Skin conductance response to trauma interview as a candidate biomarker of trauma and related psychopathology in youth resettled as refugees

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ABSTRACT

Objective: Previous research in adults found that skin conductance responses to trauma interview predicted current and future PTSS. We extended this method to refugee youth exposed to civilian war trauma and forced migration, to examine associations between PTSS and skin conductance in this uniquely vulnerable child and adolescent population.

Methods: 86 refugee youth ages 7–17 years completed a trauma interview and assessment of self-reported PTSS. The mobile eSense app on an iPad was used to obtain continuous recordings of skin conductance level (SCL) during a trauma interview (trauma SCL). Skin conductance response (SCR) was calculated by subtracting the baseline SCL from the maximum amplitude of the trauma SCL.

Results: SCL during trauma was significantly greater than baseline SCL. Trauma exposure was significantly associated with SCR to trauma interview, $R^2 = 0.084$, $p = 0.042$. SCR to trauma interview was positively correlated with reexperiencing ($R^2 = 0.127$, $p = 0.028$) and hyperarousal symptoms ($R^2 = 0.123$, $p = 0.048$).

Conclusions: The present study provides evidence for feasibility of SCR to trauma interview as a candidate biomarker of PTSS in youth. This is the first study to look at SCR to trauma interview in youth resettled as refugees and is part of the limited but growing body of research to look at biomarkers in refugee cohorts more broadly. As the number of forcibly displaced persons surges, early detection and prevention of trauma-related psychology is becoming more important than ever.

KEYWORDS

Trauma; PTSD; skin conductance response; refugee health; child and adolescent

HIGHLIGHTS

• Using the mobile eSense app, we demonstrate that skin conductance is measurable in a variety of research settings and that skin conductance response may be a biological indicator of trauma and related psychopathology – namely re-experiencing symptoms – in youth resettled as refugees.

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1. Introduction

The Diagnostic and Statistical Manual of Mental Disorders (DSM-5) defines trauma as actual or threatened death, serious illness/injury, or sexual violence (American Psychiatric Association, 2013). Trauma can be experienced directly or as part of one's occupation, witnessed, or learned about. Trauma exposure confers risk for developing posttraumatic stress disorder (PTSD), which is defined by intrusive/reexperiencing, avoidance, negative cognition and mood, and arousal/reactivity (hyperarousal) symptoms. Traumatic experiences are common in childhood, in 1 in 5 youth experience trauma (Saunders & Adams, 2014). However, not everyone exposed to trauma develops PTSD – of the 20% of youth exposed, only about 5% develop PTSD (Merikangas et al., 2010) although this rate may be higher in higher-risk populations, including youth who resettle as refugees (rates of PTSD range from 9% to 54%) (Grasser et al., 2021; Perreira & Ornelas, 2013; Solberg et al., 2020). Furthermore, there is heterogeneity in symptom presentation (Galatzer-Levy & Bryant, 2013). Additionally, children with subthreshold symptoms of PTSD may still be significantly distressed and experience functional impairment (Carrion, Weems, Ray, & Reiss, 2002).

The heterogeneity of PTSD and the continuum across which persons may experience posttraumatic stress symptoms (PTSS) has motivated the query for possible biomarkers related to posttraumatic stress. Biomarkers are measurable characteristics reflecting biological function or dysfunction, therapeutic response, and/or natural disease progression (Group et al., 2001). The goal of biomarkers research is to define objective signs and states that can be observed and measured accurately and reproducibly (Michopoulos, Norrholm, & Jovanovic, 2015). Biomarkers may reflect genetic or physiological characteristics that may put an individual at risk for a disease state when confronted with certain environmental exposures, or may reflect ways in which certain exposures have affected physiology and may contribute to disease (Mayeux, 2004; McKeever & Huff, 2003). Currently, there are no validated and reliable biomarkers for any psychiatric condition (Casey et al., 2013; Macaluso & Preskorn, 2012), and
diagnoses are made based solely on clinical interviews (Casey et al., 2013; Kupfer & Regier, 2011). However, with growing widespread research in this field (Kalia & Costa e Silva, 2015), biomarkers may be leveraged to predict which individuals are at greater risk for developing PTSD in the aftermath of trauma and provide an objective method for identifying those with the disorder as well as measuring treatment efficacy (Macaluso & Preskorn, 2012). There are over 600,000 ‘ways’ to have PTSD given the 20 DSM-defined symptoms across 4 domains and an additional dissociative subtype (Galatzer-Levy & Bryant, 2013). Biological diatheses (McKeever & Huff, 2003) and the differential ways in which trauma can impact the nervous system (Lanius, Bluhm, Lanius, & Pain, 2006) may contribute to such heterogeneity in posttraumatic stress. Indeed, a recent neuroimaging study found heterogeneous stress responses in the immediate aftermath of trauma exposure (Stevens et al., 2021). These brain-based biotypes support the idea that there may be biological phenotypes related to risk, resilience, and differential presentation of posttraumatic stress (Stevens et al., 2021). This heterogeneity in response to trauma exposure and disease expression may require fine-tuned intervention, and biomarkers of specific symptoms could define specific therapeutic targets.

