AWARD NUMBER: W81XWH-17-1-0451

TITLE: The Effect of a Powered Ankle Foot Orthosis (PAFO) on Function, Safety, and Quality of Life in Military Service Members and Veterans Who Wear a Prescribed Orthosis

PRINCIPAL INVESTIGATORS: Phyllis A. Richey, PhD.

CONTRACTING ORGANIZATION: University of Tennessee
Memphis, TN 38103

REPORT DATE: September 2018

TYPE OF REPORT: Annual

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

DISTRIBUTION STATEMENT: Approved for Public Release; Distribution Unlimited

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<tbody>
<tr>
<td>Phyllis A. Richey, PhD. (Joint-PIs)</td>
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<tr>
<td>E-Mail: <a href="mailto:prichey@uthsc.edu">prichey@uthsc.edu</a></td>
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<td>This project is a 2-arm, parallel, randomized, controlled clinical trial designed to determine if a powered ankle foot orthosis (PAFO), that assists with toe clearance and provides push-off power when taking a step, will translate into enhanced function in individuals who walk with a prescribed AFO. We will assess these outcomes in 64 veterans who walk with a prescribed AFO by randomizing participants, in a 1:1 ratio, into an intervention and a comparison group. Participants in both groups will receive new shoes to be worn with their orthosis to eliminate any confounding variables presented by worn or inadequate shoes. Participants in the intervention group will be fitted with and trained to use a PAFO while the comparison group will continue with their current prescribed orthosis. All participants will be followed with weekly contact over a 7-month period of time and receive physical therapy training. All outcome measures will be evaluated three times during the 7-month study period. Local IRB approval was received August 2018 and HRPO application was submitted. We are currently awaiting HRPO approval. Recruitment resources via Partner Orthotic clinics, Regional DAV, local area health care agencies and physician practices have been identified and approval to contact potential study volunteers is currently in progress. Recruitment, enrollment/randomization, intervention and follow-up assessments will take place during in the coming quarters/year.</td>
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<td>Ankle foot orthosis (AFO), powered ankle foot orthosis (PAFO), randomized clinical trial, functional performance, ambulatory safety, falls, quality of life, gait symmetry, foot drop, stroke, spinal cord injury (SCI), traumatic brain injury (TBI), peripheral injury.</td>
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1. INTRODUCTION:

This project is a 2-arm, parallel, randomized, controlled clinical trial designed to quantify functional performance, gait symmetry, ambulatory safety, and quality of life in 64 veterans who walk with a prescribed AFO. The cohort will be randomized in a 1:1 ratio into an intervention or a comparison group. The blocked randomization schedule will be generated by a computer program with a block size of 4; this will guarantee that we have approximately the same number of participants in each treatment group throughout the trial. Participants in both groups will receive new shoes to be worn with their orthosis to eliminate any confounding variables presented by worn or inadequate shoes. Participants in the intervention group will receive a PAFO while the comparison group will continue with their currently prescribed orthosis. All participants will be followed with weekly contact over a 7-month period of time and receive physical therapy training to minimize deviations resulting from habit or lack of training, education to maximize use of the mechanical properties of their currently prescribed AFO, strengthening and stretching based on published guidelines, balance training and training on traversing environmental barriers. All outcome measures will be evaluated three times during the 7-month study period: At baseline, at the 4-month follow up visit and at the 7-month follow up visit. We believe the immediate benefit of this project will determine if an innovative PAFO, designed to assist with toe clearance and provide push off power when taking a step, will improve functional performance, gait symmetry, ambulatory safety (risk of falls), and quality of life in the typical veteran with lower extremity impairment. We will also study whether the same variables/constructs show evidence of any carry over effect of the PAFO when the patients are not wearing an AFO. This study will have significant long-term benefit for all people who depend on an AFO to walk, both veterans and the general public, as they face medical, social and psychological complications associated with falling (broken bones, head trauma, depression, social isolation and death), decreased function and poor quality of life that directly impacting their families and caregivers.

