the established milestones, timelines, expenditures and invoicing to make sure the project is on schedule and that you voucher promptly. If you have not submitted a final SF270 and been paid before the expiration date of these funds, any excess funds will be deobligated from the award at that time.

DIVISION II – AWARD SPECIFIC RESEARCH TERMS AND CONDITIONS**Federal Interagency Traumatic Brain Injury Research Informatics System**

This award involves research in the area of traumatic brain injury (TBI). The Department of Defense, in collaboration with the National Institutes of Health, has developed the Federal Interagency Traumatic Brain Injury Research (FITBIR) Informatics System, a central repository and resource for sharing data to promote collaboration, accelerate research, and advance knowledge on the characterization, prevention, diagnosis, and treatment of TBI. FITBIR provides a common platform and standardized format for data collection, retrieval and archiving, while allowing for flexibility in data entry and analysis.

The Principal Investigator is expected to share his/her data via FITBIR in accordance with FITBIR policy and procedures found at <https://fitbir.nih.gov/jsp/about/policy.jsp>. If the PI feels (s) he cannot submit data to the system, (s) he must coordinate with the GOR.

Interim (In-Progress) Progress Review

In addition to quarterly, annual, and final technical progress reports, the PI shall prepare for and participate in at least one Interim Progress Review (IPR) for each year of the project's term of award. Generally, the IPR will last no longer than two days and require no more than two overnight stays. It most likely will be held in the Fort Detrick, Maryland area, but may occur elsewhere in the U.S. The invitation and format for the IPR will be provided by the GOR at least 90 days prior to the scheduled date.

Quarterly Technical Reports

- a. For each year of the award, the PI must submit Quarterly Technical Progress Reports covering research results (positive and negative data) over a three month period (quarter). A reporting quarter begins with the start date of the award and restarts annually from that date for the entire period of performance. A Quarterly Technical Progress Report for the fourth quarter each year is not required, as the Annual Technical Report must incorporate all four quarters of progress.
- b. Quarterly reports are the most immediate and direct contact between the PI and the Grants Officer's Representative (GOR). The reports provide the means for keeping the US Army Medical Research and Materiel Command (USAMRMC) advised of developments and problems as the research effort proceeds. The reports also provide a measure against which funding decisions are made.
- c. Prepare all Quarterly reports in accordance with the Quarterly Technical Progress Report format, available at <http://www.usamraa.army.mil/Pages/Resources.aspx>. Each item of the report format must be completed.
- d. Each report must be submitted electronically, within 30 days after the end of each quarter, to the Grants Specialist and the GOR at the e-mail addresses specified in the front pages of this award. Name your file with your award number, followed by Year X Quarter Y Report (example: W81XWH-18-1-0000 Year 1 Quarter 1 Report.) If you have questions, contact the GOR.
- e. Special Requirements for Quarterly Technical Reports (must be submitted as an appendix to the quarterly report)

Quad Charts: The Quad Chart (available on <https://www.usamraa.army.mil>) must be updated and submitted as an appendix.

Special Requirements for Annual/Final Technical Reports

Special Requirements for Annual/Final Reports (must be submitted as an appendix to the annual/final report)

Quad Charts: The Quad Chart (available on <https://www.usamraa.army.mil>) must be updated and submitted as an appendix.

Title to Equipment - Conditional

Title to the infrared diode laser (Lasmed model LS110) acquired with award funds vests upon acquisition in you, subject to the conditions of Section A of PROP Article I of the DoD R&D General T&Cs. This equipment is non-exempt property and title is conditional. Upon completion of the award or when the equipment is no longer needed in performance of the award, you must request disposition instructions from the USAMRAA Grants Officer.