When an individual is exposed to a stressful, dangerous, and/or traumatic event as well as reminders of such events, neurotransmitter and hormonal signalling cascades trigger relevant physiological and behavioural responses, including modulation of sweat secretion (Shibasaki & Crandall, 2010). A component of such sympathetic responses is increased perspiration (Seligowski, Harnett, Merker, & Ressler, 2020; Van der Kolk, 2015), and skin conductance is a biological indicator of arousal/reactivity (Christopoulos, Uy, & Yap, 2019). Amygdala signalling to the hypothalamus in response to salient stimuli or trauma-related cues triggers a cascade of defensive responses, including thermoregulatory responses mediated by the preoptic region of the hypothalamus (Shibasaki & Crandall, 2010). Direct amygdala to brainstem signalling also occurs (Shibasaki & Crandall, 2010). Resultant downstream cholinergic and adrenergic signalling at the eccrine glands stimulates sweat secretion. Sweat gland activity increases with the stress response and serves as a measure of sympathetic activation (Harker, 2013). Increased moisture due to sweat secretion can be detected as a change in conductance and electrical potentials using electrodes. Prior studies have reported alterations in autonomic function, including decreased resting parasympathetic nervous system (PNS) activity and heightened reactivity of the sympathetic nervous system (SNS) during mental stress in PTSD (Buckley & Kaloupek, 2001; Park et al., 2017; Pole, 2007), including elevated sweating, or skin conductance. Here, elevated skin conductance as a marker of increased autonomic reactivity, or arousal, may reflect a predisposing physiological phenotype that increases risk for posttraumatic stress symptoms when confronted with trauma. Alternatively, elevated skin conductance as a marker of autonomic reactivity may reflect effects of trauma exposure – particularly chronic and severe trauma exposure, as in civilian war trauma – on the nervous system. Here, persistent activation of the SNS as described above may contribute to persistent increased skin conductance and confer greater reactivity to novel stressors or reminders thereof following exposure.

