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TITLE: Novel strategies to improve immunomodulation and non-invasive clinical monitoring in VCA

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Novel strategies to improve immunomodulation and non-invasive clinical monitoring in VCA

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- **14. ABSTRACT**: Safely minimizing the risks associated with vascularized composite allotransplantation (VCA) is crucial for functional restoration of wounded warriors. Our overarching goal is enabling functional and aesthetic restoration to patients with severe, unreconstructable vascularized composite tissue defects by safe VCA protocols with minimal side effects. Our specific aims are: (1) Establishing the efficacy of a low-dose IL-2 protocol at enabling minimalization of immunosuppression to sirolimus monotherapy in recipients of VCA. (2) Exploring correlations between cellular and molecular immunoassays performed in specimens from VCA recipients (and their donors) with clinical observations of stability and rejection. In future trials, these assays can be developed into tools that prospectively predict rejection and tolerance in VCA, and (3) Implementing next-generation methods to supplement and potentially overcome limitations of established methods such as histology and ultrasound biomicroscopy (UBM). We are enrolling 5 subjects for VCA. <3 months after VCA, once recipient and allograft are stable, we will administer an IL-2 based protocol intended to enable minimalization of immunosuppression to sirolimus monotherapy. Afterwards, immunosuppression will be weaned.
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1. Introduction

Many individuals lose parts of their faces, their limbs or their abdomen in traumatic incidents such as active combat, burns, gunshot wounds, violent attacks, and motor vehicle accidents, amongst others. People with these types of traumatic injuries have decreased quality of life, and are often disabled. Although they may receive the best of the available conventional reconstruction therapies, they continue to suffer from chronic pain, psychological distress, social isolation, and limitations in their ability to perform daily activities such as bathing, dressing, ambulating, and eating without substantial help. “Vascularized composite allotransplantation”, or “VCA” for short, is a new promising therapy for these types of patients. Face transplants, hand transplants and abdominal wall transplants are examples of types of VCA.

The most significant disadvantage of VCA is that patients who receive this therapy must take immunosuppressive drugs for the rest of their lives in order to prevent their bodies from rejecting the transplant. Immunosuppressive drugs pose significant health risks. As VCA is not a life-saving therapy, the risks of immunosuppressive drugs are given much more consideration than in the case of, for example, a heart transplant. Therefore, many people who would benefit from VCA end up not receiving the therapy due to concerns about immunosuppression. We have developed a novel, safe treatment that may enable patients who receive VCA to drastically reduce or even completely eliminate immunosuppressive drugs in the months after transplantation. The objective of this study is to test this novel treatment in 5 patients who will receive VCA. At least 3 months after their VCA operations, our patients will receive our novel treatment which is based on low doses of “interleukin-2” or “IL-2” for short, over a period of 3-4 months. After receiving IL-2 treatment, we will try to minimize or possibly stop immunosuppressive drugs in our patients. If, however, we see signs of rejection, we give standard immune suppression back, which stops rejection successfully in the vast majority of VCA patients. We will follow the progress of our patients for 24 months thereafter. Using state of the art molecular, cellular and imaging technologies, we monitor the subjects’ immune status to identify patients who can safely minimize immune suppression and those who are likely to suffer rejection.

VCA will give many patients the opportunity to improve their quality of life and regain social participation and independence. Our study is carefully designed to thoroughly inform the patients about risks and benefits of participation, to minimize the incidence of complications, and if it is not possible to avoid them, have a safe treatment plan.
2. **Key Words:**

Vascularized composite allotransplantation, immune modulation, immune tolerance, IL-2

3. **Accomplishments.**

Our main accomplishment this year was to obtain IRB approval of a new protocol designed to precisely fit the statement of work in this award, in order to proceed with recruitment of subjects for this study. We then submitted the package for HRPO approval, and received feedback from HRPO on a few requested modifications.

