Award Number: W81XWH-09-1-0722

TITLE:
High Risk Suicidal Behavior in Veterans- Assessment of Predictors and Efficacy of Dialectical Behavioral Therapy

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Approximately one third of the Army’s completed suicides last year occurred in the post-deployment period (Alvarez 2009) highlighting the importance of studying high-risk suicidal veteran populations. This project proposes two related studies. The first project is a randomized clinical trial of 120 veterans identified with high-risk suicidal behavior comparing the efficacy of Dialectical Behavioral Therapy (DBT) vs. treatment as usual (TAU) on suicidal behavior as a primary outcome measure. A second aim of the project is to examine group differences between 150 veterans at high risk (HR) for suicide and 150 veterans at low risk (LR) in a variety of symptom domains.

The randomized clinical trial was completed with 93 subjects. Results were notable for improvements in suicidality and secondary outcomes of depression, hopelessness and anxiety in both treatment groups, however there was not a significant difference between the treatment groups. DBT subjects did receive significantly more hours of treatment and for homeless, substance abusing males, there was considerable early drop-out in both arms. Our supplemental funding is to study affective startle as potential marker of suicide risk and as a marker of treatment outcome. Affective startle does significantly differ between Veteran ideators, single and multiple attempters.

We collected additional female Veteran samples in a no-cost extension in order to perform gender sub-analyses.

Borderline Personality Disorder, SUICIDE
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Introduction:

Approximately one third of the Army’s completed suicides last year occurred in the post-deployment period (Alvarez 2009), highlighting the importance of studying high-risk suicidal veteran populations. This project proposes two related studies. The first project is a randomized clinical trial of 120 veterans identified with high-risk suicidal behavior comparing the efficacy of Dialectical Behavioral Therapy (DBT) vs. treatment as usual (TAU) on suicidal behavior as a primary outcome measure. A second aim of the project is to examine group differences between 150 veterans at high risk for suicide and 150 veterans at low risk in a variety of symptom domains. The goal of this will be to identify symptoms associated with suicidal behavior that may advise future treatment.

We will assess symptom domains including mood and substance use in our veteran population by comparing symptoms in low vs. high risk veterans recently discharged from the James J Peters VAMC (JJPVAMC) psychiatric inpatient unit. In addition, we will explore indices of interpersonal function and measure features that have some evidence of offering protection from suicide, which could be viewed as resilience factors. A particular emphasis of the present project is to characterize the nature of the interpersonal dysfunction in high risk individuals, as there exists very good evidence that social isolation, or a lack of a sense of “belonging” puts people at particularly high risk for suicide, in particular in a military sample. We intend to assess the impact of DBT vs. TAU on these symptom domains in addition to their impact on suicidal behavior.

Body:

In October 2011, a supplement to this project was approved to add a physiological measure, affective startle, to the baseline assessment and post- DBT treatment.

Aim 1 relates to a randomized clinical trial of Dialectical Behavior Therapy (DBT) vs. treatment as usual (TAU) in 120 veterans recently hospitalized with high-risk suicidal behavior. This will be accomplished under the leadership of Dr. Marianne Goodman, James J. Peters VAMC, Bronx, NY 10468

Aim 1: To examine, in a randomized controlled trial (RCT), the efficacy of a 6 month treatment with standard DBT (weekly individual sessions, skills training group and telephone coaching as needed) as compared to TAU in 120 veterans recently discharged from an acute psychiatric inpatient stay with high risk suicidal behavior. The primary treatment outcome will be a quantification of suicidal events, as assessed by the Columbia Suicide Severity Rating Scale, which measures suicide attempts, plans and preparations. Our study will be powered to examine treatment assignment differences in this measure. Secondary outcomes will include suicidal ideation, parasuicidal events, treatment compliance, depressed mood, substance abuse and hopelessness.