2. KEYWORDS:
Ankle foot orthosis (AFO)
Powered ankle foot orthosis (PAFO)
Randomized clinical trial
Functional performance
Ambulatory safety
Falls
Quality of life
Gait symmetry
Foot drop
Stroke
Spinal cord injury (SCI)
Traumatic brain injury (TBI)
Peripheral injury.
3. ACCOMPLISHMENTS:

What were the major goals of the project?
The major goals of this project as stated in the approved SOW are as follows:
1. Perform Preliminary Study Requirements (Months 1-6)
2. Recruit, Coordinate and Train Study Personnel for Clinical Trial (Months 3-6)
3. Participant Recruitment, Screening Eligibility and Baseline Evaluations (Months 7-24)
4. Participant Randomization (Months 7-24)
5. Participant Fit with Powered Ankle Foot Orthosis (PAFO); Intervention Group (N=32; Months 7-24)
6. Physical Therapy Sessions and Orthosis Accommodation Period (N=64; Months 7-25)
7. 4-Month Follow Up Visit and Prosthesis Accommodation Period (N=64; Months 10-30)
8. 7-Month Follow Up Visit and subject closure (N=64; Months 13-30)
9. Assess Secondary Aims (N=64; Months 7-36)
10. Data Analysis/Dissemination of Findings (Months 28-36)

What was accomplished under these goals?
1. Perform Preliminary Study Requirements
   a. Prepare study documents and apply for Local IRB (UTHSC) – Completed 08/06/2018
   b. Apply for USAMRM Human Research Protection Office (HRPO) approval- Submitted, awaiting HRPO approval.
   c. Complete Manual of Operations finalizing procedures sections and forms for recruiting and reporting – Completed
   d. Develop database management system – Completed
   e. Develop and finalize all study data collection forms – Completed
   f. Submit amendments, adverse events and protocol deviations – None to report.
   i. Maintain, update and perform data integrity test on study DBMS – NA at this time

2. Train Study Personnel for Clinical Trial
   a. Train staff, evaluation physical therapist, treating physical therapist for project – Completed
   b. Trial run through of Screening and Baseline visits for the study – In Progress
   c. Eligibility and Randomization training – In Progress
   d. Adverse Events Training with Dr. Johnson, MD – In Progress
   e. Develop participant recruitment materials – In Progress

3. Participant Recruitment
   a. Establish participant recruitment resources with partner Orthotic Clinics, healthcare agencies, physician practices and Regional DAV – In progress
   b. Identify targeted mailings to prospective participants – In progress

What opportunities for training and professional development has the project provided?
During this reporting period a professional development seminar for all study personnel and physical therapists was conducted by ReWalk Restore engineers trainers at our study facility at the University of Tennessee Health Science Center (UTHSC) on the use of the Restore PAFO.
How were the results disseminated to communities of interest?

Nothing to Report

What do you plan to do during the next reporting period to accomplish the goals?

During the next reporting period we will continue to perform the following actions to accomplish the goals and objectives listed:

1. Complete development and approval of study recruitment materials
2. Identify prospective participants for targeted recruitment
3. Participant Recruitment and enrollment, phone (pre-) screening, schedule in-person screening eligibility visit
4. Confirm pre-screening information at in-person Screening Eligibility Visit
   a. Sign Informed Consent
   b. Confirm pre-screening information
   c. Perform screening evaluation including evaluation of functional level of participant
   d. Evaluate orthotic fit
5. Participant Randomization
   a. Randomize participants into Intervention (N=32) or Comparison (N=32) Groups
   b. Schedule physical therapy visits
   c. Provide all participants new pair of shoes
6. Participant Fit with PAFO
   a. Conduct physical therapy sessions
   b. Provide all participants 2 sessions per week of physical therapy for 4 weeks
   c. Provide weekly phone visits during 8-week following completion of PT sessions to all participants in both groups
7. Perform 3-month evaluation
   a. Perform repeat of all baseline evaluation measures
   d. Provide weekly phone visits during 12-week following completion of 3-month follow-up visit to all participants in both groups
8. Perform 6-month evaluation and subject closure
   a. Perform repeat of all baseline evaluation measures
9. Continue to perform ongoing study requirements
   a. Submit amendments, adverse events and protocol deviations as necessary
   b. Maintain, update and perform data integrity test on study DBMS
4. **IMPACT:** Describe distinctive contributions, major accomplishments, innovations, successes, or any change in practice or behavior that has come about as a result of the project relative to:

