AWARD NUMBER: W81XWH-14-2-0150

TITLE: Improving Balance in TBI Using a Low-Cost Customized Virtual Reality Rehabilitation Tool

PRINCIPAL INVESTIGATOR: Denise Krch, PhD

CONTRACTING ORGANIZATION: Kessler Foundation Inc
West Orange, NJ 07052

REPORT DATE: October 2016

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

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.
### 4. TITLE AND SUBTITLE
Improving Balance in TBI Using a Low-Cost Customized Virtual Reality Rehabilitation Tool

### 6. AUTHOR(S)
Denise Krch, PhD and Karen J. Nolan, PhD

### 7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES)
Kessler Foundation, 0705
120 Eagle Rock Ave., Suite 100
East Hanover, NJ 07936-3147

### 9. SPONSORING / MONITORING AGENCY NAME(S) AND ADDRESS(ES)
U.S. Army Medical Research and Materiel Command
Fort Detrick, Maryland 21702-5012

### 12. DISTRIBUTION / AVAILABILITY STATEMENT
Approved for Public Release; Distribution Unlimited

### 14. ABSTRACT
The proposed study will implement and evaluate a novel, low-cost, Virtual Reality (VR) rehabilitation tool (Island Quest; IQ) targeting somatosensory, vestibular, and vision systems through a double-blind RCT. Given the importance of dual-task skills for real-world functioning, we will also evaluate the relative effectiveness of dual task (balance and cognitive) VR training to improve balance.

A total of 180 participants (Service Members, Veterans, civilians) with mild to severe TBI and documented balance impairments will be randomly assigned into one of three balance treatment groups: 1) Standard of care (control condition); 2) IQ; 3) IQ dual task (balance plus cognitive). All groups will undergo 2 treatment sessions/week x 6 weeks. Following completion of the treatment protocol, participants in the IQ training group will be randomly assigned to a maintenance training group (2 sessions/month x 4 months) or a non-maintenance group. All participants will undergo baseline, immediate (6 weeks), and long-term (4 months) follow-up assessments of: 1) static and dynamic balance and 2) community integration, self-efficacy, quality of life, and cognitive function. This design will allow us to assess the efficacy of IQ as a customizable balance treatment in TBI, and to evaluate the impact of this remediation program on overall functioning.

### 15. SUBJECT TERMS
Virtual reality, balance dysfunction, dual task, traumatic brain injury, multisensory, cognitive, motor

### 16. SECURITY CLASSIFICATION OF:
- a. REPORT Unclassified
- b. ABSTRACT Unclassified
- c. THIS PAGE Unclassified

### 17. LIMITATION OF ABSTRACT
Unclassified

### 18. NUMBER OF PAGES
20

### 19a. NAME OF RESPONSIBLE PERSON
USAMRMC

### 19b. TELEPHONE NUMBER (include area code)

**Table of Contents**

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Introduction</td>
<td>3</td>
</tr>
<tr>
<td>2. Keywords</td>
<td>3</td>
</tr>
<tr>
<td>3. Accomplishments</td>
<td>3</td>
</tr>
<tr>
<td>4. Impact</td>
<td>13</td>
</tr>
<tr>
<td>5. Changes/Problems</td>
<td>14</td>
</tr>
<tr>
<td>6. Products</td>
<td>15</td>
</tr>
<tr>
<td>7. Participants &amp; Other Collaborating Organizations</td>
<td>17</td>
</tr>
<tr>
<td>8. Special Reporting Requirements</td>
<td>19</td>
</tr>
<tr>
<td>9. Appendices</td>
<td>19</td>
</tr>
</tbody>
</table>
1. INTRODUCTION:

The proposed study will implement and evaluate a novel, low-cost, Virtual Reality (VR) rehabilitation tool (Island Quest; IQ (recently renamed from Mystic Isle)) targeting somatosensory, vestibular, and vision systems through a double-blind RCT. Given the importance of dual-task skills for real-world functioning, we will also evaluate the relative effectiveness of dual task (balance and cognitive) VR training to improve balance. A total of 180 participants (Service Members, Veterans, civilians) with mild to severe TBI and documented balance impairments will be randomly assigned into one of three balance treatment groups: 1) Standard of care (control condition); 2) IQ; 3) IQ dual task (balance plus cognitive). All groups will undergo 2 treatment sessions/week x 6 weeks. Following completion of the treatment protocol, participants in the IQ training group will be randomly assigned to a maintenance training group (2 sessions/month x 4 months) or a non-maintenance group. All participants will undergo baseline, immediate (6 weeks), and long-term (4 months) follow-up assessments of: 1) static and dynamic balance and 2) community integration, self-efficacy, quality of life, and cognitive function. This design will allow us to assess the efficacy of IQ as a customizable balance treatment in TBI, and to evaluate the impact of this remediation program on overall functioning.