This aim involves recruiting 120 veterans off the JJPVA “high-risk” suicide list; a designation made primarily after psychiatric inpatient admission for serious suicidal behavioral. High-risk (HR) suicide subjects will undergo a comprehensive diagnostic interview prior to entering the treatment study. Subjects will receive 6 months of TAU vs. DBT but both groups will continue to receive standard psychopharmacology and case management services from their clinic providers. Subjects will receive a battery of assessments at month 6, 12 and 18.

Aim 2 relates to a comparison of high-risk and low-risk suicidal veterans in interpersonal functioning and resilience, in an effort to identify intermediate symptoms that are closely associated with HR suicidal behavior. This will be accomplished under the leadership of Dr. Marianne Goodman, James J. Peters VAMC, Bronx NY 10468

Aim 2: To recruit veterans recently discharged from an acute psychiatric inpatient stay comparing 150 veterans with HR suicidal behavior to 150 veterans without such behavior (LR) in symptom domains focusing on interpersonal functioning and resiliency.
Aim 3 is exploratory and examines the effect of treatment (DBT or TAU) on the putative intermediate symptom domains associated with HR suicidal behavior of interpersonal functioning and resiliency. This will be accomplished under the leadership of Dr. Marianne Goodman James J. Peters VAMC, Bronx NY 10468

**Aim 3:** To explore the effect of DBT on the candidate intermediate symptoms of interpersonal functioning and resiliency associated with HR suicidal behavior.

**Promised work:**

**Parent Project**

The first 3 months is devoted to training the raters on our assessment and diagnostic battery while we await regulatory approvals. During months 3-6, we expect to perform thirty baseline assessments and 15 high-risk subjects will be randomized to treatment. During months 6-12, 12-18, 18-24, 24-30, we expect that thirty high-risk and thirty low-risk suicidal subjects will receive baseline assessments during each 6 month block. We anticipate that 25 of the high-risk subjects will proceed into treatment during each one of the time blocks. Months 30-36 will target 30 total additional assessments for baseline high and low-risk subjects with 5 of the HR individuals being randomized for treatment. The baseline assessment is a more comprehensive evaluation and we estimate that it will take approximately 6-7 hours with follow-up assessments requiring 1-2 hours.

While we met recruitment goals for Aim 1 of the study, our recruitment for the RTC fell behind. In order to continue recruitment we requested and were granted a fifth year, no cost extension. The Table below reflects promised work, and new numbers with the 5th year added.

<table>
<thead>
<tr>
<th>Months 0-3</th>
<th>Baseline assessments (50% HR, 50% LR)</th>
<th>Randomized to treatment (HR only)</th>
<th>Follow-up assessments 6mo 12 mo 18mo</th>
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<tr>
<td>Months 4-6</td>
<td>30</td>
<td>15</td>
<td>----</td>
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<td>Months 7-12</td>
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<td>25</td>
<td>12</td>
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<tr>
<td>Months 13-18</td>
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<td>19 11</td>
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<td>19 17 10</td>
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<td>Months 25-30</td>
<td>60</td>
<td>25</td>
<td>19 17 15</td>
</tr>
<tr>
<td>Months 31-36</td>
<td>30 –</td>
<td>5 –</td>
<td>19 17 15</td>
</tr>
<tr>
<td>Months 37-48 (year 4)</td>
<td><strong>New year 4 target- 60</strong>&lt;br&gt;Actual 65&lt;br&gt;293 to date (goal 300)</td>
<td><strong>New year 4 target-25</strong>&lt;br&gt;Actual 15&lt;br&gt;90 to date (goal 120)</td>
<td>28 19 30</td>
</tr>
<tr>
<td>Months 49-60 (year 5)</td>
<td><strong>New year 5 target- 10</strong>&lt;br&gt;Recruitment just to meet RTC goals</td>
<td><strong>New year 5 target-10</strong>&lt;br&gt;15</td>
<td>15 20 30</td>
</tr>
</tbody>
</table>

**Progress Parent Study:**
IRB approval
Towards accomplishing these aims, we received approval from our local IRB 7/9/09 and local Research and Development approval on 7/15/2009; prior to official funding of the project. This allowed us to pilot the intervention, assessments and randomization procedure. Dept of Defense approval was obtained on 4/27/2010; almost four months later that we had projected in our initial statement of work.