Elevated skin conductance response (SCR) to stimulation is a sensitive measure of hyperarousal and predictive of PTSD when using script-driven imagery (Pineles et al., 2013), a method in which individuals listen to trauma-related scripts. Trauma-related scripts may be individualized to one’s own experiences; standardized traumatic scripts may also be used for comparison, in addition to personalized and standardized positive and neutral scripts (Hoge et al., 2012; Pitman, Orr, Forgue, de Jong, & Claiborn, 1987). These protocols have indicated that physiological arousal was specifically associated with individualized trauma-related scripts compared to the standardized trauma scripts in individuals with PTSD (Orr, Metzger, & Pitman, 2002; Pitman et al., 1987). Additionally, neutral and positive scripts have not been shown to generate physiological responses in neither individuals with nor without PTSD (Orr et al., 2002; Pitman et al., 1987). Resultantly, elevated SCR to trauma reminders could serve as a candidate biomarker of PTSD and its specific symptoms. Recent studies have built on these methods using the mobile eSense application, to improve accessibility and cost-effectiveness for clinical and research settings. Using this technology, previous research found that SCRs elicited by a standardized trauma interview – in which participants described a traumatic event during the eSense recording – were positively associated with severity of posttraumatic stress symptoms and significantly predicted presence of the disorder 6 months later (Hinrichs et al., 2017; Hinrichs et al., 2019). Increased SCR to trauma interview is emerging as a candidate biomarker of PTSD and its specific symptoms. Mobile collection methods further the utility of this biomarker, especially in settings where clinicians may be unavailable for full diagnostic interviews and mental health screenings, in the immediate aftermath of trauma, and in challenging and dynamic research contexts. One such example is with persons who are resettling as refugees in host countries who may have encountered various stressors such as war, natural disaster, and forced migration, where data must be collected in homes rather than laboratories.
Ongoing conflicts in Syria and Iraq have exposed millions of youth to trauma and caused many to be displaced. Such youth have experienced potential separation from friends and family members, death of loved ones and strangers alike, loss of belongings, lack of basic living needs, and additional stressors in resettlement camps. Following resettlement, youth may in turn experience new stressors – e.g. being considered a minority group in a host country, discrimination and bullying, language barriers, and sociocultural transitions. As a result, youth who resettle as refugees experience higher rates of PTSD compared to the general United States population (Grasser et al., 2021; Javanbakht, Rosenberg, Haddad, & Arfken, 2018). This underscores a need for objective assessment of trauma-related psychopathology and pathophysiology in youth who have resettled as refugees. Such objective assessments could be feasible for use in not only research labs, but also in real-world settings where they may have the greatest utility and clinical impact (Loś & Waszkiewicz, 2021) – including resettlement agency offices, medical clinics, community centres, and even patient homes with appropriate consultation and permissions (Hiba Abu Suhaiban, Grasser, & Javanbakht, 2019). Therefore, the present study aimed to investigate the feasibility of collecting SCR during a trauma interview using the mobile eSense system in a cohort of youth ages 7–17 who resettled in the US as refugees of Syria and Iraq. SCR has previously been used to monitor threat and safety learning in infants (Ingram & Fitzgerald, 1974), healthy non-trauma exposed children and adolescents (Michalska et al., 2016; Miller & Shields, 1980; Morrow, Boring, Keough, & Haesly, 1969; Neumann, Waters, & Westbury, 2008), and youth exposed to maltreatment (physical, and sexual abuse, as well as domestic violence) (McLaughlin et al., 2016). However, this is one of the first studies to use SCR in tandem with a trauma interview and as a biomarker for PTSS in a child and adolescent sample (Wiltshire et al., 2022). We also examined SCR as part of a Research Domain Criteria (RDoC) approach to investigate pathophysiology of mental illness, specifically the negative valence domain construct of arousal, using psychophysiological and self-report data as units of analysis (Cuthbert & Insel, 2013; Cuthbert & Kozak, 2013; Morris & Cuthbert, 2012). We hypothesized that severity of PTSS would be positively correlated with SCR to trauma interview. In addition, we examined association with the specific symptom clusters of re-experiencing and hyperarousal, as those have been previously linked with physiological responses in PTSD (Jovanovic et al., 2010; Norrholm et al., 2015, 2011). Our present study leverages data from a largely under-represented group in mental health – youth of Middle Eastern/North African (MENA) ethnicity – and provides recommendations for future application of this method.

2. Methods

2.1. Participants

86 youth (36 female, mean age = 12.419, +/- 2.737 SD, range = 7–17) were recruited from an existing cohort of persons resettled as refugees of Syria and Iraq in Southeast Michigan (Grasser et al., 2021; Javanbakht et al., 2018). Participants were initially recruited as families at the Arab American and Chaldean Council Clinic, where all refugees resettling from Middle Eastern nations, received mandatory health screenings within one month of migration to the U. S. Inclusion criteria were as follows: (1) males and females between the ages of 7 and 17, (2) refugee status, originating from Syria or Iraq, and (3) willing and able to give oral assent (ages 7–12) or written assent (ages 13–17) with parent/guardian willing and able to give written informed consent. Exclusion criteria were as follows: (1) inability or unwillingness to consent, (2) wardens of the court, and (3) current diagnosis of autism or psychosis spectrum disorders. Siblings were permitted to participate in the study. All study procedures described herein were approved by the Institutional Review Board at Wayne State University (IRB#012416B3F).

2.2. Self-report measures

All self-report assessment measures were administered by bilingual clinicians and research assistants and were available in both English and Arabic. Questionnaires were translated from English to Arabic by a native speaker, back-translated to English by a separate Arabic speaker to ensure consistency and accuracy in translations, and finally certified and approved by external reviews. Internal consistency was assessed with Cronbach’s alpha and included for each measure below.