2. KEYWORDS: Virtual reality, balance dysfunction, dual task, traumatic brain injury, multisensory, cognitive, motor

3. ACCOMPLISHMENTS:
   - What were the major goals of the project?

<table>
<thead>
<tr>
<th>Project Milestones &amp; Deliverables</th>
<th>Timeline</th>
<th>Complete</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Phase I - Project Kick-off</strong></td>
<td>9/30/14 - 3/30/15</td>
<td>%</td>
</tr>
<tr>
<td><strong>Subtasks Phase I:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Submit Administrative Approval requests - regulatory review and approval processes to include local Institutional Review Board (IRB) and DoD Human Research Protection Office.</td>
<td>09/30/14 - 03/30/15</td>
<td>100%</td>
</tr>
<tr>
<td>2. Coordinate with CRMRP, USC ICT, NICO ISO (Fort Belvoir) and VANJHCS.</td>
<td>09/30/14 - 03/30/15</td>
<td>100%</td>
</tr>
<tr>
<td>3. Purchase study equipment and supplies, configure for study methods, and set up at study sites.</td>
<td>09/30/14 - 03/30/15</td>
<td>100%</td>
</tr>
<tr>
<td>4. Advertise for, interview, and hire study personnel.</td>
<td>09/30/14 - 03/30/15</td>
<td>100%</td>
</tr>
<tr>
<td>5. Prepare study assessment and outcome measures, organize participant folders (e.g., case report forms) and paperwork.</td>
<td>09/30/14 - 03/30/15</td>
<td>100%</td>
</tr>
<tr>
<td>6. Train study personnel in study methods, including evaluation of balance, global functioning, and cognition.</td>
<td>12/31/14 - 03/30/15</td>
<td>95%</td>
</tr>
<tr>
<td>7. Train study personnel in double-blind RCT procedures.</td>
<td>12/31/14 - 03/30/15</td>
<td>100%</td>
</tr>
<tr>
<td>8. Train study personnel in administering study treatment conditions.</td>
<td>12/31/14 - 03/30/15</td>
<td>95%</td>
</tr>
<tr>
<td>9. Set up study database.</td>
<td>01/31/15 - 03/30/15</td>
<td>100%</td>
</tr>
<tr>
<td>10. Finalize project-related modifications to the balance treatment protocols.</td>
<td>01/31/15 - 03/30/15</td>
<td>100%</td>
</tr>
</tbody>
</table>
### Phase II - Clinical Trial (Years .5 to 3.5)

<table>
<thead>
<tr>
<th>Subtasks Phase II:</th>
<th>3/31/15 - 03/30/18</th>
<th>% Complete</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Conduct telephone and in-person screening to evaluate for inclusion/exclusion criteria.</td>
<td>03/31/15 - 09/29/17</td>
<td>30%</td>
</tr>
<tr>
<td>2. Begin Clinical Trial Recruitment and Enrollment.</td>
<td>03/31/15 – 11/29/17</td>
<td>20%</td>
</tr>
<tr>
<td>3. Randomize participants into Standard of Care Balance (control), Mystic Isle (experimental), or Mystic Isle Dual Task (experimental) treatment.</td>
<td>03/31/15 – 11/29/17</td>
<td>10%</td>
</tr>
<tr>
<td>4. Conduct Balance, Global Functioning, and Cognition baseline assessments.</td>
<td>04/30/15 - 03/30/18</td>
<td>10%</td>
</tr>
<tr>
<td>5. Review sessions to evaluate treatment fidelity.</td>
<td>03/31/15 - 03/30/18</td>
<td>5%</td>
</tr>
<tr>
<td>6. Conduct immediate follow-up Balance, Global Functioning, and Cognition assessments.</td>
<td>07/31/15 - 03/30/18</td>
<td>5%</td>
</tr>
<tr>
<td>7. After completion of the treatment protocol, randomize single task IQ group participants into Maintenance or Non-Maintenance group.</td>
<td>07/31/15 - 01/30/18</td>
<td>5%</td>
</tr>
<tr>
<td>8. Conduct Maintenance sessions.</td>
<td>07/31/15 - 03/30/18</td>
<td>5%</td>
</tr>
<tr>
<td>9. Conduct long-term follow-up Balance, Global Functioning, and Cognition assessments.</td>
<td>07/31/15 - 03/30/18</td>
<td>2%</td>
</tr>
</tbody>
</table>

### Phase III: Project Completion (Final 12 Months)

<table>
<thead>
<tr>
<th>Subtasks Phase III:</th>
<th>09/30/15 - 09/29/18</th>
<th>% Complete</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Conclude data collection.</td>
<td>09/30/17 - 03/30/18</td>
<td>0%</td>
</tr>
<tr>
<td>2. Conduct data analysis.</td>
<td>03/31/18 - 09/29/18</td>
<td>0%</td>
</tr>
<tr>
<td>3. Prepare final report and manuscripts for publication, and other dissemination efforts to military and civilian consumers and professionals.</td>
<td>03/31/18 - 09/29/18</td>
<td>5%</td>
</tr>
</tbody>
</table>

### Phase I, II, and II Outcomes, Products and Deliverables:

<table>
<thead>
<tr>
<th>% Complete</th>
</tr>
</thead>
<tbody>
<tr>
<td>09/30/14 – 9/29/18</td>
</tr>
<tr>
<td>Personnel hired and trained.</td>
</tr>
<tr>
<td>Equipment and methods set up and implemented at study sites.</td>
</tr>
<tr>
<td>Full IRB approval and DoD Human Research Protection Office.</td>
</tr>
<tr>
<td>Subjects run according to the methodological plan.</td>
</tr>
<tr>
<td>Data entered, analyzed, interpreted and presented (progress reports, manuscripts).</td>
</tr>
</tbody>
</table>
What was accomplished under these goals?