Recruitment
No-cost extension year; During our no-cost extension year, we focused solely on increasing our female Veteran subject pool in order to be able to do gender analyses. In addition, we realized that our LR pool of female veterans needed to be augmented. During this time period, we recruited and successfully assessed an additional 11 females who completed our baseline assessment and affective startle paradigm.

For the treatment trial, during our no cost extension year, we completed follow-ups.

Over the entire course of the study, 384 veterans have signed consent (248 high-risk subjects and 136 low-risk subjects) and 340 (221 high-risk and 119 low-risk) have completed baseline assessments. With these numbers, we have surpassed our goal of 300 completed assessments. However, recruitment for the treatment trial remained limited and we randomized a total of 93 subjects to the trial and 53 completed the 6-month treatment.

Summary of Entire Parent Study to date

<table>
<thead>
<tr>
<th></th>
<th>High Risk</th>
<th>Low Risk</th>
</tr>
</thead>
<tbody>
<tr>
<td># consented</td>
<td>248</td>
<td>136</td>
</tr>
<tr>
<td># completed</td>
<td>221</td>
<td>119</td>
</tr>
<tr>
<td># randomized</td>
<td>93</td>
<td></td>
</tr>
<tr>
<td>#complete 6 month</td>
<td>53</td>
<td></td>
</tr>
<tr>
<td>12 month f/up</td>
<td>49</td>
<td></td>
</tr>
<tr>
<td>18 month f/up</td>
<td>43</td>
<td></td>
</tr>
</tbody>
</table>

Randomized Treatment Trial:
Over the course of the project- 93 subjects have been randomized (to 45 TAU/48 to DBT). 53 subjects have completed the 6-month treatment trial; 25 for DBT and 28 for TAU.

Longitudinal Follow-up at 12 and 18 months:
49 subjects have completed 12-month follow up to date including- 22 DBT subjects and 27 TAU subjects and 43 subjects have completed the entire study (e.g. 18-month completers) including 19 DBT and 24 TAU subject.

Total: 340 of 300 completed baseline assessments (exceeded goals)
93 of 120 randomized to clinical trial

Progress Pertaining to Aim #1
Our Statement of work projected that by study completion we will have 300 baseline assessments finished. We completed 340, and during our no-cost extension year targeted females in order to perform gender sub-analyses. Data analysis was the focus of our no-cost extension year and we generated several manuscripts. Interim analyses have yielded findings pertaining to the importance of Axis I diagnoses of substance abuse, Axis II diagnoses of borderline personality disorder and responses on the interpersonal psychological survey as important risk factors for identifying "high-risk" veterans (see Figure 1).
The identification of the interpersonal psychological survey as a critical instrument has led us to further examine its contents through a computerized implicit task assessment that we will be piloting in year 5 (see Figure 2).

### Figure 1- Predictors of Suicide Risk in Veterans

#### Table 1: Logistic Regression predicting high-risk vs. low-risk subjects using diagnostic variables and self-report measure variables

<table>
<thead>
<tr>
<th>Predictor</th>
<th>B</th>
<th>S.E.</th>
<th>Wald</th>
<th>df</th>
<th>Sig</th>
<th>Exp(B)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPS score (Interpersonal psychological survey)</td>
<td>.215</td>
<td>.057</td>
<td>14.126</td>
<td>1</td>
<td>.000</td>
<td>1.239</td>
</tr>
<tr>
<td>SIDP, Borderline Personality Disorder, diagnosis</td>
<td>2.394</td>
<td>.631</td>
<td>14.407</td>
<td>1</td>
<td>.000</td>
<td>10.955</td>
</tr>
<tr>
<td>Constant</td>
<td>-4.066</td>
<td>.958</td>
<td>18.024</td>
<td>1</td>
<td>.000</td>
<td>.017</td>
</tr>
</tbody>
</table>

