Effect of transcutaneous cervical vagus nerve stimulation on the pituitary adenylate cyclase-activating polypeptide (PACAP) response to stress: A randomized, sham controlled, double blind pilot study

Nil Z. Gurel,1 Yunshen Jiao,1 Matthew T. Wittbrodt, Yi-An Ko, Allison Hankus, Emily G. Driggers, Stacy L. Ladd, Lucy Shallenberger, Nancy Murrah, Minxuan Huang, Ammer Haffar, Mhmtjamil Alkhalaif, Oleksiy Levantsevych, Jonathon A. Nye, Viola Vaccaroni, Amit J. Shah, Omer T. Inan, J. Douglas Bremner, Bradley D. Pearce*

a School of Electrical and Computer Engineering, Georgia Institute of Technology, Atlanta, GA, USA
b Department of Epidemiology, Rollins School of Public Health, Emory University, Atlanta, GA, USA
c Department of Psychiatry and Behavioral Sciences, Emory School of Medicine, Atlanta, GA, USA
d Department of Biostatistics and Bioinformatics, Rollins School of Public Health, Emory University, Atlanta, GA, USA
e Department of Radiology, Emory School of Medicine, Atlanta, GA, USA
f Department of Medicine, Division of Cardiology, Emory School of Medicine, Atlanta, GA, USA
g Atlanta VA Medical Center, Decatur, GA, USA
h Coulter Department of Bioengineering, Georgia Institute of Technology, Atlanta, GA, USA

ABSTRACT

Pituitary adenylate cyclase-activating peptide (PACAP) is a neuropeptide that plays a key role in the neurobiology of the stress response, and prior studies suggest that its function is dysregulated in post-traumatic stress disorder (PTSD). Transcutaneous cervical vagus nerve stimulation (tcVNS) acts through PACAP and other neurobiological systems to modulate stress responses and/or symptoms of PTSD. In this pilot study, we examined the effects of tcVNS on PACAP in a three-day chronic stress laboratory paradigm involving serial traumatic and mental stress exposures in healthy individuals with a history of exposure to psychological trauma (n = 18) and patients with PTSD (n = 12).

Methods: A total of 30 subjects with a history of exposure to psychological trauma experience were recruited (12 with PTSD diagnosis) for a three-day randomized double-blinded study of tcVNS or sham stimulation. Subjects underwent a protocol that included both personalized trauma recall and non-personalized mental stressors (public speaking, mental arithmetic) paired to tcVNS or sham stimulation over three days. Blood was collected at baseline and multiple time points after exposure to stressors. Linear mixed-effects models were used to assess changes in PACAP over time (in response to stressors) and its relation to active tcVNS or sham stimulation.

Results: PACAP blood levels increased over the course of three days for both active tcVNS and sham groups. This increase was statistically-significant in the sham group at the end of the second (Cohen’s dₘ = 0.35, p = 0.04), and third days (dₘ = 0.41, p = 0.04), but not in the active tcVNS group (dₘ = 0.21, dₘ = 0.18, and p > 0.20).

Conclusion: These pilot findings suggest tcVNS may attenuate this neurobiological stress-response. Larger studies are needed to investigate gender and interaction effects.

1. Introduction

Post-Traumatic Stress Disorder (PTSD) is a neuropsychiatric condition triggered by experiencing or witnessing a terrifying event, with a lifetime prevalence of 6.4% in the United States [92]. Treatment modalities for PTSD primarily involve trauma-focused psychotherapies including cognitive behavior therapy (CBT) and medication [4,22,36,37,57,106]. A range of medications, which were generally not developed for PTSD specifically, have also shown efficacy in some cases [27,118]. Despite the tremendous burden of PTSD, the treatments are still not

* Corresponding author.
E-mail addresses: nil@gatech.edu (N.Z. Gurel), bpearce@emory.edu (B.D. Pearce).
1 co-first authors.

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widespread, have high rates of non-completion, and/or limitations in efficacy [64,65,77]; U.S. Department of Veterans [Affairs 2019 [111, 118]]. Moreover, regardless of which treatment method is chosen, about 40% of the patients have a recurrence of symptoms within one year, and the risk of five-year recurrence is about 20% [7]. Therefore, new approaches are needed for the treatment of PTSD.