DIVISION III- USAMRAA ADDENDUM TO THE DoD GENERAL TERMS AND CONDITIONS AND USAMRAA PROGRAMMATIC REQUIREMENTS

Preamble

This award incorporates by reference the Department of Defense (DoD) Research and Development Terms and Conditions available at <https://www.onr.navy.mil/Contracts-Grants/submit-proposal/grants-proposal/grants-terms-conditions> . The USAMRAA Addendum to the DoD R&D General Terms and Conditions provides additional content relevant to USAMRAA awards for sections of specified articles from those general research terms and conditions. **The five asterisks indicate that there is content from the DoD R&D General Research Terms and Conditions within the identified parts and articles that remains unchanged and is not restated in this document. To understand the requirement for a given article, the DoD R&D General Research Terms and Conditions must be read in tandem with this USAMRAA Addendum.** The second portion of this addendum is comprised of the programmatic requirements portion of the general terms and conditions that apply to USAMRAA awards subject to the DoD R&D General Terms and Conditions.

USAMRAA Addendum to the DoD R&D General Terms and Conditions

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Part I: Definitions

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Section D. Definitions

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43. Intangible Property

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c. For purposes of USAMRAA awards, software is also considered intangible property.

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Part 2: Financial and Program Management (FMS Articles)

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FMS Article II. Payments

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Section C. Electronic Funds Transfer and other payment procedural instructions of information

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2. Other payment procedural instructions or information

a. Request for Payments

- i. Payments. Payments will be made to you upon receipt of a “grant voucher” (used for both grants and cooperative agreements) submitted through the Wide Area Work Flow (WAWF) e-Business Suite in accordance with the Contract Line Item Number (CLIN) structure set forth in this award. The Defense Finance and Accounting Service (DFAS) will generally make payments within 30 calendar days after we receive the request for reimbursement unless we reasonably believe the request is improper.

- ii. You must select “advance” or “reimbursement” on the grant voucher in WAWF.
- iii. In order to conserve administrative resources for both parties, you are encouraged to voucher no more frequently than monthly. Failure to voucher at least quarterly may raise concerns about research progress and the need for continued funding.
- iv. All payments will be made by Electronic Funds Transfer (EFT) to the bank account registered in the System for Award Management (SAM) (available at <https://www.sam.gov>). You must maintain the currency about yourself in SAM, including information necessary to facilitate payment via EFT. We cannot be held responsible for any misdirection or loss of payment which occurs as a result of your failure to maintain correct/current EFT information within your SAM registration. Failure to update SAM ensuring active account status will result in nonpayment.

b. Electronic Payment Instructions

- i. The Wide Area Work Flow (WAWF) e-Business Suite is the required method to electronically process your requests for payments. Once on the WAWF e-Business Suite web site, select the Invoicing, Receipt, Acceptance, and Property Transfer (iRAPT) button to electronically submit “grant vouchers” (used for both grants and cooperative agreements). You must (i) register to use WAWF at <https://wawf.eb.mil> and (ii) ensure an electronic business point of contact (POC) is designated in the System for Award Management (SAM) site at <https://www.sam.gov> within ten (10) calendar days prior to requesting a payment for this award. The Award specific Research Terms and Conditions will include additional instructions on how to submit grant vouchers and who to contact for assistance if needed.
- ii. Questions concerning specific payments should be directed to the Defense Finance and Accounting Service (DFAS), Indianapolis, at 1-888-332-7366, unless a different office is specified in Division II in your award specific terms and conditions. **You can also access payment and receipt information using the “myInvoice” button in WAWF at <https://wawf.eb.mil>.** The award number or grant voucher number will be required to inquire about the status of the payment.
- iii. The following codes and information are required to initiate the grant voucher and assure successful flow of WAWF documents.