<table>
<thead>
<tr>
<th>Phase 1 – Project Kick-off</th>
<th>% Complete</th>
<th>Specific Objectives Achieved</th>
</tr>
</thead>
<tbody>
<tr>
<td>Major Activities</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| 1. Submit Administrative Approval requests - regulatory review and approval processes to include local Institutional Review Board (IRB) and DoD Human Research Protection Office. | 100%        | • Kessler’s initial IRB application submitted to Kessler Foundation (KF) IRB (05/5/2014); Approval received (6/13/14).  
• IRB amendment submitted to reflect changes in protocol consistent with DoD grant application methodology (08/26/2014); e.g., addition of veteran and military personnel to protocol; Approval received (9/3/14).  
• IRB amendment submitted with minor clarification changes (9/11/14); Approval Received (9/18/14).  
• IRB amendment submitted with changes to be in compliance with the requirements of the U. S. Army Medical Research and Material Command (USAMRMC) (9/24/14); Approval received (9/29/14).  
• Kessler’s initial IRB application submitted to HRPO (11/4/2014); Received request for clarification from HRPO (1/13/15); Responded to HRPO’s requests for clarification (1/30/15) and submitted memo to local IRB to request risk determination (1/30/15) in reference to HRPO’s 1/13/15 email correspondence; IRB determined non-significant risk (3/2/15); Submitted IRB non-significant risk determination to HRPO (3/2/15); Received additional requests for clarification from HRPO (3/4/15); Responded to HRPO’s requests for clarification (4/8/15); Received additional requests for clarification from HRPO (5/14/15); Responded to HRPO’s requests for clarification (6/4/2015); Received permission from HRPO to submit changes to local IRB (6/15/15); Submitted local IRB approval of changes to HRPO (6/23/15); Received request for clarification of protocol version number from HRPO (6/30/15).  
• Established IRB Agreement with USC ICT, with USC ICT acting under Kessler’s FWA (04/01/15).  
• Submitted recruitment flyer to local IRB (4/7/15); Received approval from local IRB for flyer (4/8/15)  
• Submitted flyer to VANJHCS IRB contact person to seek guidance on steps to gain approval to post flyer for Veteran recruitment on VANJHCS campus.  
• Submitted yearly review/continuation application to local IRB (5/1/15); Received continuation approval (5/5/15).  
• Submitted amendment with HRPO changes to local IRB (6/19/15); Received approval of changes from local IRB (6/23/15).  
• Submitted amendment to add required VANJHCS language to flyers to local IRB (7/16/15). Received approval (7/17/15).  |
<table>
<thead>
<tr>
<th>Date</th>
<th>Event Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>7/30/15</td>
<td>Ft. Belvoir site specific amendment (SSA) submitted to Ft. Belvoir IRB Manager for initial review (7/30/15) and forwarded for administrative review on 7/31/15.</td>
</tr>
<tr>
<td>7/31/15</td>
<td>Teleconference between the Defense Health Agency (DHA), Ft. Belvoir Research Staff, and Dr. Zhang in order to discuss the need for a Data Sharing Agreement (DSA) between the DHA and Kessler Foundation/System Security Verification (SSV) for data capture system (8/6/15). It was later established that neither a DSA or an SSV would be required.</td>
</tr>
<tr>
<td>9/3/15</td>
<td>Submitted amendment adding names of recently hired physical therapists and personnel from collaborating sites to local IRB (9/3/15). Received approval (9/4/15).</td>
</tr>
<tr>
<td>9/25/15</td>
<td>Sarah Rule, NICOE ISO Fort Belvoir Community Hospital’s (FBCH) Research Compliance Officer agreed to rely on Kessler’s IRB review for NICOE ISO approval (9/25/15). The IRB reliance agreement (IAIR) is currently being routed for signature at the FBCH Command Suite level.</td>
</tr>
<tr>
<td>9/9/15</td>
<td>IRB manager compiled a list of suggested revisions and additional documentation required for the new project submission and sent it to the Fort Belvoir (FB) Research Coordinator for review and editing (9/9/15).</td>
</tr>
<tr>
<td>9/18/15</td>
<td>Received draft marketing project (study advertisement to be displayed in hospital/TBI NICOE ISO clinic and on electronic display board in hospital), (9/18/15). FB RC made final edits to this document and received final version on 9/22/15.</td>
</tr>
<tr>
<td>9/25/15</td>
<td>DRP Administrative Review is completed for NICOE ISO (9/25/15).</td>
</tr>
<tr>
<td>10/8/15</td>
<td>The Office of the Undersecretary of Defense for Personnel and Readiness Research Regulatory Oversight Office (R2O2) delegated that the Component Level Administrative Review (CLAR) be performed by Sarah Rule, Acting Chief Department of Research Programs, Human Protections Administrator, and Research Oversight &amp; Compliance Officer at Fort Belvoir Community Hospital (FBCH) (10/8/15).</td>
</tr>
<tr>
<td></td>
<td>A request was submitted from the Fort Belvoir Department of Research Programs to the Kessler IRB for clarification regarding the risk determination of the protocol and Fort Belvoir study staff was subsequently notified that the protocol was determined to be greater than minimal risk and as a result of this determination a DoD Research Monitor (RM) would need to be assigned to oversee the</td>
</tr>
</tbody>
</table>
A research monitor was identified by the PI and study coordinator at Fort Belvoir and following completion of human subjects training (CITI) was added to the protocol. The updated SSA and supporting documentation were then submitted to the Fort Belvoir IRB Manager for review (12/3/15).