The etiology of PTSD is complex and likely involves interconnected molecular pathways of the sympathetic nervous system, the hypothalamic pituitary adrenal (HPA) axis, inflammatory response systems, and other neuropeptidel and neurohormonal systems [10,87]. Pituitary adenylate cyclase activating polypeptide (PACAP) is a highly conserved neuropeptide that connects these systems and regulates and integrates adaptive responses to stress [47]. A growing body of literature has pointed to dysregulation of PACAP along with its selective PAC1 receptor in PTSD [28,30,47,59,62,78,79,100]. Within the brain, PACAP participates in neural circuits relevant to PTSD and other stress disorders [55,106]. In the hypothalamus and hippocampus, PACAP acts as an important neuromodulator [34,75]. Anatomical and physiological studies established the importance of PACAP neurotransmission in the amygdala, which underscored the role of this peptide transmitter in fear responses and potentially PTSD [93,109]. Studies suggest that PACAP is anxiogenic and may play a role in symptoms of PTSD [55,100]. PACAP also has neuroprotective and anti-inflammatory properties [29,98] and is well-positioned for modulation by vagus nerve activity [88,99].

Electrical stimulation of the vagus nerve, also referred to as vagus nerve stimulation (VNS), has been demonstrated to be efficacious for the treatment of epilepsy [44,54], major depression [1,43,104], heart failure [96], and migraine [76]. VNS also results in improvements in cognition, memory, and adverse conditions such as tinnitus when paired with a stimulus or task [26,31,32,56,95]. Preclinical work suggested that VNS may be an effective adjunct to exposure therapy for the treatment of PTSD [82,89]. Considering the potential side effects and complications of implant surgeries, new medical devices were developed providing noninvasive transcusaterminal stimulation [11,45]. Transcutaneous VNS devices target the auricular (taVNS) or cervical (tcVNS) portions of the vagus nerve. There has been a growing literature in recent years suggesting that these devices modulate central and peripheral physiology, as investigated by brain imaging [2,38,39,116,117], inflammatory serum cytokines [8,17,71], and peripheral physiological measures [34,49,51] in preclinical models or in humans with or without adverse chronic conditions including PTSD and paired with traumatic and mental stress tasks [5,15,24,25,41,49,51,53,69,101,112,117].

Of particular interest to the vagus nerve, PACAP innervation of the lateral central amygdala is thought to arise from PACAP containing neurons in the vagus nerve brainstem complex [93]. A growing literature on the anatomical location and physiology of PACAP suggested a close association with systems that are also regulated by VNS [66,86,88]. Therefore, PACAP is well-positioned as a mediator in pathways of the cholinergic anti-inflammatory axis and its potential modulation by VNS. PACAP may potentially serve as a dynamic and objective biochemical biomarker that could measure PTSD severity, hence the longitudinal changes in PACAP may indicate therapy response to potential treatments targeted at PTSD. The relationship between PACAP levels, stress stimuli, and VNS (either direct or noninvasive stimulation) has not been examined in humans. We previously investigated the physiological [42, 48–50,52], inflammatory [8], and neural [115] effects of tcVNS as a part of a larger study that included three days of chronic stress applied to traumatized subjects with or without PTSD, who were randomized to either tcVNS or sham stimulation (control). In the current pilot investigation, we evaluated the effect of tcVNS treatment by comparing PACAP concentration across timepoints spanning the three days of exposure to personalized traumatic scripts and mental stress (public speaking, mental arithmetic) tasks.

2. Methods

2.1. Overview

The study was reviewed and approved by the Emory Institutional Review Board (#IRB00091171), Georgia Institute of Technology (#H17126), SPAWAR Systems Pacific, and the Department of Navy Human Research Protection Program. Data collection took place in Emory University School of Medicine between May 2017 and October 2019. This study used blood draw data from the project, Closed Loop Vagal Stimulation in Patients with Posttraumatic Stress Disorder funded by N6600116-2-4054 (phases 1 and 2), which aimed to investigate the effects of tcVNS paired with acute stress in the context of traumatic recall and mental stress for PTSD and non-PTSD traumatized controls (ClinicalTrials.gov Identifier # NCT02992899). This project was a parallel study, in which tcVNS or sham stimulation were randomly conducted in both patients with PTSD and healthy subjects. The study included dynamic monitoring of physiological signals, high resolution position emission tomography (HR-PET) of the brain [48,49,51], and blood biochemical collection, PACAP investigation was considered exploratory due to the relevance of PACAP in PTSD, hence it was not a part of the primary or secondary outcomes. A limited number of blood draws (baseline and end of each day) were analyzed for this study, as a pilot, phase 0 investigation.

