MEMORANDUM FOR SGVT
ATTN: CAPT BRITTANY LENZ

FROM: 59 MDW/SGVU

SUBJECT: Professional Presentation Approval

1. Your paper, entitled HLA-B Sequencing in Patients with Stevens-Johnson Syndrome and Toxic Epidermal Necrolysis presented at/published to AAD Annual Meeting, Orlando, FL, 3-7 March 2017 in accordance with MDW 41-108, has been approved and assigned local file #16399.

2. Pertinent biographic information (name of author(s), title, etc.) has been entered into our computer file. Please advise us (by phone or mail) that your presentation was given. At that time, we will need the date (month, day and year) along with the location of your presentation. It is important to update this information so that we can provide quality support for you, your department, and the Medical Center commander. This information is used to document the scholarly activities of our professional staff and students, which is an essential component of Wilford Hall Ambulatory Surgical Center (WHASC) internship and residency programs.

3. Please know that if you are a Graduate Health Sciences Education student and your department has told you they cannot fund your publication, the 59th Clinical Research Division may pay for your basic journal publishing charges (to include costs for tables and black and white photos). We cannot pay for reprints. If you are 59 MDW staff member, we can forward your request for funds to the designated wing POC.

4. Congratulations, and thank you for your efforts and time. Your contributions are vital to the medical mission. We look forward to assisting you in your future publication/presentation efforts.

[Signature]
PAUL T. BARNICOTT, GS-13-DAF
Deputy Director, Clinical Research Division

Warrior Medics – Mission Ready – Patient Focused
INSTRUCTIONS

1. The author must complete page two of the 59 MDW Form 3039 (this form):
   a) In Section 2, add the funding source for your study [e.g., 59 MDW CRD Graduate Health Sciences Education (GHSC) [SG5 O&M]; SG5 R&D; Tri-Service Nursing Research Program (TSNRP); Defense Medical Research & Development Program (DMRDP); NIH; Congressionally Directed Medical Research Program (CDMRP); Grants; etc.]

2. Print your name, rank/grade, sign and date the form in the author's signature block or use electronic signature.

3. Attach a copy of the 59th MDW IRB or IACUC approval letter for the research related study. If this is a technical publication/presentation, state the type (e.g., case report, QA/QI study, program evaluation study, informational report/briefing, etc.) in the "Protocol Title" box of the 59 MDW Form 3039.

4. Attach a copy of your abstract, paper, poster and other supporting documentation.

5. Save and forward, via email, the processing form and all supporting documentation to your Unit Commander, Program Director or immediate supervisor for review/approval.

6. On page 2, have either your Unit Commander, Program Director or immediate supervisor:
   a) Print their name, rank/grade/title, sign and date the form in the approving authority's signature block or use electronic signature.

7. Contact the 59th CRD/Publications and Presentations Section at (292-7141) for instructions for submitting the request form.

8. The 59th CRD/Publications and Presentations Section will route the request form to clinical investigations and public affairs and forward you a final letter of approval or disapproval.

9. Once your manuscript, poster or presentation has been approved for a one-time public release, you may proceed with your publication or presentation submission activities, as stated on the 59 MDW Form 3039. [NOTE: For each new release of medical research or technical information as a publication/presentation, a new 59 MDW Form 3039 must be submitted for review and approval.]

10. If your manuscript is accepted for scientific publication, please contact the 59th CRD/Publications and Presentations Section (292-7141). This information is reported to the 59 MDWICC. All medical research or technical information publications/presentations must be reported to the Defense Technical Information Center (DTIC). See 59 MDWI 41-108, Presentation and Publication of Medical and Technical Papers for additional information.

NOTE: All abstracts, papers, posters, etc., should contain the following disclaimer statement:

"The views expressed are those of the [author(s)] [presenter(s)] and do not reflect the official views or policy of the Department of Defense or its Components."

NOTE: All abstracts, papers, posters, etc., should contain the following disclaimer statement for research involving humans:

"The voluntary, fully informed consent of the subjects used in this research was obtained as required by 32 CFR 219 and DoD 3216.02, Protection of Human Subjects in Biomedical and Behavioral Research."