### Figure 2- Interpersonal Psychological Survey components

Progress Pertaining to Aim #2
We have randomized 93 subjects to the treatment trial and completed data analyses. Results are pictured below. Our findings indicate that both dialectical behavior therapy (DBT) and enhanced treatment as usual (TAU) were beneficial to suicidal Veterans, but there were no significant differences between treatment groups in our primary or secondary outcome measures (see Aim 2 Figures 1-5). DBT utilized considerably more treatment sessions (see Aim 2-Figure B). Survival Curves for suicide ideation and re-hospitalization also showed no differences (see Aim 2 Figures 6-7). These results were accepted for publication, citation below:

Aim 2- Figure A: # weeks in Treatment

Weeks in TAU: 28 Completed
Weeks in DBT: 25 Completed

Aim 2- Figure B: # Mental Health Visits by Treatment Group

Mental Health Individual Visits

- Significant Difference
  - F=23.012
  - p<.001
- DBT
  - Mean=34.8
  - SD=8.5
- TAU
  - Mean=15.0
  - SD=6.5

Aim 2- Figures 1-5:
Primary Outcome Measures by Treatment Group:
- Figure 1- Suicide Symptomatology (measured by the CSSRS)
- Figure 2- Suicide Ideation (measured by the Beck Suicide Ideation scale)

Secondary Outcome Measures by Treatment Group:
- Figure 3- Depression (measured by the Beck Depression Scale)
- Figure 4- Hopelessness (measured by the Beck Hopelessness Scale)
- Figure 5- Anxiety (measured by the Beck Anxiety Scale)

** Note improvements across time in both treatments, however no significant difference between treatment groups
ADDITIONAL ANALYSIS- SURVIVAL ANALYSES

Aim 2 - FIGURES 6 and 7:
Survival Analysis – Suicide Ideation
Survival Analysis- Re-Hospitalizations

** Note almost identical survival curves between the two treatment groups. Note very high rate of re-hospitalization.

### Survival Analysis: Suicidal Ideation

DBT 50% w/o ideation @ 18 months
TAU 45% w/o ideation @ 18 months

### Survival Analysis: Hospitalizations

Both treatment groups @ about 51% w/o hospitalizations.
Progress Pertaining to Aim #3
There were no significant differences found between treatment groups in the domains of interpersonal functioning nor resiliency.

Problems Accomplishing Tasks

With Hurricane Sandy, the Manhattan VA hospital was closed for upwards of 5 months. This lead to disruptions of care at our facility, as Manhattan patients sought treatment temporarily at our hospital. This complicated RCT recruitment efforts as pts were less likely to enroll in a longitudinal study that would require changing the location of their outpatient care beyond the expected time of Manhattan VA’s closure.

SUPPLEMENT:

In addition to our three aims for the parent study, we added three additional aims for the supplemental study:

Supplement Aim 1 is to conduct a nonverbal and objective psychophysiological assessment of emotion processing using the affective startle paradigm to test whether it might serve as a potential biomarker for differentiating levels of suicidality. This will be accomplished under the leadership of Drs. Marianne Goodman and Erin Hazlett, James J. Peters VAMC, Bronx NY 10468

Aim 1. To examine the magnitude, time course, and rate of habituation of the startle eyeblink response during unpleasant, neutral, and pleasant pictures in 150 veterans with varying levels of suicidality; 60 veterans with a recent suicide attempt (during past 3 months), 60 veterans with suicidal ideation but no history of attempts, and 30 healthy veteran controls (i.e. no current psychiatric diagnosis).