2.2. Eligibility criteria

a. Non-PTSD traumatized subjects: Subjects aged 18-70 with a history of exposure to psychological trauma, but did not meet criteria for PTSD as determined by the Structured Clinical Interview for DSM-5 (SCID) [35] and Clinician Administered PTSD Scale (CAPS) interview [6].

b. Patients with PTSD: Subjects aged 18-70 met criteria for PTSD according to CAPS.

2.3. Exclusion criteria

Exclusion criteria were: 1) positive pregnancy test; 2) meningitis; 3) traumatic brain injury; 4) neurological disorder or organic mental disorder; 5) history of loss of consciousness greater than 1 min; 6) alcohol abuse or substance abuse or dependence based on the SCID within the past 12 months; 7) current or lifetime history of schizophrenia, schizoaffective disorder, or bulimia, based on the SCID; 8) a history of serious medical or neurological illness, such as cardiovascular, gastrointestinal, hepatic, renal, neurologic or other systemic illness; 9) evidence of a major medical or neurological illness on physical examination or as a result of laboratory studies (Complete blood count (CBC), blood urea nitrogen (BUN), creatinine, blood sugar, electrolytes, liver and thyroid function tests, urinalysis, and electrocardiogram (ECG)); 10) active implantable device (i.e. pacemaker); 11) carotid atherosclerosis; and 12) cervical vagotomy. Women were counseled about the risks of pregnancy during the course of the study and pregnancy tests were conducted before the study started for each female subject.

2.4. Baseline assessments

Each subject informed, provided written consent, after which they underwent a psychological and health assessment. Sociodemographic factors (age, sex, race/ethnicity, marital status, education level) and clinical information (current medications, medical history, SCID) were collected. Table 1 summarizes demographic characteristics and psychological assessment scores grouped by PTSD status and treatment. Additional psychological assessment was performed using structured interviews and standard questionnaires, including PTSD Checklist.
The subjects who did not self-identify as either White or African American identified as multiracial, Asian, or chose not to self-identify race, and missing education data were coded as “other” in analysis. Psychometric scores were reported per the literature recommendation of each test.

2.5. Blinding and stimulation administration

The subjects were randomized into tcVNS and sham groups with a double-blind approach. The research staff and patients were blinded to the stimulus type. The “tcVNS” group was administered stimulation using the electroCore® GammaCore-S non-invasive VNS devices. The tcVNS stimulus intensity was adjusted by the researcher to the maximum tolerable level without causing pain with a burst frequency of 5 kHz, and envelope frequency of 25 Hz (five sine waves at 5 kHz for 1 ms, repeating every 40 ms, amplitude 0–30V peak, adjustable roll switch 0-5 a.u.). The sham devices were identical to the active tcVNS devices in appearance and operation, with different waveform characteristics (0.2Hz biphasic square-like waves, amplitude 0-14V peak, adjustable roll switch 0-5 a.u.), resulting in a tingling sensation. Supplementary Fig. 1 shows both active and sham stimulation waveforms.

For ensuring sham stimulation will not be as powerful as active, we used a low frequency sham waveform to minimize (possible) stimulation with sham device. From an electrical engineering standpoint, high frequency voltage signals (such as the active stimulus, 25Hz with 5 kHz bursts, waveform seen in Supplementary Fig. 1) pass through the skin with minimal power dissipation due to the low skin-electrode impedance (in the range of ohms) at kHz frequencies; in contrast, lower frequency signals (such as the sham stimulus, 0.2Hz, waveform seen in Supplementary Fig. 1) are mainly attenuated at the skin-electrode interface due to the high impedance (in the range of megaohms) [102]. Accordingly, the active device operating at higher frequencies may deliver substantial energy to facilitate nerve stimulation, while the voltage levels appearing under the skin would be expected to be orders of magnitude lower for the sham device and thus vagus nerve stimulation is unlikely with sham.