NOTE: All abstracts, papers, posters, etc., should contain the following disclaimer statement for research involving animals, as required by AFMAN 40-401_IP, The Care and Use of Laboratory Animals in DoD Programs:

"The experiments reported herein were conducted according to the principles set forth in the National Institute of Health Publication No. 80-23, Guide for the Care and Use of Laboratory Animals and the Animal Welfare Act of 1966, as amended."
1. TITLE OF MATERIAL TO BE PUBLISHED OR PRESENTED
HLA-B SEQUENCING IN PATIENTS WITH STEVENS-JOHNSON SYNDROME AND TOXIC EPIDERMAL NECROLYSIS

2. FUNDING RECEIVED FOR THIS STUDY? □ YES □ NO
FUNDING SOURCE:

3. IS THIS MATERIAL CLASSIFIED? □ YES □ NO

4. IS THIS MATERIAL SUBJECT TO ANY LEGAL RESTRICTIONS FOR PUBLICATION OR PRESENTATION THROUGH A COLLABORATIVE RESEARCH AND DEVELOPMENT AGREEMENT (CRADA), MATERIAL TRANSFER AGREEMENT (MTA), INTELLECTUAL PROPERTY RIGHTS AGREEMENT ETC.? □ YES □ NO
NOTE: If the answer is “YES” then attach a copy to the Agreement to the Publications/Presentations Request Form.

5. MATERIAL IS FOR (Check appropriate box or boxes for approval with this request.)
□ PUBLICATION/JOURNAL (List intended publication/journal)
□ PUBLICATION ABSTRACT (List intended journal.)
□ POSTER (To be demonstrated at meeting/Name of Meeting, City, State, and Date of Meeting)
AAD ANNUAL MEETING, ORLANDO, FL. 03-07 MARCH 2017
□ PLATFORM PRESENTATION (At civilian institutions/Name of Meeting, State, Date of Meeting)
□ OTHER (Describe: Name of Meeting, City, State, and Date of Meeting)

6. WHAT IS THE EXPECTED DATE YOUR PRESENTATION/PUBLICATION WILL BE SUBMITTED TO THE DEFENSE TECHNICAL INFORMATION CENTER (DTIC)?

7. WHO IS THE PRIMARY 59 MDW POINT OF CONTACT? (Last, First, Ml.) (Include email)
LENZ, BRITTANY L.
brittany.lenz@us.af.mil

AUTHORSHIP AND CO-AUTHOR(S) (List in the order they will appear in the manuscript)

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<tr>
<th>LAST NAME</th>
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I CERTIFY ANY HUMAN OR ANIMAL RESEARCH RELATED STUDIES WERE APPROVED AND PERFORMED IN STRICT ACCORDANCE WITH 32 CFR 219, AFMAN 40-401, IP AND 59 MDW 41-108. I HAVE READ THE FINAL VERSION OF THE ATTACHED MATERIAL AND CERTIFY THAT IT IS AN ACCURATE MANUSCRIPT FOR PUBLICATION AND/OR PRESENTATION.

AUTHOR’S PRINTED NAME/RANK/GRADE
Brittany Lenz, Capt, O-3

AUTHOR’S SIGNATURE

APPROVING AUTHORITY’S PRINTED NAME, RANK, TITLE
Wendy Wohltmann, Lt Col, Program Director

APPROVING AUTHORITY’S SIGNATURE

DATE
Nov 16, 2016

DATE
Nov 16, 2016
| TO: Clinical Research Division  
| (59 MDW/SGVU)  
| (Contact 292-7141 for email instructions) | 1. DATE RECEIVED | 2. ASSIGNED PROCESSING REQUEST FILE NUMBER |
| | | Nov 17, 2016 | 16399 |
| 3. DATE REVIEWED | Dec 6, 2016 | 4. DATE FORWARDED TO PA |

5. AUTHOR CONTACTED FOR RECOMMENDED OR NECESSARY CHANGES

- [ ] NO
- [ ] YES (If yes give date: Nov 17, 2016)
- [ ] N/A

6. COMMENTS

- [ ] APPROVED
- [ ] DISAPPROVED

IRB approved retrospective poster with appropriate disclaimers.