The CLAR was completed by Sarah Rule at Fort Belvoir (12/7/2015) and then forwarded to R2O2 for review (12/8/2015).

Submitted amendment adding alternate test (Bilingual Aphasia Test: Verbal Comprehension) to Token Test for individuals with color vision impairment (12/9/15). Received approval (12/15/15).

Fort Belvoir forwarded IRB documents to KF IRB for review (12/22/15).

Fort Belvoir SSA was approved by the Kessler IRB on 12/24/2015 and approval documents were sent to Fort Belvoir (1/15/2016).

Face-to-face PI Responsibilities meeting between Sarah Rule and Dr. Purohit, FB RC also in attendance (1/29/2016).

Sarah Rule sent email to Kessler PI, Karen Nolan, requesting clarification on the risk determination on 2/2, and received clarification from Dr. Greene regarding the risk determination. The Kessler IRB determined the risk of the research protocol to be no greater than minimal risk (2/8/16).

IRB amendment submitted correcting medical therapy section of the protocol (2/9/16); Approval Received (2/11/16).

Submitted protocol amendment (Amendment #1) after receiving clarification in risk from Kessler; updated SSA to reflect this change from greater than minimal risk to minimal risk and to remove DoD Research Monitor. Also, updated Dr. Chae’s status from Collaborator to Associate Investigator (2/11/16).

Received required revisions back from FB IRB Manager along with notification that adding Dr. Chae as an AI on the protocol would require the leadership signature to go up a level to LTC Waits, the Director of Behavioral Health and Dr. Chae’s supervisor at FBCH (2/17/16).

LTC Waits signed off on the protocol and all required revisions and documentation were submitted to the Fort Belvoir IRB (2/26/16).

Amendment #1 and all supporting documents sent to Kessler IRB for review (3/3/16).

Amendment #1 was approved by the Kessler IRB on 3/3/2016 and the approval letter was forwarded to
| 2. Coordinate with CRMRP, ICT, NICoE ISO and VANJHCS. | 100% | • Established communication with DoD Science Officer (07/29/2014).
• Contract negotiations completed; award date established by DoD Contracting Officer (09/17/2014).
• A subcontract was established with the University of Southern California, Institute for Creative Technologies (USC ICT; agreement executed 11/19/2014).
• A subcontract was initiated with Geneva for collaboration with NICoE ISO (signed by Geneva on 12/1/2014).
• Conducted first site visit (3/11/2015) at Fort Belvoir (Karen Nolan and Denise Krch, Co-PIs; Irene Ward, Treatment Intervention Liaison).
• Established communication with VANJHCS regarding recruitment through consultant Glenn Wylie
• Began discussing steps required to obtain IRB approval to post Veteran recruitment flyer on VANJHCS campus as well as those steps required to submit an IRB application to gain access to the VANJHCS subject recruitment database.
• Supporting IRB application preparation activities at NICoE ISO through regular communication with NICoE ISO’s research coordinator (RC). |
| 3. Purchase study equipment and supplies, configure for study methods, and set up at study sites. | 100% | • Purchase orders for KF neuropsychological tests submitted end of December, 2014.
• Created neuropsychological testing administration binder.
• Created data collection worksheets, sample subject binder, clinical trial regulatory binder, and IRB communication binder.
• Conducted ongoing meetings with KF, Kessler Institute for Rehabilitation (KIR), and USC ICT regarding study methods. |
### Methodology
- Completed POs for balance intervention equipment.
- Received office supplies, computer equipment (including monitor and Microsoft Kinect), patient hi-low table, and Mini Mental Status Examination to determine capacity to consent.
- Balance intervention equipment ordered for KF. Most equipment has been received.