Nevertheless, since the sham device does deliver relatively high voltage under the skin it would be expected to be orders of magnitude lower for the sham device and thus vagus nerve stimulation is unlikely with sham. Nevertheless, since the sham device does deliver relatively high voltage and current levels directly to the skin, it activates skin nociceptors, causing a similar feeling to a pinch. This sensation is necessary for blinding of the participants.

The duration of delivery was 2 min for each device and every subject received a dedicated device throughout the protocol, which was determined upon randomization before recruitment by a research coordinator who did not take part in data collection or analysis. Collar electrodes were used for ease of delivery, on left carotid pulsation which was used for ease of delivery, on left carotid pulsation which was used for voltage measurement. Nevertheless, the sham device does deliver relatively high voltage and current levels directly to the skin, it activates skin nociceptors, causing a similar feeling to a pinch. This sensation is necessary for blinding of the participants.
2.6. Study design

Protocol consisted of three consecutive days with continuous physiologic monitoring, intermittent blood draws, and one day of HR-PET imaging (Fig. 1). Day 1 focused on personalized traumatic stress and HR-PET imaging, days two and three focused on mentally stressful, non-personalized tasks without HR-PET imaging. Physiologic monitoring and blood draws were carried out on each day. Prior to the protocol, subjects were asked to prepare scripts of their personal traumatic experiences in written form. The scripts were transcribed, edited to last 60 s each, and recorded by a research associate in normal voice, in the first person, present tense. On the first day, after the initial set up with physiologic monitors and intravenous catheter (IV), subjects rested for 30 min and baseline measurements were collected. Laying down in HR-PET scanner, subjects underwent 14 HR-PET scans with contents detailed in Fig. 1 in order. Six of these scans included pleasant scenery recordings (termed as “neutral script”, ~60 s, scan # 1, 2, 7, 8, 11, 12), other six included listening to the personalized traumatic stress recordings (~60 s, scan # 3, 4, 9, 10, 13, 14), each traumatic stress recording was immediately followed by stimulation (active or sham, 120 s). Lastly, the remaining two scans included only stimulation (active or sham, scan # 5, 6) without any playback or acute stress. The subjects were instructed to image each event as vividly as possible. Blood was drawn (total 194.5 cc) at baseline (referred as timepoint 0 in this manuscript), and during timepoints 4, 6, 8, 10, 12, 14 as expressed in Fig. 1 (immediately after the aforementioned event ended, such as the end of recordings or stimulation). The second and third days were identical to each other. After initial set up and resting, baseline (timepoints 15 and 19 for days 2 and 3, respectively) blood draws were taken both mornings. Afterwards, subjects underwent a public speech task and mental arithmetic tasks, as described previously [49]. Stimulations were applied immediately after each task. After two mental stressors and two stimulation administrations, the subjects were given a 90-min break. After the break, a second blood draw was taken (timepoints 18 and 22 for days 2 and 3, respectively). A total 57 cc of blood was drawn per day on these days. It should be noted that a third stimulation was applied after the blood draw of 90-min break to monitor the physiologic changes to stimulation without acute stress, however this administration was after the blood draws (after timepoints 18 and 22). Since the PACAP investigation was exploratory, PACAP analysis used baseline (day 1, timepoint 0), end of day 1 (timepoint 14), end of day 2 (timepoint 18), and end of day 3 (timepoint 22) due to budget constraints.

2.7. Biomarker measurement

Human PACAP was assayed with a competitive ELISA using kits purchased from LS Bio (Seattle, Washington) for blood draws at timepoints 0, 14, 18, 22. Fig. 2 depicts the analyzed PACAP timepoints with number of subjects and PTSD status for each timepoint. All experimental operations were in accordance with standard protocols. Intra-assay coefficient of variation (CV) was 9.89%, sensitivity was 2.55 pg/ml, and the detection range was 6.17–500 pg/ml. The polyclonal antibody used in this assay does not differentiate between the PACAP-27 and PACAP-38 isoforms. However, since the PACAP-38 is the most abundant isoform in mammals [23], it is likely that most of the detected peptide is PACAP-38.