This aim will be accomplished by adding the affective startle modulation paradigm to our current assessment battery of high- and low-risk suicidal subjects. Eligible subjects enrolled in the DoD funded parent project will participate in a 1-hour psychophysiology session at the MIRECC psychophysiology laboratory where we will record our primary variable of interest, namely the affective startle eyeblink response at baseline and 6 months for those enrolled in the DoD treatment trial. During this session, participants will view an intermixed series of unpleasant, neutral, and pleasant pictures from a standardized picture set. For each of the 3 picture conditions, we will examine three measures related to affective startle eyeblink modulation which is our psychophysiological measure of emotion processing: (1) the amplitude of the startle eyeblink response; (2) the time course of emotion processing by presenting the startle probes at different times during and post-picture processing; and (3) the rate of habituation of the startle eyeblink response.

Supplement Aim 2 is to compare startle variables across suicide groups (ideators, attempters) by presence or absence of borderline personality disorder to clarify if differences in affective startle modulation extend beyond personality disorder diagnosis. Thirty suicide attempters with BPD (SABPD+) will be compared with 30 suicide attempters without BPD (SABPD-) and 30 suicide ideators with BPD (SIBPD+) will be compared to 30 ideators without BPD (SIBPD-) across startle variables. This will be accomplished under the leadership of Drs. Marianne Goodman and Erin Hazlett, James J. Peters VAMC, Bronx NY 10468

Aim 2 investigates the relationship of Axis II diagnosis, suicidality and affective startle. The collected data for Aim 2 will be used to explore this question.

Supplement Aim 3 (No-Cost Extension) is to explore gender differences between male and female veterans regarding affective startle modulation and using lifetime number of suicide attempts as a continuous variable of interest. Fifty female veterans will be compared with fifty male veterans for this analysis. This will be accomplished under the leadership of Drs. Marianne Goodman and Erin Hazlett, James J. Peters VAMC, Bronx NY 10468
**Aim 3** investigates the relationship of gender, suicidality and affective startle. We will investigate potential differences in affective startle response across gender and suicide status.

**Supplement Promised Work:**

<table>
<thead>
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<tr>
<td>Startle assessments:</td>
<td></td>
</tr>
<tr>
<td>Pt  Pt  HC  HC</td>
<td></td>
</tr>
<tr>
<td>Baseline 6mo  Baseline 6mo</td>
<td></td>
</tr>
<tr>
<td>Months 13-18</td>
<td>Obtain IRB approval</td>
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<tr>
<td>Months 19-24</td>
<td>45  -----  12  ----</td>
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<tr>
<td>Months 25-30</td>
<td>50  23 *  12  10</td>
</tr>
<tr>
<td>Months 31-36</td>
<td>25  22 *  6  11</td>
</tr>
<tr>
<td>Months 37-42</td>
<td>5  *  ----  4</td>
</tr>
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</table>

The project was awarded funding on 9/24/2011. In its first 12 months, the research team was incredibly effective in mobilizing resources and enrolling and testing 94 subjects at baseline and completing 1 6-month follow-up. Over the next 12-month period (10/1/12 → 9/30/13), the team was similarly effective with recruitment, testing an additional 55 subjects at baseline and 13 at 6-months following the treatment trial. In the 12-month period (10/1/13 → 9/30/14), we have tested additional 21 subjects at baseline and 9 at 6-months. The vast majority of these new participants were female (18/19), as we redoubled efforts to recruit from this population at the JJPVAMC. During the no-cost extension year, we added 6 additional females. These figures bring the cumulative total of baseline and 6-month numbers to 177 subjects and 23 subjects respectively.

**Progress Pertaining to Supplement Aim #1**

Since receiving funding, we have run 177 patients at baseline, exceeding our goals, and have done 23 6-month follow-ups and therefore have met our recruitment goals for supplement Aim #1. The overall breakdowns are as follows:

<table>
<thead>
<tr>
<th>Group</th>
<th>Recruitment - Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controls</td>
<td>38 (9F/29M)</td>
</tr>
<tr>
<td>Ideators</td>
<td>46 (14F/32M)</td>
</tr>
<tr>
<td>Single Attemptors</td>
<td>44 (14F/30M)</td>
</tr>
<tr>
<td>Multiple Attemptors</td>
<td>49* (23F/26M)</td>
</tr>
<tr>
<td>6-Month Follow-Up</td>
<td>23 (6F/17M)</td>
</tr>
</tbody>
</table>