PRINTED NAME, RANK/GRADE, TITLE OF REVIEWER | DATE | SIGNATURE OF REVIEWER |
---|---|---|
Kevin Kupferer/GS13/Hum Res Subj Prot Expert | Dec 6, 2016 | KUPFERER.KEVIN.R.1086667270 |

2nd INDORSEMENT (PA Use Only)

| TO: 59 MDW OFFICE OF PUBLIC AFFAIRS (PA) | 1. DATE RECEIVED | 2. DATE forwarded TO 59 MDW/SGVU |
| | Dec 6, 2016 | Dec 8, 2016 |

6. COMMENTS

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- [ ] DISAPPROVED

PRINTED NAME, RANK/GRADE, TITLE OF REVIEWER | DATE | SIGNATURE OF REVIEWER |
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Kevin Inuma, SSgt/E-5, 59 MDW Public Affairs | Dec 8, 2016 | INUMA.KEVIN.MITSUGU.296227613 |

3rd INDORSEMENT (SGVU Use Only)

| TO: 59 MDW/SGVU | 1. DATE RECEIVED |
| | |

2. SENIOR AUTHOR NOTIFIED BY PHONE OF APPROVAL OR DISAPPROVAL:

- [ ] YES
- [ ] NO
- [ ] Could not be reached
- [ ] Left message

3. DATE WRITTEN NOTICE OF APPROVAL AND CLEARANCE MAILED TO AUTHOR:

4. COMMENTS

- [ ] APPROVED
- [ ] DISAPPROVED

PRINTED NAME, RANK/GRADE, TITLE OF REVIEWER | SIGNATURE OF REVIEWER | DATE |
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59 MDW Form 3039 20151028  Supersedes 59 MDW Form 3039, 20131018  Page 3 of 3
Introduction

Stevens-Johnson syndrome (SJS) and toxic epidermal necrolysis (TEN) are severe cutaneous adverse drug reactions (ADRs) associated with significant morbidity and mortality (Figures 1-4). Though the exact pathogenesis remains unclear, targeted genomic analysis has identified relationships with certain immunologic markers and enzymes associated with drug metabolism. HLA-B types in particular have been associated with ADRs; however, to date these associations exist for only a few specific drugs and patient populations.

Given the unique presentation and ability to obtain target tissue for analysis, severe cutaneous ADRs provide an ideal target for research using precision medicine techniques. We initially sought to examine whether certain HLA-B alleles were present at an increased frequency in patients with SJS/TEN. Now through whole genome sequencing and epigenetic analysis, we aim to identify novel biomarkers for cutaneous adverse drug reactions.

Methods

We conducted a retrospective study of SJS/TEN patients admitted to the San Antonio Military Medical Center (SAMMC) Burn Unit between 2001 and 2015. Targeted sequencing of the HLA-B gene was performed on 28 formalin-fixed paraffin-embedded (FFPE) skin biopsy samples from cases with a known offending drug. Typically, HLA-B gene alleles are determined by sequencing Exons 2 and 3 of the gene, using primers that anneal in a non-variable region of the introns. This was not possible with FFPE samples due to fragmentation of DNA; therefore we used commercial primers designed from the least variable areas of the exons possible.

Using the Sanger sequencing method, we identified the potential HLA-B alleles present in the FFPE samples and evaluated for an association with the offending drug (Table 1).

Results

Multiple potential HLA-B alleles were identified in most of our specimens. This highlights the limitations of DNA sequencing using FFPE samples, as the fragments of DNA and non-overlapping sequences limited our ability to determine an exact HLA type for every specimen. Based on these results, we were unable to determine statistically significant associations regarding frequency of HLA type or inciting drugs. However, our data show HLA-B*44 as a potential allele in 9 of the 28 samples, including 5 of 10 cases associated with Sulfonamides and SJS/TEN with severe ocular complications.

Discussion & Prospective Study

Severe cutaneous ADRs remain a significant cause of morbidity and mortality in health care and are often unpredictable. To further investigate potential genetic risk factors for SJS/TEN, we designed a prospective study using whole genome sequencing and transcriptome studies to examine epigenetic changes during ADRs (see below).

We are currently enrolling patients at SAMMC through 2018. A secondary goal of this study is the establishment of a tissue repository at the Collaborative Health Initiative Research Program at the Uniformed Services University to aid in future studies examining adverse drug reactions.

References