2.8. Statistical methods

Since the distribution of PACAP concentration was skewed, log-transformation was applied to PACAP concentrations to achieve normality and better interpretation in all the subsequent analyses. These
PACAP data were confirmed to be normal using Shapiro-Wilk test. Initially, we computed partial correlations between (log) PACAP concentration (baseline, maximum, quartiles (lower (Q1, 25th percentile), medium (Q2, 50th percentile), upper (Q3, 75th percentile)) for each subject) and psychometric scales, controlling for age, sex, BMI, race, and education and reported correlation coefficients (rho, $\rho$) and p-values. To assess the trajectory of PACAP concentration over time, we computed the delta (from baseline) at each timepoint. We used a linear mixed model to regress the delta PACAP values on timepoints (0, 14, 18, 22), PTSD status (yes/no), device type (active tcVNS/sham), as well as other covariates (BMI) with subject-specific random intercepts to account for within-subject correlations. When a significant main effect or interaction was present, pairwise post-hoc comparisons with Bonferroni-Holm corrections were used to determine location of differences. For this analysis, an a priori planned comparison approach yielded the number of tests as five: successive time points (e.g., 14 vs 18) and all numbers different from 0. Effect sizes (Cohen’s $d$) were reported for results, regardless of the significance of p-value, based on group means, standard deviation, and group sample size for device and PTSD status main effects [68]. For post-hoc analyses that include timepoints, repeated measures Cohen’s $d$ (d_{rm}) was calculated that includes a correction factor based on the correlation of PACAP concentrations between two timepoints (for example, correlation of delta PACAP between timepoints 0 and 18 and between timepoints 0 and 22 were included as a correction factor) [68]. Missing biomarker data were assumed to be missing completely at random for reasons such as hemolyzed sample or missing baseline data, and all analyses were based on available data. P-values less than or equal to 0.05 were considered statistically significant. Statistical analyses were done using R (v 3.6.0) and MATLAB R2020a.

3. Results

A total of 36 subjects who completed or at least partly completed the trauma tasks were involved in these analyses. Fig. 2 depicts the CONSORT diagram of the study, Table 2 breaks down the PACAP concentrations per PTSD status and treatment groups, and Supplementary Fig. 2 shows normalized PACAP concentration separated by device and PTSD status. Subject groups were similar in age, body mass index, race, education level and marital status (Table 1). Although subjects were randomly assigned to the tcVNS treatment or sham treatment groups, only female PTSD patients received tcVNS treatment. The gender proportion in the other groups was similar. The average age of this population was 30.3 (SD = 7.9) years, and the average BMI was 27.0 (SD = 5.5) kg/m$^2$. Among all the 30 subjects, 15 (50.0%) were female, 13 (43.3%) White/Caucasian, 17 (56.7%) had a Bachelor’s or higher degree, and 20 (66.7%) had never married.

First, the association between PACAP baseline concentration and scores of psychological and functional scales were examined. Table 3 lists

![Fig. 2. Consolidated Standards of Reporting Trials (CONSORT) diagram of the study.](image-url)
Partial correlation coefficients and p-values between each psychological scale and related PACAP concentrations. Baseline PACAP concentrations were significantly positively correlated with total PTSD Symptom Score (PTSS-SS) \((p = 0.45, p = 0.04)\) and significantly negatively correlated with Baecke Sports Index (\(p = -0.46, p = 0.05\)). The maximum PACAP concentrations (from timepoints 0 to 22 for each subject) were significantly positively correlated with Hamilton Anxiety Scale (HAM-A) \((p = 0.43, p = 0.05)\) and Hamilton Depression Scale (HAM-D) \((p = 0.45, p = 0.04)\). Upper quartile PACAP concentrations (from timepoints 0 to 22 for each subject) were significantly negatively correlated with Baecke Leisure Index \((p = -0.53, p = 0.03)\). None of the other psychological and functional scales were statistically significant with the examined PACAP concentrations.

When examining the PACAP values within the overall sample, the main effect of device type \((p = 0.26, d = 0.46)\) and PTSD status \((p = 0.13, d = 0.12)\) along with interactions of device type by time \((p = 0.53)\), device type by time by PTSD status \((p = 0.59)\), PTSD status by time \((p = 0.48)\), and PTSD status by device type \((p = 0.63)\) were not significant. However, the main effect of time was statistically significant \((p = 0.008)\). Post-hoc comparisons revealed elevated PACAP at timepoints 18 \((p = 0.03, d_{\text{em}} = 0.27)\) and 22 \((p = 0.03, d_{\text{em}} = 0.28)\), but not at timepoint 14 \((p = 0.27, d_{\text{em}} = 0.18)\). No other significant differences were observed at any timepoint for the overall sample \((p > 0.05)\).