Trauma exposures were assessed using a modified version of the Harvard Trauma Questionnaire (HTQ), which queries traumatic events more specific to refugee and non-Western cohorts (Mollica et al., 1992; Rasmussen, Verkuilen, Ho, & Fan, 2015). The modified version of the HTQ included 21 items, to which participants would respond ‘yes’ or ‘no’ as to whether they had experienced the event, regardless of frequency. The item responses were then summed to provide the total number of unique traumatic events youth had been exposed to, with a possible range of 0–21. The sections from the original HTQ on head injury, trauma symptoms, and torture history were not included in this modified version – no participants reported head injury at initial screening, and the sections on torture history were inappropriate for use in children and adolescents. The UCLA PTSD Reaction Index was used as a measure of trauma...
symptoms. The modified version of the measure is available in the Supplementary Materials, with a list of the traumatic events that were excluded in the modification. Cronbach’s alpha for the 21-item HTQ trauma inventory in this sample was 0.887.

The severity of PTSS was assessed using the UCLA PTSD Reaction Index for Children and Adolescents (UCLA PTSD RI) for DSM 5 (Kaplow et al., 2020). The UCLA PTSD RI is a 31-item questionnaire to which participants indicate how much they have been bothered by particular symptoms within the last month, ranging from none of the time (0) to most of the time (4). The measure also generates subscales for re-experiencing, avoidance, negative cognitions, and hyperarousal symptoms. The UCLA PTSD RI has been recommended for use with refugee populations, including those with non-Western backgrounds (Miller, Brown, Shramko, & Svetaz, 2019), and has been found to be reliable in MENA refugee populations (Grasser et al., 2021; Javanbakht et al., 2018). Cronbach’s alpha in this sample for the full measure was 0.933. For re-experiencing symptoms, Cronbach’s alpha was 0.826; for hyperarousal symptoms, alpha was 0.734. Given that these are the symptom clusters most strongly linked to the neurobiological pathways associated with PTSS and related to the biomarker of interest, we made a priori hypotheses about these two symptom clusters and did not do exploratory analyses for the others to avoid Type I error.

2.3. Skin conductance response to trauma interview

Data were collected either in participants’ homes or in a university lab setting, to minimize barriers to research participation. The procedures were administered the same way in both settings. Skin conductance level (SCL) was measured using the eSense app (Mindfield Biosystems, Inc., Berlin, Germany) on an iPad. Two Ag/AgCl electrodes were placed on the index and middle fingers of the non-dominant hand. Isotonic paste was added to electrodes prior to attachment to the middle phalanges of the fingers to increase signal quality and ensure contact with skin. A two-minute baseline was obtained at rest, during which time the participant was seated in a quiet room refraining from talking or engaging with any stimuli. This was followed by a measurement of SCL while the individual was administered the HTQ interview. Throughout the recordings, participants remained seated and still with their hands resting on a table to reduce movement. An example recording from a single participant session is depicted in Figure 1, with an inset image of the data collection software. Data were exported to Excel for initial processing of raw data and scored values were imported to R for statistical analysis. No artifacts were identified in the dataset, and no additional filtering was applied to raw values. As in prior studies, baseline SCL was calculated as the average of the last 30 s of the two-minute baseline recording taken at rest. Trauma SCL was calculated as the average of full trauma interview. SCR was calculated as the baseline SCL subtracted from the peak skin conductance value during the trauma interview. Trauma SCL reflected the continuous level of electrodermal activity throughout the entire interview. SCR reflected the peak response during this trauma interview (maximum amplitude) from which the baseline SCL was subtracted to normalize across participants.