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

Nothing to Report

**What was the impact on other disciplines?**

Nothing to Report

**What was the impact on technology transfer?**

Nothing to Report

**What was the impact on society beyond science and technology?**

Nothing to Report

5. **CHANGES/PROBLEMS:** The Project Director/Principal Investigator (PD/PI) is reminded that the recipient organization is required to obtain prior written approval from the awarding agency Grants Officer whenever there are significant changes in the project or its direction. If not previously reported in writing, provide the following additional information or state, “Nothing to Report,” if applicable:

**Changes in approach and reasons for change**

During this reporting period, we were informed that BionX Medical Technology, manufacturer of the emPOWER powered ankle foot orthosis (PAFO) device originally proposed in our study design, was sold to another company which decided not continue production of the emPOWER therefore requiring we use a different device for our study intervention. After an exhaustive search for an alternative PAFO device and were able to reach an agreement with ReWalk Robotics Inc. to supply our study with their new product, Restore Exosuit PAFO to carry out the study intervention. A request to change the study device was submitted and approved 01/24/2018.

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

As referenced above a delay in executing our timeline was incurred during the first two quarters of the reporting period while searching for and awaiting approval to use the Restore device. After approval was received we established confidentiality and data use agreements with the device manufacturer, ReWalk, for use of the Restore Exosuit PAFO device and proceeded with local UTHSC IRB application. UTHSC IRB approval was received August 6, 2018 and HRPO application was promptly submitted. During the next reporting period we will coordinate acquisition of the Restore Exosuit PAFO device from with ReWalk Robotics Inc. and continue as planned with study recruitment.
Changes that had a significant impact on expenditures

As described above in actual delays during this reporting period, we experienced significant delays 1) identifying a replacement study device for the one originally proposed that was discontinued by the manufacturer, 2) receiving approval to use the replacement device, and 3) receiving local IRB approval. In response to this delay we choose to provide funding support to only the essential staff necessary during the study start up period. Therefore, only the joint PIs, Drs. Richey and Singhal and the study staff, Mr. Hood and Ms. Leone, were supported during the reporting period. This intentional staffing support decision, along with a delay in initiation of recruitment has translated into a delayed expenditure of participant-related study items (e.g. the Restore Exosuit, participant incentives, etc) and allowed a significant savings in the expenditures originally anticipated during the reporting period. By taking these expenditure-saving measures during this reporting period we will have the funds available in the coming reporting periods to fulfill the goals of this project.

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

Significant changes in use or care of human subjects

None

Significant changes in use or care of vertebrate animals.

Not Applicable

Significant changes in use of biohazards and/or select agents

Not Applicable

6. PRODUCTS:

- Publications, conference papers, and presentations
  Nothing to Report

- Journal publications
  Nothing to Report

- Books or other non-periodical, one-time publications.
  Nothing to Report

- Other publications, conference papers, and presentations
  Nothing to Report

- Website(s) or other Internet site(s)
  Nothing to Report
• **Technologies or techniques**  
Nothing to Report

• **Inventions, patent applications, and/or licenses**  
Nothing to Report

• **Other Products**  
Nothing to Report

7. **PARTICIPANTS & OTHER COLLABORATING ORGANIZATIONS**

What individuals have worked on the project?