TYPE OF DOCUMENT: **Grant Voucher** (*Used for both grants and cooperative agreements*)

CAGE CODE: **Enter Your Cage Code**

ISSUE BY DODAAC: **W81XWH**

ADMIN BY DODAAC: **W81XWH**

INSPECT BY DODAAC: **W81XWH**

ACCEPT BY DODAAC: **W81XWH**

SHIP TO DODAAC: **W81XWH**

LOCAL PROCESSING OFFICE DODAAC: **Not Applicable**

PAYMENT OFFICE FISCAL STATION CODE: **Unless otherwise specified in Division II in your award specific terms and conditions enter Fiscal Station DODAAC as HQ0490 = DFAS Indianapolis**

EMAIL POINTS OF CONTACT LISTING:

INSPECTOR: **Submit to Assistance Agreement Branch Email identified in the Division I, 10.v.**
 ACCEPTOR: **Submit to Assistance Agreement Branch Email identified in the Division I, 10.v.**
 RECEIVING OFFICE POC: **Submit to Assistance Agreement Branch Email identified in the Division I, 10.v.**
 GRANT ADMINISTRATOR: **Leave Blank**
 GRANTS OFFICER: **Leave Blank**
 ADDITIONAL CONTACT: **Submit to Assistance Agreement Branch Email identified in the Division I, 10.v.**

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FMS Article IV. Revision of budget and program plans.

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Section B. Revisions requiring prior approval.

1. Non-Construction Activities

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e. USAMRAA Specific Prior Approval Requirements

- i. The transfer (relocation) of the PI and or research project to another entity.
- ii. Reimbursing a DoD Military Treatment Facility (MTF) for costs incurred if the MTF is involved in the award. Reimbursing these costs is generally prohibited and only approved under unusual and extraordinary circumstances.

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Section C. Pre-award costs, carry forward of unobligated balances, and one-time no-cost extensions.

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3. No-cost Extension of the Period of Performance

- a. You may initiate one time, without prior approval, a no-cost extension to the expiration date of the award for a period of up to 12 months, as long as the no-cost extension does not involve a change in the approved objectives or scope of the project. You must notify the USAMRAA Grants Officer in writing at least 30 calendar days prior to the expiration date of the award. The notification must state the additional time needed, the reasons for the extension, and the work to be completed during the extension period. You must be current with all financial and technical reporting requirements and be in compliance with all other terms and conditions of the award. This one-time no-cost extension may not be exercised merely for the purpose of using unobligated balances. An official modification to the award document must be issued by the USAMRAA Grants Officer to extend the period of performance.
- b. Reference “Expiration of Funds” in Division I Award Cover Pages to understand the impact of the availability of funds on award extensions.
- c. Collaborating awards (two or more USAMRAA-issued awards completing the same Statement of Work) may have to have identical periods of performance. Each collaborating recipient’s business office must contact the Grants Officer assigned to the awards regarding extensions.
- d. Any subsequent no-cost extensions require prior approval from the USAMRAA Grants Officer.

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Part 5. Financial Programmatic, and Property Reporting (REP Article)

REP Article I. Performance management, monitoring, and reporting.

Section A. Required reporting form, format, or data elements for interim and final performance reports.

1. Annual Technical Report

- a. Annual reports are required and must be prepared in accordance with the Research Performance Progress Report (RPPR). The RPPR is the uniform format for reporting performance progress on Federally-funded research projects and research-related activities. Annual reports must provide a complete summary of the research results (positive or negative) to date in direct alignment to the approved Statement of Work (SOW). The importance of the report to decisions relating to continued support of the research cannot be over-emphasized.
- b. Special Requirements for Annual Reports-Refer to Division II.

2. Final Technical Report

- a. A final report must be prepared in accordance with the RPPR. The report must summarize the entire research effort, citing data in the annual reports and appended publications.
- b. Special Requirements for Final Reports-Refer to Division II.

3. Format

Prepare the annual and final reports in accordance with the RPPR format, available at <http://www.usamraa.army.mil/Pages/Resources.aspx>. Although there is no page limitation for the reports, each report must be of sufficient length to provide a thorough description of the accomplishments with respect to the approved SOW

Section B. Frequency, reporting periods, and due dates for interim performance reports.

An annual technical report must be submitted within 30 calendar days of the anniversary date of the award for the preceding 12 month period. If the award period of performance is extended by the USAMRAA Grants Officer, then an annual report must still be submitted within 30 days of the anniversary date of the award.