2.4. Data analysis

All analyses were performed in R statistical package version 4.0.3. Data were screened for potential artifacts in skin conductance recordings; we did not identify any artifacts and there were no instances of measurements in which a recording was unable to be obtained. Data were then screened for univariate outliers based on median absolute deviation (Kwak & Kim, 2017; Leys, Ley, Klein, Bernard, & Licata, 2013); outliers for baseline SCL (n = 5), trauma SCL (n = 3) trauma exposure (n = 3), SCR (n = 6), reexperiencing symptoms (n = 8), and hyperarousal symptoms (n = 2) were identified and corrected using Winsorization (Dixon, 1960; Kotz, Read, Balakrishnan, Vidakovic, & Johnson, 2006). Missing data were identified for baseline SCL (4.7% missing), trauma SCL (4.7% missing), SCR (5.8% missing), total PTSD symptoms (17% missing), reexperiencing symptoms (13% missing), and hyperarousal symptoms (9.3% missing). Little’s MCAR indicated that data were missing completely at random, p = .634. Patterns of missing data were also inspected, and no significant patterns of concern emerged. Missing data were imputed using multiple imputation with 8 iterations, given that no variables exceeded 20% missingness and the sample size. Multilevel models were fit to accommodate the nested structure of this dataset. In cases where there were repeated measures (i.e. comparing skin conductance level from baseline to trauma interview), data were nested by individuals, as 2 measurements were obtained – one for baseline and one for trauma SCL – and individuals were nested by family – for 172 total observations across 32 groups. In cases where there were not repeated measures (i.e. assessing relations between variables), data were nested by families only for 86 total observations across 32 groups. Random intercepts, fixed slopes models were selected, as we did not have any reason to hypothesize that patterns would differ by family. This was confirmed by ANOVAs and model fit indices (AIC, BIC) indicating that
random intercepts, fixed slopes models were of better fit than random intercepts, random slopes. Sex and location of assessment were included as covariates in the models. Models were screened to ensure that the assumptions of the model were met – namely the assumption of normality of the residuals. Adjusted alpha for four models \((p < .0125)\) was based on the Bonferroni correction for multiple comparisons.

3. Results

3.1. Demographic Characteristics

Data were collected from 86 youth (36 female) from 32 families with an average of 2.69 (range: 1-6) participating youth per family. Most of the participants had at least 1 sibling in the study; only 4 participants did not have any siblings in the study sample. 31 individuals were assessed in their homes and 55 at the lab. The number of unique traumatic exposures per individual ranged from 0 to 15. Participants were between the ages of 0 and 9 (\(M_{age} = 3.43, SD\pm2.57\)) at the time of trauma exposure; ages 7–17 (\(M_{age} = 12.42, SD\pm2.74\)) at the time of assessment. Descriptive statistics are shown in Table 1.

Age was significantly associated with number of traumatic events endorsed on the HTQ, \(R^2 = .28, p < .001\). Older age was associated with more trauma exposure; however, age was not associated with severity of post-traumatic stress symptoms, all \(p\) values > .10. The number of traumatic events endorsed on the HTQ was positively correlated with posttraumatic stress symptoms, \(R^2 = .27, p < .001\). Age was not associated with baseline SCL \((p = .58)\) or SCR to trauma interview \((p = .29)\), and neither baseline SCL nor SCR differed by gender \((t = 1.09, p = .28\) and \(t = -1.24, p = .22\) respectively). Neither age \((t = .64, p = .52)\) nor trauma exposure \((t = -.12, p = .91)\) significantly differed between girls and boys. Girls had higher re-experiencing symptoms than boys \((M_{girls} = 4.29, M_{boys} = 2.88; t = -1.97, p = .05)\); total PTSS \((t = -1.30, p = .20)\) and hyperarousal symptoms \((t = -1.14, p = .26)\) did not differ. While baseline and trauma SCL were higher in the group sampled at home compared to the group sampled in the lab \((t = -2.57, p = .013\) and \(t = -2.74, p = .008\) respectively), SCR did not differ based on location \((t = -1.20, p \)
= .23). There was no significant effect of interview time on trauma SCL \((p = .20)\) nor SCR, \((p = .90)\).

### 3.2. Feasibility of measuring skin conductance responses in a sample of youth resettled as refugees

To determine whether the trauma interview evoked changes in skin conductance, we compared baseline and trauma SCL. A random intercepts fixed slopes MLM with skin conductance level as the dependent variable and interview segment as the predictor. The MLM indicated a significant effect of phase on skin conductance, \(B = 1.08, t(139) = 5.72, \ p < .001, \ R^2 = .16\) (moderate effect), ICC = .49. Skin conductance levels during the trauma interview \((M = 9.52, SD = 5.27)\) were greater than during baseline \((M = 8.43, SD = 4.86)\) by an average of 1.08 µS, see Figure 2.

### 3.3. Skin conductance, trauma exposure, and psychopathology

We then examined whether SCR to trauma interview (difference between SCL during trauma and baseline SCL) was significantly associated with trauma exposure and PTSS. A multi-level model (MLM) in which participants’ data was nested within families was used to account for shared variance across siblings in the dataset, controlling for the effects of sex and location.