*Four patients reclassified as a Single Attemptors

Preliminary analyses on the first 108 subjects demonstrated a significant interaction between affective startle and suicide risk. (see Supplement figure 1). Multiple ideators, in the unpleasant picture condition, had significantly elevated affective startle % change values as compare to single attempters and ideators (both active and passive).
Supplement/Figure 1.
Affective Startle and Suicide Risk in veterans with passive ideation, active ideation, and a history of single and multiple suicide attempts (preliminary data, n=108).

Supplement Figure 2.
The data was also analyzed by collapsing suicide groups to ideators vs. attempters. Using a larger data set of 126 of the Veterans, attempters demonstrated significantly greater levels of depression, anxiety, borderline PD symptoms and emotion dysregulation scores (see demographic slide).

Supplement Figure 3.
Affective startle in attempters vs. ideators
Startle amplitude to emotional pictures in suicide attempters was significantly different from ideators in the unpleasant picture condition only.
Supplement Figure 4. Correlations with Affective Startle. This figure shows that the affective startle finding of increased amplitude to unpleasant pictures collapsed across all subjects correlates with the number of suicide attempts and increased scores of the difficulty of emotion regulation scale. These findings suggest that affective startle may be a biomarker of suicide risk.

**In addition, using data from the treatment trial participants who received affect startle, amplitude to unpleasant pictures was used prospectively to examine prospective suicide attempt risk. Increased amplitude to unpleasant pictures was significantly associated with prospective suicide attempt in a small sample of treatment participants.**

Progress Pertaining to Supplement Aim # 2
See information pertaining to Aim #1, specifically figure #4 above.

Progress Pertaining to Supplement Aim # 3 (NCE)
A newly added Aim for our approved No-Cost Extension was to continue to gather data from female participants. Female sub-analyses are still being conducted.

Problems Accomplishing Tasks
We did not experience any difficulty recruiting for this project and are in fact were ahead of schedule.

Key Research Accomplishments for both Parent and Supplement Projects
We have just completed year 5 of the parent study and no-cost extension for the affective startle supplement.

- Since DoD IRB approval (4/27/10): parent subject recruitment has been brisk. 384 Veteran subjects have signed consent.
- 340 (out of promised 300) subjects completed baseline assessments.
- 93 HR patients were randomized to the treatment trial,
- 53 HR patients have completed the 6-month treatment trial,
- 177 subjects have completed baseline affective startle
- Treatment trial results suggest that DBT is no more effective than enhanced treatment as usual for suicidal Veterans.

** however both groups had significant drop out rates and difficulty with engagement.
Future directions for research suggest alternative modalities for early engagement post hospitalization.

- Affective startle is a potential biomarker of suicide risk.
- Preliminary data shows affective startle may predict prospective suicide attempts.

Reportable Outcomes

Publications

Treatment trial results:

Data from the baseline study:

Data from the supplement study:

Dissemination/Presentations

1) American Psychiatric Association, May 2012
2) DOD/VA Joint Conference Suicide Prevention, June 2012
3) Society Psychophysiology, Research, September 2012*, Sept 2014*,
4) Veterans Integrated Service Network (VISN) 3 Conference on Addressing Mental Health Needs of OEF/OIF Soldiers, October 2012
5) American Psychiatric Association, May 2013
6) North American Society for Personality Disorders (NASSPD), April 2013* (2 presentations), 2015

Posters

1) International Society of Psychoneuroendocrinology (ISPNE) special meeting on Biomarkers of PTSD, September 2012*
2) NASSPD, April 2013, April 2014, March 2015*
3) Biological Psychiatry May 2013*, May 2014*
4) International Society Psychophysiology, September 2013*

- DoD supplement

Conclusion:
Our preliminary baseline data highlights the importance of Axis II psychopathology, in particular, borderline personality disorder as a risk factor for high-risk suicidal behavior. This is relevant as the disorder is often
under recognized in VA settings and not even listed in the Uniform Services Package, the document listing required services for Veterans.