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Section F. Performance Reporting Procedures

Annual and Final Technical Reports, in electronic format (PDF or Word file only), must be submitted to <https://ers.amedd.army.mil>.

Additional information is available on the Researcher Resources website, available at https://mrmc.amedd.army.mil/index.cfm?pageid=researcher_resources.technical_reporting

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REP Article II. Financial Reporting.

Section A. Required reporting form, format, or data elements for interim and final financial reports.

You must submit financial reports on the Standard Form 425 (SF425) "Federal Financial Report."

Section B. Interim financial reports; frequency, reporting periods, and due dates.

The Federal Financial Reporting period end dates fall on the end of the calendar year for annual reports (12/31). You must submit annual reports no later than 90 days after the end of the calendar year.

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Section E. Where and how to submit financial reports.

1. You must submit all interim SF425 reports electronically through the web site <https://www.usamraa.army.mil/Pages/SF425.aspx>. The form and instructions can be obtained on this site.
2. Do not report multiple awards on one report. Each award must be reported separately on its own SF425.
3. Do not combine multiple SF425s into one submission. Each form must be saved as a separate PDF and submitted individually.
4. You must submit Final SF425 reports electronically to usarmy.detrick.medcom-usamraa.mbx.closeout@mail.mil

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REP Article III. Reporting on Property.

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Section D. Intangible Property.

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1. Inventions developed under this award.

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a. Patents and Inventions Reporting Requirements

- i. iEdison and annual reporting. You must electronically file Invention Disclosures and Patent Applications using the Interagency Edison (iEdison) system through the National Institutes of Health (<https://s-edison.info.nih.gov/iEdison>) within the times specified for reporting.
- ii. Report of Inventions and Subcontracts. A final DD Form 882 is required and must be submitted electronically within 120 days of end of the term of award. List all inventions made during the term of the award or state “none,” as applicable. The award will NOT be closed until you have met all reporting requirements. Submit the final DD882 reports electronically to usarmy.detrick.medcom-usamraa.mbx.closeout@mail.mil

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Part 6: Other Administrative Requirements (OAR Articles)

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OAR Article III Remedies and termination.

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Section B. Remedies for non-compliance.

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- f. If you are delinquent on technical reporting requirements for other USAMRAA-sponsored awards, no new awards will be issued to you until all delinquent reports have been submitted.
- g. Failure to submit required Technical Reports or Federal Financial Reports (SF425s) may delay payments or result in nonpayment.

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OAR Article IV. Claims, disputes, and appeals.

Section A. Definitions

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2. Grant Appeal Authority- Lamont G. Kapec Deputy Chief of Staff, Procurement and Head of the Contracting Activity HQDA Office of the Surgeon General and U.S. Army Medical Command, 7700 Arlington Boulevard Falls Church, VA 22042-5140

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OAR Article VI. Closeout

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Section B. Refunds of Unobligated balances.

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- a) Make check payable to the U.S. Treasury and mail to:
USAMRAA
Attn: MCMR-AAP-C
(Insert Federal Award No. W81XWH-16-1-0098
820 Chandler Street
Fort Detrick, Maryland 21702-5014

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Section D. Accounting for Property.

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- a) **Property Acquired with Award Funds, if applicable** [Reference PROP Article IV of the DoD R&D General Terms and Conditions (September 2017).]
 - i. If equipment under this award is exempt property, you must provide a cumulative listing of exempt equipment acquired with award funds. Submit this on your organization's letterhead. Submit to: Assistance Agreement Branch Email identified in the Division I, 10.v.
 - ii. If supplies under this award are exempt, you must submit a statement that: (i) there is, or is not, a residual inventory of unused supplies exceeding \$5,000 in total aggregate value; and (ii) if there is, state whether or not the unused items will be needed on other Federally sponsored

projects or programs. Submit this on your organization's letterhead. Submit to Assistance Agreement Branch Email identified in the Division I, 10.v.