MLMs indicated that trauma exposure was directly correlated with SCR (Figure 3a), see Table 2. SCR was positively associated with re-experiencing (Figure 3c) and hyperarousal symptoms (Figure 3d), but not total post-traumatic stress symptoms (Figure 3b), see Table 3.

### 4. Discussion

The present study examined skin conductance in response to a trauma interview in order to assess associations between autonomic function and trauma and stress in youth resettled as refugees. We also aimed to test the utility of a novel method for measuring skin conductance using an app on an iPad in this unique population. Our results indicated that mobile SCL eSense data collection is easy, low-cost, and feasible in a variety of environments – not just emergency rooms, clinical settings, and research labs as has been previously shown (Hinrichs et al., 2017; Hinrichs et al., 2019), but also natural settings like homes. We found that a higher degree of trauma exposure was associated with greater SCR to trauma interview indicating potential SNS hyperactivation to stress and threat in youth at risk for psychopathology. This finding is supported by the strong positive association between re-experiencing symptom severity and SCR to trauma interview. Such autonomic hyperactivation may be an early risk factor for future mental and physical health problems, and youth who experience
heightened SCR may benefit from mindfulness-based stress reduction and creative arts/movement therapies (Feen-Calligan et al., 2020; Grasser & Javanbakht, 2021; Grasser & Jovanovic, 2021; Grasser, Al-Saghir, Wanna, Spinei, & Javanbakht, 2019). The association between trauma exposure and SCR to trauma interview replicates that from another child and adolescent sample from a different demographic background – Black Americans – further supporting the notion that trauma may affect autonomic reactivity in highly exposed youth, however, causal conclusions cannot be made from either dataset (Wiltshire et al., 2022).

In contrast to previous findings from adults exposed to trauma, (Hinrichs et al., 2017; Hinrichs et al., 2019), our data did not indicate a significant association between total PTSS and SCR to trauma interview. Future studies in larger samples may be better powered to detect a significant relation replicating the findings in adults, given the small effect we observed in this sample. Alternatively, this method may show specificity towards posttraumatic stress symptoms related to autonomic reactivity – namely interview.

**Table 2.** Effect of trauma exposure on skin conductance response.

| Predictors            | Estimates | 95% CI       | p    |
|-----------------------|-----------|--------------|------|
| Intercept             | 3.25      | 1.99 - 4.51  | <.001|
| Sex (Ref = 0; Female) | −0.68     | −1.78 - 0.43 | .235 |
| Location (Ref = 0; Home) | −0.71   | −1.84 - 0.42 | .223 |
| Trauma Exposure       | 0.14      | 0.01 - 0.27  | .042 |

Note: R² = .084, ICC < .001. The estimate for trauma exposure reflect the degree of change in SCR per exposure. Results did not survive Bonferroni correction for multiple comparisons at p < .0125.

**Table 3.** Effects of SCR on total PTSS (R² = .095, ICC = .13), reexperiencing (R² = .127, ICC = .03), and hyperarousal symptoms (R² = .123, ICC = .20).

| Predictors | Total PTSS  | Reexperiencing | Hyperarousal |
|------------|-------------|----------------|--------------|
|            | Estimates   | 95% CI         | Estimates    | 95% CI       | Estimates | 95% CI       |
| Intercept  | 15.25       | 5.22 - 25.28   | 2.36         | 0.64 - 4.08  | 3.50      | 1.08 - 5.92  |
| Sex (Ref = 0; Female) | −4.28     | −12.06 - 3.49  | −1.20        | −2.58 - 0.18 | −0.84     | −2.67 - 0.99 |
| Location (Ref = 0; Home) | 8.39     | −0.46 - 17.25  | 1.38         | −0.06 - 2.82 | 2.31      | 0.10 - 4.51  |
| SCR        | 1.22        | −0.24 - 2.69   | 0.30         | 0.04 - 0.65  | 0.36      | 0.01 - 0.70  |

Note: Results did not survive Bonferroni correction for multiple comparisons at p < .0125. Estimates for sex and location reflect the mean difference in the outcome variable between groups (female/male sex and home/lab assessment location). The estimates for SCR reflect the degree of change in the outcome variable for every one unit change (1 µS) in SCR.
reexperiencing symptoms, as well as hyperarousal symptoms. As this is the first application of the mobile eSense method in refugee youth, these data are promising and merit further inquiry.

