Finding the golden lining
Assessment, self-help and treatment after trauma
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Screening for trauma-related symptoms via a smartphone app: the validity of Smart Assessment on your Mobile (SAM) in referred police officers

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Chapter 2

Abstract

Background: To facilitate easily accessible screening for trauma-related symptoms, a web-based application called Smart Assessment on your Mobile (SAM) was developed. In this study, we examined whether SAM was able to accurately identify posttraumatic stress disorder (PTSD) and depression in adults.

Methods: Eighty-nine referred police officers completed SAM, containing the PTSD Checklist for DSM-5 (PCL-5) and the Depression Anxiety and Stress Scale (DASS-21), on their own device prior to a diagnostic interview where the Clinician-Administered PTSD Scale for DSM-5 (CAPS-5) and Structured Clinical Interview for DSM-IV (SCID-I/P) were administered.

Results: Results showed a substantial agreement between SAM and the diagnostic interview in the assessment of PTSD and depression. An optimal trade-off between sensitivity (89%) and specificity (68%) levels was found at a cut-off score of 31 on the PCL-5 (area under the curve = .845 [95% CI .765 to .925], diagnostic odds ratio = 15.97).

Conclusions: This is one of the first studies to support the validity and reliability of a mobile screener following trauma. SAM may facilitate screening for trauma-related symptoms on a large scale and could be a first step in a stepped care model for trauma survivors to help identify individuals who need further diagnostics and care.
Introduction

It is widely acknowledged that most of us will be confronted with a potential traumatic event (PTE) during our life, such as loss, serious injuries, or (sexual) violence (Benjet et al., 2016; Breslau, 2009; de Vries & Olff, 2009; Kilpatrick et al., 2013). After experiencing a PTE, individuals are at increased risk to develop profound psychological problems, like posttraumatic stress disorder (PTSD) and depression (Benjet et al., 2016; Breslau, 2009; de Vries & Olff, 2009). PTSD and depression are both listed at the top of the most common psychiatric disorders (Kessler, Petukhova, Sampson, Zaslavsky, & Wittchen, 2012; O'Donnell, Creamer, & Pattison, 2004; Shalev et al., 1998). Despite the fact that a considerable proportion of people suffers from these psychological problems after trauma, the majority of people do not seek professional help, which can lead to deterioration and under-treatment of these symptoms (Bijl & Ravelli, 2000; Brackbill, Stellman, Perlman, Walker, & Farfel, 2013; Brewin et al., 2010; Graves et al., 2011; Grubaugh et al., 2005; Shalev, Ankri, Peleg, Israeli-Shalev, & Freedman, 2011).

This treatment gap is unfortunate since effective treatments for trauma-related symptoms exist (Bisson & Andrew, 2007; Cusack et al., 2016). Moreover, it has been proven that the sooner individuals receive treatment after trauma, the fewer symptoms will arise (Roberts, Kitchiner, Kenardy, & Bisson, 2009). Therefore, it is of crucial importance to (timely) detect trauma-related symptoms, such as PTSD and depression, and refer individuals who need professional care as soon as possible. This is however a great challenge. Besides personal factors like embarrassment, avoidant behaviour and the fear of stigma (Corrigan, 2004), regular health care institutions frequently do not have the capacity and resources to reach all trauma survivors and identify the ones who need further diagnostics or care (Brewin et al., 2010; Shalev et al., 2011). Easily accessible and low-cost screening tools could make an important contribution to the (early) detection and appropriate referral of individuals with mental health needs in the aftermath of trauma (Grubaugh et al., 2005; Olff, 2015; Price, Kuhn, Hoffman, Ruzek, & Acieno, 2015; Price, Sawyer, Harris, & Skalka, 2016; Price, Yuen, et al., 2014).

Currently, there is an explosive worldwide growth in smartphone and applications (apps) usage (Donker et al., 2013; Olff, 2015), which offers new possibilities in reaching and delivering care to a wide range of people. Mobile Health (mHealth) is a specific, upcoming field that focuses on using apps to improve medical or mental health care (Istepanaian & Zhang, 2012; Olff, 2015; Price, Yuen, et al., 2014). mHealth provides a great opportunity to enhance the process of (timely) detection of trauma-related symptoms, both after personal, small scale traumatic events and after major disasters (Donker et al., 2013; Olff, 2015; Price, Yuen, et al., 2014).
2015; Price, Yuen, et al., 2014). Via apps, easily accessible and low-cost trauma-related screening instruments can be delivered which may help individuals recognize symptoms, stimulate help seeking behaviour and contribute to appropriate referral (Brewin et al., 2010; Bush, Skopp, Smolenski, Crumpton, & Fairall, 2013; Olff, 2015; Price, Ruggiero, et al., 2014; Price et al., 2016; Price, Yuen, et al., 2014). Although the great potential of mHealth tools in improving (mental) health care is widely recognized, research on the validity, reliability and effectiveness of these tools is rarely conducted and described (Donker et al., 2013; Olff, 2015). In comparison to the increased knowledge on electronic health (eHealth), scientific support for mHealth tools is considerably lacking behind (Donker et al., 2013). Donker et al. (2013) reviewed the empirical literature on mobile apps in general mental health care and identified nearly 5000 studies mainly describing the development and content of the apps. Strikingly, the majority of these studies did not address the validity, reliability or efficacy of the apps. Finally, only 8 studies describing 5 different apps targeting depression, stress and substance use were included in the review as they were the only studies that examined the efficacy of the tools, included a pre- and post-assessment, and described psychological outcome measures. Of these 5 apps, 4 were efficacious in reducing psychological problems (Donker et al., 2013). In addition, some studies have been conducted on mobile apps that screen for the general mental health (BinDhim et al., 2015; Bush et al., 2013), but none of these were specifically aimed at trauma survivors. Regarding eHealth interventions in general and in the psychotrauma field, scientific support is growing and results are promising. For instance, a meta-analysis on the scientific evidence for eHealth targeting PTSD symptoms showed that internet-based therapies (i.e. cognitive behavioural therapy and expressive writing) were more efficacious in reducing PTSD symptoms than control conditions (i.e. wait list, psychoeducation and control writing task) (Donker et al., 2013; Kuester, Niemeyer, & Knaevelsrud, 2016).