Enhanced Treatment as Usual and DBT are both effective treatments for suicidal veterans, however there is no significant difference between treatments and DBT results in enhanced treatment delivery needs. Future directions for research should be targeted to better engagement of high risk suicidal veterans, who do not follow up with outpatient psychotherapy treatments as both groups had high drops out in this subset. These individuals were primarily male, homeless and abusing substances.

Regarding our supplemental funding for affective startle, a psychophysiological assessment tool, there does appear to be a significant relationship between the affective startle response to negative picture probes and number of suicide attempts. These results suggest that the processing of negative emotion is a critical element of suicide attempt history. Future studies using affective startle include whether this measure can predict those at suicide risk prospectively. We are currently piloting a portable version of the affective startle assessment which will allow us to assess individuals in different setting (e.g. emergency room, inpatient unit, off campus locations).

We continue to analyze our data for gender differences.

References:

Appendices: none included

Supporting Data: none included
Affective Startle Modulation in Suicidal Veterans
DMRDP Proposal Number WX81XWH-09-1-0722

PI: Goodman, Marianne
Org: James J. Peters VAMC, Bronx NY
Award Amount: $452,834

Study/Product Aim(s)

Aim 1 is to examine the magnitude, time course, and rate of habituation of the startle eye blink response during unpleasant, neutral, and pleasant pictures in 150 veterans with varying levels of suicidality; 60 veterans with a recent suicide attempt (during past 3 months), 60 veterans with suicidal ideation but no history of attempts, and 30 healthy controls.

Aim 2 is to compare startle variables across suicide groups (ideators, attempters) by presence or absence of borderline personality disorder to clarify if differences in affective startle modulation extend beyond personality disorder diagnosis. Thirty suicide attempters with BPD (SABPD+) will be compared with 30 suicide attempters without BPD (SABPD-) and 30 suicide ideators with BPD (SIBPD+) will be compared to 30 ideators without BPD (SIBPD-) across startle variables.

Aim 3 is exploratory and will examine whether (a) magnitude, time course and/or rate of habituation to unpleasant, neutral and pleasant pictures predicts treatment response to six-month Dialectical Behavioral Therapy (DBT) for suicidal behavior; and (b) magnitude, time course and/or habituation of affective startle improves with 6 months of DBT in treatment responders compared with non-responders.

Goals/Milestones

CY11 Goal – obtain IRB approval, begin recruitment and assessment of subjects for baseline startle assessment
✓ Assess 25 high risk and 12 healthy control subjects with startle paradigm by end of calendar year (revised to 10 HR/5 LR due to 5 month delay in funding)

CY12 Goals – ✓ complete data collection of baseline startle assessments
✓ Complete startle assessments of suicidal subjects (total n=100)
✓ Begin to obtain 6 month follow-up startle

CY13 Goal – Complete data collection of 6 month startle assessments and data analysis (we have collected 177 of 150)
✓ Complete data collection on 6-month follow up startle assessments (collected 23 to date)
✓ CY14 Goal- finish startle follow-ups, prepare startle data for analysis and conduct analyses. Manuscript generation (paper submitted and under review)

Activities

<table>
<thead>
<tr>
<th>Activities</th>
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<th>CY 12</th>
<th>CY 13</th>
<th>CY 14</th>
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<td>Obtain IRB approval, start data collection</td>
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<tr>
<td>Complete baseline startle assessments</td>
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<tr>
<td>Complete follow-up startle assessments</td>
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<tr>
<td>Data preparation and analysis</td>
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<tr>
<td>preparation of manuscripts and presentations</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
</tbody>
</table>

Estimated Budget ($K) indirect and direct

| Estimated Budget ($K) indirect and direct | $168 | $142 | $143 |

Updated: 4/2014

Accomplishment: The supplement funding was not received until after month 24, five months later than the original target date. However, IRB approval was accomplished rapidly by month 26 and study recruitment commenced shortly afterward. We have currently assessed 177 baseline subjects and 23 6-month follow-up. Data analysis is complete, manuscript has been developed and currently under review for publication. Multiple presentations have occurred.