Looking at specific symptom clusters, we initially found an association between SCR to trauma interview and re-experiencing symptoms, as well as hyperarousal symptoms, with a stronger effect observed for re-experiencing symptoms. Given the nature of the trauma narrative, it is not surprising that those with greater re-experiencing symptoms had higher SCR throughout the trauma interview. Recalling traumatic memories in the interview setting may model a re-experiencing event; therefore, greater severity of re-experiencing symptoms may be associated with reminders of the trauma and predict physiological arousal in this context. Whereas re-experiencing symptoms may predict task-specific autonomic arousal, hyperarousal symptoms may be linked to more tonic autonomic states. In adults, salivary alpha amylase and heart rate variability have also been associated with re-experiencing symptoms (Rombold et al., 2016a, 2016b; Rombold-Bruehl et al., 2017; Schultebraucks et al., 2019); more broadly, startle response and fear-potentiated startle have been correlated with trauma-related psychopathology (Grasser & Jovanovic, 2021), and other physiological responses of the autonomic nervous system – including respiratory rate and blood pressure (Michopoulos et al., 2015) – may constitute candidate biomarkers of PTSS. Future research that investigates these and other biomarkers alongside SCR to trauma interview may provide convergent validity (Krabbe, 2017) for SCR to trauma interview as a biomarker of re-experiencing symptoms and for SCR to trauma interview as a biomarker of PTSS more broadly.

The focus on biomarkers research has been motivated by the need to rapidly and objectively identify individuals in need of clinical care, both for early intervention and in cases where resources may be limited. This is of particular importance in psychiatry, as the mental health provider shortage affects 124 million Americans (Weiner, 2018). Importantly, the aim of biomarkers research and application to clinical practice is not to replace psychometrics and structured clinical interviews (Loś & Waszkiewicz, 2021). Rather, biomarkers could add to evaluation strategies and may enhance detection – particularly prodromal detection – of psychiatric illness that may, in part, arise from alteration or dysfunction of the nervous system that can be detected through biomarkers (Kalia & Costa e Silva, 2015). Additionally, with substantive validation and replication, biomarkers may be more rapidly obtained than full clinical interviews to provide an initial assessment of which individuals require further follow-up with clinicians in situations where there is limited clinician availability (Kalia & Costa e Silva, 2015) and when even standardized measures fail to identify some disorders in up to 50% of affected individuals (Eack, Greeno, & Lee, 2006). Objective biomarkers could not only help us deliver care to those who need it most, but also identify what treatments may be of best fit for patients with greater specificity than diagnoses alone as there is heterogeneity within psychiatric disease states (Galatzer-Levy & Bryant, 2013; Malhi, 2014; Woods et al., 2012), track treatment efficacy, understand underlying mechanisms of interventions, and define new therapeutic targets (Lydiard & Nemeroff, 2019; Loś & Waszkiewicz, 2021; Scarr et al., 2015). Yet despite the growing number of biomarker studies, the field remains inconclusive. Major limitations in this area of research have included small sample size and unmeasured or unmodeled variables that may account for variation in findings – including biological sex, geographic ancestry, and environmental exposures. As a result, there are currently no existing biomarkers for any psychiatric condition, including anxiety and PTSD (Prasadt, Chaichi, Parker Kelley, Fancis, & Ranjan Garcia, 2019). The ability to identify individuals who may be at risk for a trauma-related disorder like PTSD with greater sensitivity and earlier detection than psychological assessments and clinical interviews alone (Mayeux, 2004) would be of great benefit so that treatment could be delivered early on after trauma exposure or during development, which may enhance treatment outcomes or even prevent the development of psychiatric disease (H. Abu Suhaiban et al., 2019; Prasadt et al., 2019). Despite this, there are limited biomarker studies in youth, although the field continues to grow. The present study provides evidence for feasibility of SCR to trauma interview as a candidate biomarker of autonomic alterations related to reexperiencing and hyperarousal symptoms. The developmental sample described herein provides the unique and important opportunity to assess a candidate biomarker of autonomic functioning linked to trauma-related psychopathology in youth. This is the first study to look at SCR to trauma interview in youth resettled as refugees and is part of the limited but growing body of research to look at biomarkers in refugee cohorts more broadly. We found that not only were symptoms associated with SCR, but also cumulative trauma exposure. Given that trauma exposure was highly correlated with symptoms, and the associations between symptoms and SCR were small, the present data are unable to disentangle whether SCR to trauma interview may be uniquely associated with symptoms above and beyond that of trauma exposure, or whether the elevation in SCR to trauma interview may possibly be, in part, affected by trauma exposure. This elevation in autonomic
functioning associated with higher trauma exposure may precede trauma-related psychopathology, however, this has yet to be measured.