### Advertise for, Interview, and Hire Study Personnel
- Kathleen Goworek Chervin was assigned as the Research Coordinator (RC) at KF.
- Lea Frank, Research Assistant (RA), was hired at KF.
- NIcoE ISO placed ad for RA.
- Fort Belvoir hired Caitlin Jones, RC (start date 3/30/15).
- Kelli Sullivan was assigned the RA at NIcoE ISO.
- Advertised for Physical Therapist position at KF.
- Hired PTs Adam Kesten and Christina Cording at KF.

### Prepare Study Assessment and Outcome Measures, Organize Participant Folders (e.g., Case Report Forms) and Paperwork
- Created scoring algorithm spreadsheet and hard copy summary sheet for patient testing.
- Study statistician completed first version of electronic case report form system.
- Study statistician optimized electronic case report form system for data collection and randomization.

### Train Study Personnel in Study Methods, Including Evaluation of Balance, Global Functioning, and Cognition
- All KF and KIR personnel completed CITI training.
- Kessler RC and RA trained to use Mystic Isle.
- KF RC completed training the RA and engineer on balance and mobility assessments.
- KF RA completed training on administration of cognitive and global functioning evaluation tools.
- NIcoE ISO Site PI and RC completed CITI training.
- KF PTs completed CITI training.

### Train Study Personnel in Double-Blind RCT Procedures
- Reviewed RCT procedures with Kessler study staff; briefed Fort Belvoir on double-blind procedures during site visit.
- Finalized RCT procedures with KF study staff.

### Train Study Personnel in Administering Study Treatment Conditions
- Kessler study staff was briefed on administration of treatment conditions.
- Continued progress in treatment protocol manual to be provided to all study staff to ensure standardization of treatment administration across personnel and sites.
- Finalizing implementation of treatment conditions using IQ with USC ICT.
- Finalized manualization of Standard of Care treatment.
- KF PTs were trained to use IQ. Clinical review of SOC and IQ treatment conditions resulted in additional required software refinements. Coordinated with USC ICT to begin implementing these refinements.
- Completed software refinements.
- Finalized manualization of IQ treatment conditions.
- NIcoE ISO PT completed onsite training at KF to review SOC and IQ treatment conditions.
9. **Set up study database.** 100%

- Coordinated with NICoE ISO study staff to prepare for enrollment and data collection launch at NICoE ISO.
- Study statistician completed first version of electronic case report form system.
- Study statistician optimized electronic case report form system for data collection and randomization.
- Neuropsychological data entry sheets were added to the electronic data capture system.
- Secondary randomization time point was implemented.
- KF RC and RA completed initial beta testing of data entry and randomization of subjects.

10. **Finalize project-related modifications to the balance treatment protocols.** 100%

- Finalizing implementation of treatments conditions using Mystic Isle with USC ICT.
- Finalized manualization of Standard of Care treatment.
- Continued progress in treatment protocol manual to be provided to all study staff to ensure standardization of treatment administration across personnel and sites.
- Initial delivery of the updated Mystic Isle software from USC was delayed. Upon delivery, KF’s study team conducted a thorough review of the software and identified areas in need of refinement. Since then, we have been working diligently with USC to implement these refinements to bring the software in line with the SOC treatment.

### Phase II - Clinical Trial

<table>
<thead>
<tr>
<th>Major Activities</th>
<th>% Complete</th>
<th>Specific Objectives Achieved</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Conduct telephone and in-person screening to evaluate for inclusion/exclusion criteria.</td>
<td>30%</td>
<td>• Ongoing telephone and in-person screening</td>
</tr>
<tr>
<td>2. Begin Clinical Trial recruitment and enrollment.</td>
<td>20%</td>
<td>• 15 participants have been enrolled to date.</td>
</tr>
<tr>
<td>3. Randomize participants into Standard of Care Balance (control), Island Quest (IQ; experimental), or IQ Dual Task (experimental) treatment.</td>
<td>10%</td>
<td>• 14 participants have been randomized into treatment</td>
</tr>
<tr>
<td>4. Conduct Balance, Global Functioning, and Cognition baseline assessments.</td>
<td>10%</td>
<td>• 14 participants have completed baseline assessments</td>
</tr>
<tr>
<td>5. Review sessions to evaluate treatment fidelity.</td>
<td>5%</td>
<td>• KF PT is completing clinical documentation after each treatment session to allow the PIs to monitor treatment fidelity and ensure systematic treatment delivery</td>
</tr>
<tr>
<td>6. Conduct immediate follow-up Balance, Global Functioning, and Cognition assessments.</td>
<td>5%</td>
<td>• 7 participants have completed immediate follow-up assessments</td>
</tr>
</tbody>
</table>
7. After completion of the treatment protocol, randomize single task IQ group participants into Maintenance or Non-Maintenance group. 5%

8. Conduct Maintenance sessions. 5%

9. Conduct long-term follow-up Balance, Global Functioning, and Cognition assessments. 2%

**Phase III - Project Completion**

<table>
<thead>
<tr>
<th>Major Activities</th>
<th>% Complete</th>
<th>Specific Objectives Achieved</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Conclude data collection.</td>
<td>0%</td>
<td></td>
</tr>
<tr>
<td>2. Conduct data analysis.</td>
<td>0%</td>
<td></td>
</tr>
</tbody>
</table>

- What opportunities for training and professional development has the project provided?
  - KF PT, Adam Kesten, conducted vestibular rehabilitation and technology inservice to Kessler Institute for Rehabilitation Brain Injury PTs July, 2016.
  - Co-PI Krch presented Virtual Reality didactic lecture for Rutgers, KF, and Children’s Specialized Hospital post-doctoral fellows April, 2016.