Comments/Challenges/Issues/Concerns- n/a

Budget Expenditure to date

Projected Expenditure: $452,834 total
Actual Expenditure: $452,759
**Aim 1:** To examine, in a randomized controlled trial (RCT), the efficacy of a 6 month treatment with standard DBT (weekly individual sessions, skills training group and telephone coaching as needed) as compared to TAU in 120 veterans recently discharged from an acute psychiatric inpatient stay with high risk suicidal behavior. High-risk (HR) suicide subjects will undergo a comprehensive diagnostic interview prior to entering the treatment study. Subjects will receive 6 months of TAU vs. DBT but both groups will continue to receive standard psychopharmacology and case management services from their clinic providers. Subjects will receive a battery of assessments at month 6, 12 and 18.

**Aim 2:** To recruit veterans recently discharged from an acute psychiatric inpatient stay comparing 150 veterans with HR suicidal behavior to 150 veterans without such behavior (LR) in symptom domains focusing on interpersonal functioning and resiliency.

**Aim 3:** is exploratory and examines the effect of treatment (DBT or TAU) on the putative intermediate symptom domains associated with HR suicidal behavior of interpersonal functioning and resiliency.

<table>
<thead>
<tr>
<th>Activities</th>
<th>CY 10</th>
<th>CY 11</th>
<th>CY 12</th>
<th>CY 13</th>
<th>CY 14</th>
</tr>
</thead>
<tbody>
<tr>
<td>Obtain IRB approval, start data collection</td>
<td>✔</td>
<td>✔</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Complete 300 baseline assessments</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
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</tr>
<tr>
<td>Perform randomized clinical trial (n=120)</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
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<tr>
<td>Complete follow-up assessments</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
</tr>
<tr>
<td>Data preparation and analysis</td>
<td>✔</td>
<td>✔</td>
<td>✔</td>
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<td>✔</td>
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<tr>
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**Estimated Budget (K)**

<table>
<thead>
<tr>
<th></th>
<th>CY 10</th>
<th>CY 11</th>
<th>CY 12</th>
<th>CY 13</th>
<th>CY 14</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indirect &amp; Direct</td>
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<td>$339K</td>
<td>$353K</td>
<td>$376K</td>
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</tbody>
</table>

**Goals/Milestones**

- **CY10 Goal** – obtain IRB approval, hire and train staff, begin recruitment targeting 90 baseline assessments and 40 randomized to treatment trial. Initial DoD IRB approval was delayed 5 months behind projected date. For the remainder of 2010, we completed 53 baseline assessments and randomized 23 high risk suicidal veterans for treatment.

- **CY11 Goal** – continue data collection of baseline assessments (n=120) and randomization to clinical trial (n=50), and follow up assessments (n=76). Actual numbers were 96 consented subjects, 66 completed baseline assessments and 26 randomized to the tx trial.

- **CY12 Goal** – continue data collection of baseline assessments (n=90) and randomization to clinical trial (n=30), and follow up assessments (n=102).

- **CY 13 Goal** - continue data collection of baseline assessments (n=75) and 50 subjects randomized to RTC by increasing efforts at our second recruitment site (Manhattan VAMC). Recruitment to date thru 2013 - we have completed 329 baseline assessments and randomized 93 subjects to the treatment trial. Our recruitment for baseline assessments has increased due to addition of the second site, but clinical trial enrollment remains behind.

- **CY 14 Goal** - Complete data collection and data analysis & manuscript generation.

**Comments/Challenges/Issues/Concerns** - recruitment for RCT as described above, addressed through addition of 5th year and no cost extension.

**Budget Expenditure to date**

- Projected Expenditure:$1.279M
- Actual: (through 3/14) $1.279