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Part 8: National Policy Requirements

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NP Article III. National policy requirements concerning live organisms.

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Section B. Other requirements concerning live organisms

1. Research Involving Recombinant DNA Molecules By signing the award or accepting funds under the award, you assure that all work involving the use of recombinant DNA will be in compliance with guidance provided at <https://osp.od.nih.gov/biotechnology/biosafety-and-recombinant-dna-activities/>.

2. Prohibition of Use of Laboratory Animals

Notwithstanding any other terms and conditions contained in this award or incorporated by reference herein, the recipient is expressly forbidden to use or subcontract for the use of laboratory animals in any manner whatsoever without the express written approval of the USAMRMC, Animal Care and Use Review Office (ACURO). Written authorization to begin research under applicable protocol(s) proposed for this award will be issued in the form of an approval letter from the USAMRMC ACURO to the recipient with a copy to the USAMRAA Grants Officer. Furthermore, modifications to already approved protocols require approval by ACURO prior to implementation. For each fiscal year, the recipient must maintain, and upon request from ACURO, submit animal usage information.

Noncompliance with any of these terms and conditions may result in withholding of funds and/or the termination of the award.

The Animal Care and Use Office requirements can be accessed at https://mrmc.amedd.army.mil/index.cfm?pageid=research_protections.acuro.

3. Prohibition of Use of Human Subjects

Research under this award involving the use of human subjects, to include research involving the secondary use of human biospecimens and/or human data, cannot begin until the USAMRMC's Office of Research Protections (ORP) provides authorization that the research may proceed. The USAMRMC ORP will issue written approval to begin research under separate notification to you. Written approval to proceed from the USAMRMC ORP is also required for any subrecipient that will use funds from this award to conduct research involving human subjects.

The USAMRMC ORP conducts site visits as part of its responsibility for compliance oversight. Accurate and complete study records must be maintained and made available to representatives of the USAMRMC as a part of their responsibility to protect human subjects in research. Research records must be stored in a confidential manner so as to protect the confidentiality of subject information.

The recipient is required to adhere to the following reporting requirements:

Submission of substantive modifications to the protocol, continuing review documentation, and the final report as outlined in the USAMRMC ORP approval memorandum.

Unanticipated problems involving risks to subjects or others, subject deaths related to participation in the research, clinical holds (voluntary or involuntary), and suspension or termination of this research by the IRB, the institution, the Sponsor, or regulatory agencies, must be promptly reported to the USAMRMC ORP.

Change in subject status when a previously enrolled human subject becomes a prisoner must be promptly reported to the USAMRMC ORP HRPO.

The knowledge of any pending compliance inspection/visits by the FDA, ORP, or other government agency concerning this clinical investigation or research, the issuance of Inspection Reports, FDA Form 483, warning letters or actions taken by any Regulatory Agencies, and any instances of serious or continuing noncompliance with regulatory requirements that relate to this clinical investigation or research, must be reported immediately to the USAMRMC ORP.

Non-compliance with these terms and conditions may result in withholding of funds and/or the termination of the award.

DoD requirements for human subjects research, including 32 CFR Part 219, DoD Instruction 3216.02, and USAMRMC ORP Human Research Protection Office submission instructions can be accessed at https://mrmc.amedd.army.mil/index.cfm?pageid=research_protections.hrpo.

4. Prohibition of Use of Human Cadavers

Research, development, testing and evaluation (RDT&E), education or training activities involving human cadaveric specimens under this award shall not begin until approval is granted in accordance with the Army Policy for Use of Human Cadavers for RDT&E, Education, or Training, 20 April 2012 (https://mrmc.amedd.army.mil/index.cfm?pageid=research_protections.overview).