- How were the results disseminated to communities of interest?
  - Although we do not yet have results to disseminate, we are actively disseminating information about the project and creating increased awareness about balance deficits in TBI to several communities of interest:
    - Industry Collaborators:
      - Parker Hannifin “Brain Injury Mobility Research”, April, 2016
    - Clinical Collaborators:
      - Kessler Institute for Rehabilitation Brain Injury PTs, July, 2016
      - Rutgers, KF, and Children’s Specialized Hospital postdoctoral fellows, April, 2016
    - Scientific Collaborators:
• **What do you plan to do during the next reporting period to accomplish the goals?**
  • Finalize organizing and setting up the balance treatment space at NICOE ISO.
  • Finalize and test out NICOE ISO study team on administration of global functioning and cognition measures.
  • Finalize and test out NICOE ISO PT in administering study treatment conditions.
  • Finalize training of NICOE ISO study staff to enter data into study database.
  • Continue collaboration between KF, KIR and USC ICT.
  • Align local study subject database with FITBIR.
  • KF study team to travel to NICOE ISO to evaluate treatment fidelity and data collection procedures.
  • Begin patient recruitment and telephone and in-person screening for inclusion/exclusion criteria at NICOE ISO.
  • Continue patient recruitment and telephone and in-person screening for inclusion/exclusion criteria at KF.
  • Continue enrollment and randomization of qualifying participants at KF.
  • Conduct preliminary data analysis with baseline data for dissemination.
  • Continue to communicate with VANJHCS and continue community outreach to recruit veterans within our study sample.

• **IMPACT:**
  • **What was the impact on the development of the principal discipline(s) of the project?**
    - *For the purposes of this project, we utilized existing balance treatment strategies and synthesized them into a multisensory treatment protocol to be delivered systematically through a virtual environment approach. Balance dysfunction is the result of damage or deficits to multiple systems, however, these integrated systems are often not treated systematically. Our experimental protocols treat the various components of balance dysfunction individually, and then as integrated system, thus enabling us to target impairments in their individual domains as well as holistically. The systematic delivery of this approach is accomplished through the use of virtual reality technology. These features are what elevates the treatment protocol to have greater potential than existing treatments for balance dysfunction.*
What was the impact on other disciplines?

- The additional utilization of a dual task treatment protocol will enable us to extend the research question to the field of neuropsychology. Implementing a dual task condition will enable us to better understand whether challenging the brain to attend to cognitive and motor demands will effect a significantly greater change in the target system of interest (i.e., balance) relative to treatment of that system alone.

What was the impact on technology transfer?

- We believe the prototype system that we now have would be considered to be at DOD Technology Readiness Level (TRL) 7: “System prototype demonstration in an operational environment”. We anticipate that the results from this investigation will produce evidence for the IQ system at TRL 9 through empirical clinical and objective support for its widespread application as a standard efficacious clinical and research tool. A customizable tool, such as IQ, could be offered as a rehabilitation treatment to clinics or health care providers. A number of health care providers and small businesses have demonstrated interest in the existing VR-based prototype tool. We expect IQ’s greater efficacy and cost effectiveness, decreased lab space requirement, and decreased requirement for sophisticated equipment and skilled technicians, to further adoption/transition of our system as a standard treatment tool for balance.

What was the impact on society beyond science and technology?

- Island Quest has implications as a telerehabilitation application, which would enable Service Members and Veterans in distant locations to independently use the training system with remote clinical supervision. This would also represent a great benefit to rural patients as well as patients with transportation barriers. The ability to reach far more patients than would ordinarily be able to present themselves to a rehabilitation facility translates into significantly improved overall quality of care and health care outcomes, and thus, is beneficial in reducing healthcare costs and burden to the healthcare system.

• CHANGES/PROBLEMS:

- Changes in approach and reasons for change
  - Nothing to report.
1. We have been experiencing challenges recruiting veterans through VANJHCS. As such, we have explored additional avenues and are working closely with the dedicated recruitment coordinator at KF to put together a strong recruitment plan. This coordinator has previous experience recruiting veterans with TBI for Kessler Foundation. Thus, we expect recruitment of veterans to improve over the next two quarters. Since the last quarterly report, we have recruited and enrolled one veteran. We anticipate continuing to increase veteran participation in this study.