### Table 3: Partial correlations (\(\rho\), p-value) controlling for age, sex, BMI, race, education, \(p\) partial correlation coefficient, \(^* p \leq 0.05\)

| Psychological Assessment Total Score | PACAP data | \(\rho\) | \(p\) | \(p\) value |
|--------------------------------------|------------|--------|------|-----------|
| PTSD Symptom Scale (PTSD-SS)         | Baseline   | 0.45   | 0.04*|           |
| PTSD Checklist (PCL-C)               | Baseline   | 0.21   | 0.35 |           |
| Clinician-Administered PTSD Scale (CAPS) | Baseline   | 0.72   | 0.49 |           |
| Early Trauma Inventory (ETI)         | Baseline   | 0.22   | 0.49 |           |
| Adulthood Trauma Inventory (ATI)     | Baseline   | 0.24   | 0.29 |           |
| Hamilton Anxiety Scale (HAM-A)       | Maximum    | 0.43   | 0.05*|           |
| Hamilton Depression Scale (HAM-D)   | Maximum    | 0.45   | 0.04*|           |
| Beck Depression Inventory (BDI)      | Baseline   | 0.28   | 0.21 |           |
| ESSI (Social Support Inventory)      | Baseline   | 0.28   | 0.21 |           |
| Baecke Questionnaire (Sports)        | Baseline   | -0.46  | 0.05*|           |
| Baecke Questionnaire (Work)          | Baseline   | -0.15  | 0.54 |           |
| Baecke Questionnaire (Leisure)       | Upper Quartile (Q3) | -0.53  | 0.03*|           |

Although the device type by time interaction was not significant, a subsequent analysis was completed using an isolated dataset with just sham or active across time to better determine the contributions of treatment group to the overall time effect (Fig. 3). In our stratified analysis, a significant main effect of time was observed in the sham group \((p = 0.04)\). Post hoc tests revealed significant differences at timepoints 18 \((p = 0.04, d_{\text{em}} = 0.35)\) and 22 \((p = 0.04, d_{\text{em}} = 0.41)\) compared to timepoint 0. No other significant differences were observed in sham including timepoint 14 \((p = 0.13, d_{\text{em}} = 0.27)\).

In contrast, no significant main effects or interactions were observed for active tcVNS \((p = 0.21)\). The tcVNS active group effect sizes were \(d_{\text{em}} = 0.12, d_{\text{em}} = 0.21,\) and \(d_{\text{em}} = 0.18,\) for days 1, 2, and 3, respectively.

### 4. Discussion

In this randomized, double blind, pilot study of trauma recall and other psychological stressors and longitudinal assessment of plasma PACAP levels, we found that PACAP increased over the course of the stress protocol—an effect attenuated by tcVNS (but not sham stimulation) in traumatized individuals. Increased PACAP concentrations were correlated with elevated PTSD symptoms at baseline, replicating earlier findings \([100]\). We also found that elevated PACAP was associated with increased symptoms of anxiety and depression and impairments in social and physical function. As PACAP is known to regulate stress response \([30,100]\), longitudinal evaluation of PACAP may be helpful in tcVNS treatment monitoring.

This investigation touches on two main points. First, to our knowledge, this is the first report of PACAP in humans undergoing a trauma recall and mental stress paradigm over multiple days. Trauma recall and stressful tasks were associated with a steady increase in PACAP blood levels, regardless of the treatment status. Second, notably, the sham group’s PACAP increase was higher, compared to active tcVNS group, which suggests that tcVNS may reduce PACAP elevation in response to stress. These results, along with correlations between PACAP and psychological scales (PTSS-SS, HAM-A, HAM-D, Baecke Sports Index, Baecke Leisure Index) suggest that PACAP may play an integral role in stress and PTSD, supporting relevant literature \([30,100]\).