In the meantime, we have:

- Reported to the FDA and maintained IND approval for use of IL-2 in this patient population
- In a study funded by AFIRM II, which administers IL-2 to patients who ALREADY had VCA interventions (i.e. retrospective trial), we have learned important lessons and made important scientific findings based on the recruitment and trial of one individual subject. These lessons/findings are expected to result in improvements to the study protocol that will need to be approved by the IRB, FDA and HRPO.

In addition, we have kept up with our monthly teleconference calls with the sponsor, as well as maintained up to date reporting requirements.

4. **Impact**

Active combat is inflicting multiple devastating injuries to unprotected body areas such as the face and limbs with alarming incidence, and resulting in limb amputation, facial disfigurement, and loss of abdominal wall. Conventional reconstructive surgery is limited in its ability to restore form and function after these injuries. Disability with associated long-term medical care and disability benefit costs is common. Considering the high incidence and devastating consequences of these complex injuries to American Service members, there is a clear need to improve their treatment outcomes. Vascularized composite allotransplants provide a mean to functionally and cosmetically restore these tissues; however, at the cost of lifelong immunosuppression. If successful, these studies will facilitate induction of immunologic tolerance to the transplanted tissues thus improving the rate of return to duty, deployment and function of American service members and veterans recovering from combat-related limb loss, with associated improvements in quality of life, mental health, social participation and the Americaan economy.
5. Changes/Problems
Our most significant roadblock in this project has been to elucidate a path to HRPO approval of this study. We have addressed this issue by submitting an entirely new IRB protocol that is specific and inclusive only of the work stated in the Statement of Work for this award. This protocol has been approved by our IRB, and submitted to the HRPO.

6. Products
Nothing to report at this time.

7. Publications, Abstracts and Presentations
We presented at the Department of Defense’s meeting in December 2016.

8. Inventions, Patents and Licenses
Nothing to report at this time.

9. Reportable Outcomes
Nothing to report at this time.

10. Other Achievements
Nothing to report at this time.

11. Participant and other collaborating organizations
Our collaborations with the Massachusetts Institute of Technology and the Beth Israel Medical Deaconness Center remain in place and active; we have obtained ceded review from their institutional IRBs so that their contribution to our studies remains under oversight by the Partners Human Research Committee.

12. Special Reporting Requirements
None.
13. Appendices

None.
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W81XWH-15-2-0031
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PI: Bohdan Pomahac, MD
Org: Brigham and Women’s Hospital
Award Amount: $2,487,729

**Study/Product Aim(s)**
- **SA1.** To develop a safe and feasible regimen for minimization of immune suppression in recipients of VCA through daily subcutaneous low-dose rIL-2.
- **SA2.** To explore correlations between cellular and molecular immune markers in VCA and clinical observations of immune stability and rejection.
- **SA3.** To develop non-invasive technologies to monitor for VCA rejection, such as next-generation MR methods.

**Approach**
Exploratory, open-label, prospective safety and feasibility clinical trial that will enroll 5 candidates for VCA.
Five subjects will be recruited and enrolled for VCA. Following VCA, they will receive an IL-2 drug protocol. Specimens and imaging data from these VCA recipients will be used towards SA2 and SA3.

**Timeline and Cost**

<table>
<thead>
<tr>
<th>Activities</th>
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<th>CY 18</th>
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<td>Task 2. Enrollment of 5 subjects</td>
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<td>Task 3. VCA surgeries</td>
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<td>Task 4. Administration of IL-2 protocol</td>
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**Goals/Milestones**
- **CY17 Goal** – IRB/HRPO approval
  - ☑ IRB approved
  - ☑ HRPO submitted
- **CY17 Goal** – Enrollment of 5 subjects
  - ☐ Informed consent
  - ☐ Screening
- **CY18 Goal** – VCA surgeries
  - ☐ 5 subjects
- **CY18 Goal** – Administration of IL-2
  - ☐ In 5 subjects

**Comments/Challenges/Issues/Concerns**
- Timelines changed with respect to original proposal because of delays in obtaining IRB/HRPO approval.
- Off in spending due to delays described above.

**Budget Expenditure to Date**
Projected Expenditure: $1,658,486
Actual Expenditure: $734,548

Updated: August 2017