2. We experienced a delay hiring the NICoE ISO PT; Kendra Reid was hired on 2/16/16 and was fully trained by the completion of this report.

3. We experienced a slight delay due to the decision to invest additional time to improve and refine the software before initiating the clinical trial. Modifications are now completed and we have begun recruitment and enrollment.

4. One of the study Research Assistants left KF for graduate school. However, we have already hired a replacement, who is now fully trained.

Despite delays in these areas, we have continued to move forward on other major project goals and have begun to look toward clinical trial milestones.

Changes that had a significant impact on expenditures

- Nothing to report.

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

- Nothing to report.

Significant changes in use or care of human subjects

- Nothing to report.

Significant changes in use or care of vertebrate animals.

- Nothing to report.

Significant changes in use of biohazards and/or select agents

- Nothing to report.

PRODUCTS:

Publications, conference papers, and presentations

- Journal publications.
  Nothing to report.

- Books or other non-periodical, one-time publications.
  Nothing to report.
Other publications, conference papers, and presentations

- KF PT, Adam Kesten, conducted vestibular rehabilitation and technology inservice to Kessler Institute for Rehabilitation Brain Injury PTs July, 2016.
- Co-PI Krch presented Virtual Reality didactic lecture for Rutgers, KF, and Children’s Specialized Hospital post-doctoral fellows April, 2016.

- Website(s) or other Internet site(s)
  www.kesslerfoundation.org - Official website of Kessler Foundation, a non-profit research organization dedicated to improving the lives of persons with disabilities. This website provides information about current research (with links to related press releases), publications and presentations, and community outreach. (Kessler Foundation is the primary research site).

- Technologies or techniques
  Nothing to report.

- Inventions, patent applications, and/or licenses
  Nothing to report.

- Other Products
  - Software: For the purpose of this project, Island Quest software was modified from a game-based exercise/rehabilitation tool to a multisensory balance treatment software that can be systematically delivered to individuals with neurological conditions.
  - Clinical interventions: For the purposes of this project, a Standard of Care multisensory balance treatment protocol was synthesized and manualized.
• PARTICIPANTS & OTHER COLLABORATING ORGANIZATIONS
  
  o What individuals have worked on the project?

<table>
<thead>
<tr>
<th><strong>Kessler Foundation</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Name:</strong></td>
</tr>
<tr>
<td><strong>Project Role:</strong></td>
</tr>
<tr>
<td><strong>Researcher Identifier:</strong></td>
</tr>
<tr>
<td><strong>Nearest person month worked:</strong></td>
</tr>
<tr>
<td><strong>Contribution to Project:</strong></td>
</tr>
</tbody>
</table>

| **Name:** | Karen J. Nolan, PhD |
| **Project Role:** | Co-Principal Investigator |
| **Researcher Identifier:** | orcid.org/0000-0002-4667-0873 |
| **Nearest person month worked:** | 3.6 |
| **Contribution to Project:** | Dr. Nolan contributed to personnel hiring and training, study organization and set-up, and acted as a liaison between personnel across study sites. Dr. Nolan provided guidance and oversight to treatment study software refinements. Dr. Nolan (unblinded) oversees treatment intervention sessions. |

| **Name:** | Kathleen Goworek Chervin, PhD |
| **Project Role:** | Research Coordinator |
| **Researcher Identifier:** | N/A |
| **Nearest person month worked:** | 10.2 |
| **Contribution to Project:** | Ms. Chervin managed administrative and IRB tasks as well as organized the regulatory and IRB documentation for KF and HRPO. Ms. Chervin trained RAs and engineers on the mobility outcome measures. Ms. Chervin provides guidance for all study activities at NCoE ISO. She also manages the electronic capture system. |

| **Name:** | Lea Frank, BS |
| **Project Role:** | Research Assistant |
| **Researcher Identifier:** | N/A |
| **Nearest person month worked:** | 6 |
| **Contribution to Project:** | Ms. Frank assisted Ms. Chervin in administrative activities and ordering study supplies. She created the neuropsychological testing binder and became proficient in administering the study balance assessments. Ms. Frank conducts screening and study balance and cognitive assessments. She is responsible for entering data into the data capture system. |

| **Name:** | Adam Kesten, DPT |
| **Project Role:** | Physical Therapist |
| **Researcher Identifier:** | N/A |
| **Nearest person month worked:** | 1.2 |
| **Contribution to Project:** | Adam Kesten worked with the study team to refine the treatment protocols. He contributed to creation of the treatment protocol manual. Mr. Kesten trained RAs and engineers on safety and spotting techniques for balance assessments. Mr. Kesten is currently |
Melvin Mejia, B.S. is responsible for administering all balance treatment sessions at KF.

**Name:** Melvin Mejia, B.S.
**Project Role:** Biomedical Engineer
**Researcher Identifier:** N/A
**Nearest person month worked:** 4.8
**Contribution to Project:** Melvin Mejia conducts balance assessments and assists the PT with treatment administration. His technological expertise is utilized in various aspects of this technology-based research study.