PTSD is associated with poor health behaviors, notably physical inactivity \([119]\), also recognized in our study with negative correlations of baseline PACAP with Baecke Sports Index. We did not find significant correlations with other assessments of PTSD symptoms (CAPS, PCL-C, ETI, ATI), we believe that this could be due to small sample size. For example, only patients with PTSD can have a total CAPS score, which significantly decreased the sample size for this correlation. Regardless of the p-values, all psychological surveys indicating increased severity with higher scores were positively correlated with baseline PACAP. Similarly, physical activity scales (Baecke Questionnaire) were negatively correlated.

The source of PACAP in circulating blood or plasma is not known \([66,67,105]\). PACAP has a close association with systems that are also modulated by VNS. PACAP is involved in a number of processes including limbic, autonomic, neuroendocrine functions, and regulation of circadian pacemaker \([93]\). Another review study noted that PACAP is an important regulator of hippocampal circuits \([58]\). Our HR-PET results from the current study, and our observations during sham stimulation compared to active tcVNS within the limbic and other brain areas involved in stress, including bilateral prefrontal and orbitofrontal cortex, premotor cortex, temporal lobe, parahippocampal gyrus, insula, and left anterior cingulate \([115]\). In addition, we investigated physiologic measures that reflect autonomic nervous system activity for the same study, and our findings suggest that active tcVNS attenuates autonomic reactivity to stress, as observed in data obtained during stimulation or within minutes after stimulation \([42,48,49,52]\). The attenuation in limbic and autonomic activity could potentially have
led to less induction of PACAP in the tcVNS group, though it would largely be a speculation given the unclear source of PACAP in circulating blood. If we consider the measurement timescales, our PACAP results have a delay as we processed the blood samples at baseline (day 1 morning) and at the end of each day (days 1, 2, 3). As tcVNS attenuated both limbic and autonomic activity based on our previous investigations within seconds (physiological activity) and minutes (HR-PET scan resolution), one might think that the attenuation in these activities decreased PACAP as PACAP due to dampening stress responses. Nevertheless, understanding the source of PACAP and the details of a sequential relationship is beyond the scope of this paper.

The use of traumatic stress scripts or imagery is a standard practice in trauma and PTSD research [9,12,14,16,70,72,85,97]. As a part of the brain imaging investigation of this protocol, we previously reported the effects of traumatic stress scripts on brain’s fear circuitry, regardless of the device status (active and sham collapsed together). Our results indicated that listening to the personalized trauma scripts elicited similar brain activation patterns to what was previously reported in healthy traumatized individuals and patients with PTSD.

We paid substantial attention for the selection of the sham device. Our study does not use an inert sham device (i.e., no power delivery as sham). Use of inert devices as controls could lead to participants’ perception that they were not getting active interventions, increasing the risk of positive results that are only due to a placebo effect, a constant risk in device research. Furthermore, the use of “active” sham stimulation controls has been used in other studies to date, including for transcutaneous auricular VNS, so our study is consistent with the literature and standard practice in use of sham devices [71,116].

In our study, stimulations occurred after the traumatic or mental stressors, not during or before. The discussion surrounding timing (i.e., when to stimulate when paired with stress) is unclear given the clinical application and current state of knowledge surrounding tcVNS. Realistically, patients undergoing traumatic flashbacks can only use tcVNS during or following a traumatic memory, as they cannot predict the onset. While previous studies indicate lasting effects of stressful protocols, and thus presenting theoretical basis for stimulation after the traumatic memory cessation, it is unclear whether the effects are similar when compared to stimulating during traumatic stress. We applied stimulation after the trauma recall because we wanted to assess whether the groups had similar traumatic reactivity when the stimuli was first initiated, as detailed in our physiological outcomes work [49]. Unlike physiological signals, PACAP measurement does not have a second-based time resolution, and is likely capturing PACAP levels due to cumulative effect of these stressors. Previous studies in animals (direct VNS) applied stimulation both during — which also may be applicable in humans — and before traumatic stress paradigms [33,89]. No current study has examined the efficacy and/or changes in any sort of outcome when stimulation is applied either during or following traumatic stress in humans, and therefore it is unclear how the timing affects the findings in the current study. Given that the effects of traumatic stress, as autonomic parameters, persists beyond the initial presentation of the adverse stimulus, it is possible that stimulation during or after a traumatic script elicits similar changes in arousal. This is consistent with findings across various studies reporting autonomic dampening with implanted or noninvasive VNS, despite variations within stimulation timing and protocols [15,19,20,69,83,90,91].