Of great community and public health significance, the method used in this study can be used in multiple settings – not only labs and clinics, but also homes and other community sites. Our data indicated that regardless of variation due to location of assessment, the main effects remain significant. When persons resettling as refugees first arrive in the United States, a mandatory health screening is conducted within one month of arrival. While some clinics will conduct mental health screenings, this is not a required component of the screening, and full clinical interviews are not possible given time constraints and lack of access to mental health clinicians. Here, rapidly obtained, objective biomarkers may provide a time-sensitive and cost-effective option to identify those in need of further screening and additional mental health resources. Objective assessments may be especially important when working with immigrant and refugee communities, where verbal presentation of symptoms may differ based on cultural norms and language may be a barrier. Additionally, screening for trauma exposure and autonomic reactivity in child and adolescent medical settings will provide an opportunity for early detection, intervention, and treatment (Gilgoff, Singh, Koita, Gentile, & Marques, 2020).

4.1. Limitations

The major limitation of this study is the small sample size, as we have discussed above. Notably, while our data provided evidence for small to moderate effects, our results did not survive correction for multiple comparison and therefore results should be interpreted with caution. Another limitation of this study is the nature of self-report questionnaires and the lack of clinical interviews. The present data were collected distally from the initial warzone trauma exposure and at least one year into the resettlement period. Therefore, acute symptoms and psychophysiological changes may have subsided, potentially indicating some level of recovery from stress exposure in this sample, although this was not directly measured. At the same time, aligned with clinical observations, our findings suggest that solely being away from the warzone even after several years does not completely resolve all symptoms. Given the single timepoint data collection, we are unable to resolve whether elevated SCR is a predisposing risk factor for developing PTSS/PTSD once one has experienced a traumatic event (McKeever & Huff, 2003), or if elevated SCR results from trauma exposure, particularly cumulative trauma exposure. Regardless, elevated SCR to trauma interview may still be a meaningful indicator of PTSS. Finally, due to the limitation of single timepoint data collection, we are unable to measure whether SCL and SCR habituate upon repeated measurement using this method. This is an outstanding research question that future studies may seek to resolve, especially if this method is to be used to track longitudinal course of illness or treatment efficacy.

5. Conclusions

We found that SCL measured using mobile eSense was feasible for use in multiple settings, and trauma interviews were effective in evoking measurable elevations in SCL in youth resettled as refugees in the US. Greater trauma exposure may drive greater autonomic reactivity in youth, potentially putting them at risk for future mental and physical health conditions. As such, youth should be screened not only for trauma exposure but also the impact of trauma exposure on psychophysiological markers to better inform care and prioritization of resources. The SCR to trauma interview may provide indications of trauma-related psychopathology – specifically re-experiencing and hyperarousal symptoms – and this study was the first to use this method in this especially vulnerable population. Larger future studies can build on this initial evidence to further query SCR to trauma interview as a biomarker of autonomic functioning related to posttraumatic stress in youth. Detecting and treating PTSS in refugee populations with scarce mental health resources is extremely important given the current global climate. Detection and risk assessment are essential first steps, and subsequently must be the development of culturally appropriate interventions that can be deployed to areas of conflict (Sijbrandij et al., 2017) and resettlement (Acarturk et al., 2022; de Graaff et al., 2020; Feen-Calligan et al., 2020; Grasser et al., 2019; Grasser & Javanbakht, 2021; Purgato et al., 2019; Purgato et al., 2021; Tol et al., 2018). As the needs for trauma-informed services are likely to become overwhelming in many areas – on top of recent increased burden on healthcare services – early detection of, and intervention for, trauma-related psychopathology is critical.

Data availability

Wayne State University requires data use agreements to be drafted and approved prior to any data sharing. If investigators are interested in accessing data, agreements will need to be drafted and approved between institutions and investigators in order to protect these data. Therefore, data cannot be made publicly available at this time, however, data can be made available by request to the authors and institutional approval with an appropriate data use agreement.
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