The USAMRMC Office of Research Protections (ORP) is the Action Office (usarmy.detrick.medcom-usamrmc.other.hrpo@mail.mil) for this policy. Approval must be obtained from the USAMRMC ORP. Award recipients must coordinate with the supporting/funding Army organization to ensure that proper approvals are obtained. ORP will issue written approvals to begin under separate notification to the recipient. Written approval to proceed from the USAMRMC ORP is also required for any subrecipient that will use funds from this award to conduct RDT&E, education or training involving human cadaveric specimens.

Recipients must promptly report problems related to the conduct of the activity involving cadavers or the procurement, inventory, use, storage, transfer, transportation, and disposition of cadavers to the USAMRMC ORP.

Recipients must maintain complete records of the activity.

The USAMRMC or designees must be permitted to observe the activity upon request and/or audit activity records to ensure compliance with the approved protocol or applicable regulatory requirements.

Non-compliance with these terms and conditions may result in withholding of funds and/or the termination of the award.

Programmatic Requirements Portion of the General Terms and Conditions

Publication, Acknowledgement, and Public Release

- a. Publication. You are encouraged to publish results of the research, unless classified, in appropriate media. Submit one copy of each paper to the GOR **simultaneously** with its submission for publication. Forward copies of all publications resulting from the research to the USAMRAA Grants Officer or Grants Specialist as they become available, even though publication may in fact occur subsequent to the termination date of the award. (See Section C of the DoD R&D General Terms and Conditions for the charging of publication costs incurred after the period of performance.)

- b. Acknowledgment. You agree that in the release of information relating to this award such release will include the statements below, as applicable. "Information" includes, but is not limited to, news releases, articles, manuscripts, brochures, advertisements, still and motion pictures, speeches, trade association meetings, and symposia.
- i. "The U.S. Army Medical Research Acquisition Activity, 820 Chandler Street, Fort Detrick MD 21702-5014 is the awarding and administering acquisition office" and;
 - ii. "This work was supported by the Office of the Assistant Secretary of Defense for Health Affairs through the Clinical and Rehabilitative Medicine Research Program/Joint Program Committee 8 under Award No. W81XWH-16-1-0098. Opinions, interpretations, conclusions and recommendations are those of the author and are not necessarily endorsed by the Department of Defense."
 - iii. "In conducting research using animals, the investigator(s) adheres to the laws of the United States and regulations of the Department of Agriculture."
 - iv. "In the conduct of research utilizing recombinant DNA, the investigator adhered to NIH Guidelines for research involving recombinant DNA molecules."
 - v. "In the conduct of research involving hazardous organisms or toxins, the investigator adhered to the CDC-NIH Guide for Biosafety in Microbiological and Biomedical Laboratories."
- c. Public release. Prior to release to the public, you must notify the USAMRAA Grants Officer and the GOR of the following: planned news releases, planned publicity, advertising material concerning project work, and planned presentations to scientific meetings. This provision is not intended to restrict dissemination of research information; the purpose is to inform the USAMRMC of planned public release of information on USAMRMC-funded research in order to adequately respond to inquiries and to be alerted to the possibility of inadvertent release of information.

Failure to include the above statements and adhere to the above regulations, when required, may result in loss of funding and/or termination of this award.

2. National Security

The award is intended for unclassified, publicly releasable research. You will not be granted access to classified information. We do not expect that the results of the research project will involve classified information. If, however, in conducting the activities supported under the award, you or the PI is concerned that any of the research results involve potentially classifiable information that may warrant Government restrictions on the dissemination of the results, you must promptly notify the USAMRAA Grants Officer.

3. Use of Non-Federal Personnel

Some USAMRMC program offices use contractor personnel to assist the GORs with review of technical reports. All review processes are conducted confidentially. Contractor personnel are required to sign agreements to protect the confidentiality of the information. Violations by reviewers that compromise the confidentiality of the reviews may result in suspension or debarment of the individual or contractor from Federal awards.

The following have been deleted:

USAMRAA-
XXXX-0002

AA T&C with For-Profits (Nov 2015)

NOV 2015

(End of Summary of Changes)