We believe that specific biomarkers such as PACAP or other objective measures (blood biomarkers, brain imaging, physiological biomarkers) have the potential to be more reliable than self-reported scales in clinical investigations since they include objective biological indices. While the PACAP investigation did not have self-reported PTSD symptom outcomes matching the PACAP timepoints, we recently published behavioral outcomes that suggest that patients with PTSD experience lower perceived anger with active tcVNS, compared to sham [8]. Perceived anger was measured by visual analog scores after each traumatic stress paired with stimulation on day 1, as a part of our main investigation. Recent neuroscience studies using implanted VNS in rat models trained with fear conditioning reported enhancement of fear extinction based on freezing time [89,91]. Along similar lines, fear conditioning studies on human subjects using transcutaneous auricular VNS reported acceleration in fear extinction [20], and enhanced processing of safety cues based on US expectancy ratings [18]. Objective measures will gain value in complementing. PTSD diagnosis based on clinician administered PTSD instruments. These instruments such as CAPS, PCL-C, or ATI have well-studied repeatability [6,103,113,114], however they could only be administered between long time scales (i.e., months), hence they are in the current practice most useful long term longitudinal clinical studies. Dynamic biomarkers such as PACAP that could quantify treatment effects in shorter time scales are needed for psychiatry research.

The limitations of this pilot study could be listed as follows. Our sample included 12 patients with PTSD, and all three male patients were randomly assigned to the sham group. Due to this small sample size and the fact that none of the PTSD male subjects received the tcVNS treatment, we cannot evaluate the effect of tcVNS treatment on PACAP concentrations among PTSD males. Moreover, as gender may be an effect modifier of PACAP concentrations and PTSD, this interaction could not be evaluated within the tcVNS treatment group. Prior work has suggested that blood levels of PACAP are associated with PTSD diagnosis among females and stress-regulation pathways may vary between men and women [100]. Although levels of PACAP were lower at all times points taken after traumatic stress and mental stress in the tcVNS group compared to the sham group, the interaction term for device by time was not significant. This is also likely related to the small sample size, and therefore our results should be considered preliminary and in need of replication. PACAP is involved in circadian rhythm regulation [21,40,61]. Although our blood draws were scheduled in the A.M., our study lasted for three days, and the precise blood draw time for each subject varied somewhat, as would be anticipated with a multi-day human investigation. In partial correlations between PACAP data (baseline, maximum, quartiles) and psychometric scales, we did not control for multiple testing, which may result in a Type I error. These results should be interrupted very cautiously since correction for multiple comparisons would yield all of these findings statistically non-significant. Again, this is considered an exploratory analysis, and results need to be established in a larger study.

Due to the clinical nature of this study, we did not evaluate whether vagal stimulation occurred, rather, we relied on previous literature that extensively studied to what extent vagal stimulation was achievable noninvasively. There are multiple studies that reported the ability to reach the vagal afferents using tcVNS using the same device [39,80]. We replicated the stimulation application reported in Ref. [39] throughout the protocol, by locating the carotid artery as reference. In addition, a study using the studied device was reported to reliably create vagal somatosensory evoked potentials, which — are also activated with vagal implant, associated with vagal afferent activation [84]. While these studies favor our methods, we also recognize a recent pig model study that suggested noninvasive vagal nerve stimulation (A6- and B-fiber activation) is largely not achievable with clinically tolerable current levels defined by the researchers [81], though not necessarily proving that noninvasive stimulation does not activate human vagal fibers with any certainty.

In this study, we showed that acute traumatic and mental stressors are associated with increased PACAP concentrations in the peripheral blood of traumatized individuals both with and without PTSD. PACAP appears to be a modifiable biochemical biomarker, and its temporal changes may predict tcVNS treatment effect to acute stress or neuropsychiatric disorders showing significant PACAP dysregulation. Moreover, longitudinal monitoring of PACAP may potentially be used to follow personalized, adaptive dosing strategies for larger trials, or to identify responsive and non-respondent patients to potential treatments. Future studies should investigate sex differences in PACAP concentrations with acute and longitudinal tcVNS treatment using larger sample sizes.
Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.cjence.2020.10012.

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