**NICoE ISO**

**Name:** Maulik Purohit, MD, MPH
**Project Role:** Principal Investigator
**Researcher Identifier:** N/A
**Nearest person month worked:** 1.2
**Contribution to Project:** Dr. Purohit contributed to personnel hiring, study organization and set-up, and acted as a liaison between personnel across study sites.

**Name:** Caitlin Jones
**Project Role:** Research Coordinator
**Researcher Identifier:** N/A
**Nearest person month worked:** 12
**Contribution to Project:** Ms. Jones manages all administrative and IRB tasks at NICoE ISO. She is trained to administer the mobility outcome measures and currently being trained to administer the cognitive outcome measures. Ms. Jones works closely with the KF RC, Kate Chervin to ensure standardization across sites.

**Name:** Kendra Reid
**Project Role:** Physical Therapist
**Researcher Identifier:** N/A
**Nearest person month worked:** 6
**Contribution to Project:** Kendra Reid has been training under Mr. Kesten, KF PT, to become proficient in administering the SOC and IQ treatment interventions.

---

- **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?**
  - **Kessler Institute for Rehabilitation, West Orange, NJ, USA**
    - Significant contribution to the manualization of the Standard of Care and the Mystic Isle treatment protocols.
    - Training clinical staff and refining and standardizing treatment delivery across treatment sites
University of Southern California, Institute for Creative Technologies, Los Angeles, CA, USA

- Modification of the Island Quest software from a game-based exercise/rehabilitation tool to a multisensory balance treatment
- Will provide software support and assistance with data extraction from the Island Quest system.

National Intrepid Center of Excellence, Intrepid Spirit One, Fort Belvoir Community Hospital, Fort Belvoir, VA, USA

- Study data collection site for active duty military population
- Provided input on refining treatment protocols for military populations

- SPECIAL REPORTING REQUIREMENTS.
  - QUAD CHARTS: See below in Appendices.

- APPENDICES:
  - Quad Chart.
Improving Balance in TBI using a Low-Cost Customized Virtual Reality Tool

PI: Denise Krch, PhD and Karen J. Nolan, PhD
Org: Kessler Foundation
Award Amount: $2,987,537

Study/Product Aim(s)

- **Objective 1**: Evaluate the effectiveness of Virtual Reality (VR)-based balance training using Island Quest (IQ) to improve balance in individuals with TBI.
- **Objective 2**: Evaluate the improvement on measures of global functioning following the VR balance training customized for a rehabilitation setting.
- **Objective 3**: Evaluate the effectiveness of VR-based dual task (balance and cognitive) training to improve balance in individuals with TBI.
- **Objective 4**: Evaluate the long-term efficacy of VR-based balance training through the inclusion of a 4-month, follow-up assessment examining balance and functional gains.
- **Objective 5**: Evaluate utility of maintenance training.

Approach

Participants (n=180) will be enrolled into a double-blind RCT at Kessler Foundation/Kessler Institute for Rehabilitation and the National Intrepid Center of Excellence: Intrepid Spirit One (NICoE ISO) Fort Belvoir Community Hospital. Individuals with TBI will be randomly assigned into 1 of 3 balance interventions (2 sessions/week x 6 weeks): 1) Standard of Care; 2) IQ; 3) IQ + dual task (balance and cognitive).

Timeline and Cost

<table>
<thead>
<tr>
<th>Activities</th>
<th>CY 14</th>
<th>CY 15</th>
<th>CY 16</th>
<th>CY 17</th>
</tr>
</thead>
<tbody>
<tr>
<td>IRB submittal and study prep</td>
<td></td>
<td></td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Study staff training</td>
<td></td>
<td>✔</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Recruitment and Data collection</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Data analysis</td>
<td></td>
<td></td>
<td>✔</td>
<td>✔</td>
</tr>
</tbody>
</table>

**Estimated Budget ($K)**
- CY 14: $647k
- CY 15: $804k
- CY 16: $796k
- CY 17: $741k

Updated: October 25, 2016

Goals/Milestones

- **CY 14 Goal** – Study preparation
  - IRB submittal
  - Preparation of study materials
  - Clinician Training with Island Quest system
- **CY 15 Goal** – Study preparation and staff training
  - Refine software to be aligned with initial treatment conceptualization
  - HRPO submittal
  - Training study staff in testing and intervention procedures
  - Initiate participant recruitment
- **CY 16 Goal** – Data collection
  - Recruit and test 52 participants from KF/KIR, 15 participants from VANJHCS and 22 participants from NICoE ISO
- **CY 17 Goal** – Data collection
  - Recruit and test 53 participants from KF/KIR, 15 participants from VANJHCS and 23 participants from NICoE ISO
  - Data analysis and Dissemination

Comments/Challenges/Issues/Concerns

- Invested additional time enhancing intervention software prior to initiating clinical trial. Modifications are completed and recruitment and enrollment have started.