Name: Phyllis Richey, PhD  
Project Role: Joint-Principal Investigator  
Research Identifier: 1  
Nearest person month worked: 12  
Contribution to Project: Dr. Richey is fulfilling the role of co-Principal Investigator as outlined in the SOW.

Name: Kunal Singhal, PhD, PT  
Project Role: Co-Investigator  
Research Identifier: 2  
Nearest person month worked: 12  
Contribution to Project: Dr. Singhal is fulfilling the role of co-Principal Investigator as outlined in the SOW.

Name: Kristen Leone  
Project Role: Study Coordinator  
Research Identifier: 3  
Nearest person month worked: 6  
Contribution to Project: Ms. Leone has worked with IRB submissions, HRPO submissions, MOP development, recruitment materials, staff development and training

Name: Matt Hood  
Project Role: Study Coordinator/Informatics  
Research Identifier: 4  
Nearest person month worked: 12  
Contribution to Project: Mr. Hood has worked with IRB submissions, HRPO submissions, database development, data collection form design, staff development and training

Has there been a change in the active other support of the PD/PI(s) or senior/key personnel since the last reporting period?  
Nothing to Report
**What other organizations were involved as partners?**

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Organization Name: Region One Healthcare
Location of Organization: Tennessee
Partner’s contribution to the project: Assisting with recruitment
Financial support: None
In-kind support: None
Facilities: None
Collaboration: Dissemination study informational materials to potential participants
Personnel exchanges: None
Other: None

8. SPECIAL REPORTING REQUIREMENTS

COLLABORATIVE AWARDS:
Not Applicable

QUAD CHARTS:
Attached

9. APPENDICES:
Not Applicable
Study Aim: Primary (1)
To determine if a PAFO with greater range of motion and active power improve functional performance while wearing the PAFO, as well as carry over to improved functional performance when the PAFO is not worn.

Secondary Aims (3):
To determine if use of a PAFO, with greater range of motion and active power, will improve (SA:1) gait symmetry, (SA:2) ambulatory safety (reduce risk of falls), and (SA:3) general quality of life and orthotic-related quality of life while wearing the PAFO, as well as carry over to improved gait symmetry when the PAFO is not worn.

Approach:
We will randomize 64 patients with neurologic impairment who have significant weakness of the foot and ankle musculature in a 1:1 ratio to either the intervention or comparison condition. The study is a 2-arm, parallel, randomized, controlled clinical trial.

3 Year Timeline and Cost

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Estimated Budget ($K) | $736,000 | $800,000 | $462,000

Updated September 2018

Goals/Milestones

Year 1– Project Planning, Start-Up, Recruitment, Enrollment, Intervention and Follow-Up Evaluation
- Define processes/obtain all approvals for safety and compliance. Begin participant recruitment and baseline evaluation and randomization treatment condition.
- Begin intervention including orthotic fitting period (2 weeks) and physical therapy sessions.
- Begin follow-up evaluations and retention including 3 month follow up visits and accommodation phase 1 with weekly phone visits

Year 2– Continue, Enrollment, Intervention and Follow-Up Evaluation
- Complete participant recruitment and baseline evaluation, complete orthotic fitting period and physical therapy sessions
- Continue 3 month and begin 6 month follow up visits and retention.
- Complete accommodation phase 1 and begin accommodation phase 2

Year 3– Project Completion and Close Out
- Complete 3 and 6 month follow up visits and accommodation phase 2
- Complete data analyses
- Disseminate findings in journal and conference venues

Accomplishment: We are currently enrolling Veteran trans-tibial amputees in the DoD funded “Veterans Leading pRostetic Research” (VALOR) study. This study is contributing to the improved rehabilitation of amputees classified as “community ambulators” to enhance their function and quality of life.

The effect of a powered ankle foot orthosis (PAFO) on function, safety & quality of life in military service members and veterans who wear a prescribed orthosis OP160076

Joint PI’s: Richey, P.A. & Singhal, K.  Organization: University of Tennessee Health Science Center  Budget: $1,998,325