Up to now, only a few studies have been performed on the feasibility and usability of mobile devices for assessing mental health after trauma (Donker et al., 2013; Olff, 2015; Price et al., 2015; Price, Ruggiero, et al., 2014; Price et al., 2016). Notably, the previous studies mainly focused on the tools' feasibility and usability. Regarding monitoring, Price et al. (2014) examined the feasibility of monitoring mental health via daily text messages in patients who recently experienced a traumatic injury. Results showed that the satisfaction with and response rate to this mobile monitoring system were high and that text messaging could be a useful and efficient method to communicate with traumatic injury patients, and to monitor their mental health (Price, Ruggiero, et al., 2014). In another study, the usability of a mobile app to monitor post-trauma symptoms in adults with a trauma history was investigated (Price et al., 2016). The participants indicated that usage of the app could improve their recovery from the traumatic event and they preferred an app that would provide immediate
feedback on their mental health status, was easy to use and customizable (Price et al., 2016). In addition, Price et al. (2015) investigated whether responses to a self-report measure on PTSD administered via a mobile device differed from paper administration in a sample of trauma-exposed adults. No differences were found in total and item scores on the PTSD self-report measure between the two administration methods (Price et al., 2015). Importantly, usability and feasibility results from the available studies are very promising, but so far there is no evidence for the validity and reliability of mHealth screening tools following trauma.

In order to address the abovementioned concerns and to facilitate screening for trauma-related symptoms, we designed a web-based app called Smart Assessment on your Mobile (SAM). SAM comprises modules to assess several relevant mental health domains that may be affected after trauma, as well as well-known risk and protective factors that influence mental health. As a first step to examine the validity of SAM, we investigated whether SAM was able to correctly identify PTSD and depression in a sample of referred police officers. We hypothesized that SAM would be a valid (diagnostic) screener, and therefore would be able to accurately assess PTSD and depression.

**Methods**

**Participants**

The sample consisted of trauma-exposed police officers who were referred to the police-outpatient clinic for a diagnostic interview. The period between the experienced traumatic event and the referral (and therefore the diagnostic interview) was at least one month. Between November 2014 and July 2015, 113 individuals were referred to the outpatient-clinic, of whom 94 (83.2%) consented to participate. Five participants (5.3%) did not complete the PTSD Checklist for DSM-5 (PCL-5) within SAM, and were omitted from the analyses, leaving the data of 89 participants (88 police officers and 1 ambulance worker) for statistical analyses. The 89 participants, of whom 67 men (75.3%), had a mean age of 44.8 years ($SD = 12.25$, range 21-67). Most of the participants (87.6%) had a partner, and 80.9% reported completing a medium educational level (comparable to a college degree). Thirty-nine participants (43.8%) reported that they were currently being treated for psychological problems, of whom 29 (74.4%) indicated that this concerned treatment for trauma-related issues (as determined within SAM). During the diagnostic interview, participants could indicate multiple experienced traumatic events. The most commonly experienced types of traumatic events were all work-related: witnessing a deceased person (44.6%), witnessing a severe accident (37.3%) and being confronted with violence (28.9%).
Procedure
All participants were scheduled for a diagnostic interview at the police-outpatient clinic for the assessment of trauma-related symptoms. The police-outpatient clinic is an independent national centre in the Netherlands where clinical diagnosticians, psychologist and psychiatrists are specialized in investigating trauma-related symptoms in police officers (van der Meer et al., 2016). Researchers contacted the participants by telephone prior to their diagnostic interview to provide information about the study aim and procedures. The researchers explained by phone that the aim of the study was to investigate if a mobile application was able to correctly assess mental health and identify trauma-related symptoms after stressful events. It was explained that the mobile application contained questions about mental health, particularly following trauma, and that the app did not provide feedback on mental health status. Participants were asked to complete SAM once, prior to their scheduled diagnostic interview, on their own personal device. Participants could decide when and where they wanted to fill in SAM.

Participants were informed that all obtained data would be treated as strictly confidential and that only the researchers would have access to the data. The clinical diagnosticians at the police-outpatient clinic were blind to the participants’ results in SAM. If a person orally consented to participate in the study, the researchers e-mailed a digital study information brochure and a web link to open SAM. Participants were asked to complete all the questions within SAM in one session. The mean time between completing SAM and the diagnostic interview was 5.6 days ($SD = 4.6$). Of the 89 participants, 76.4% completed SAM within a week prior to the diagnostic interview. This study was conducted in compliance with ethical principles. The Medical Ethical Committee of the Academic Medical Centre exempted this study from formal review.

Measures

**SAM**
SAM is a web-based app that can be used on a smartphone, PC or tablet. SAM assesses several mental health domains that might be affected after PTEs, including PTSD symptoms, general functioning, depression, anxiety, stress, work engagement, substance use and physical functioning. In addition, SAM evaluates well-known risk factors (trauma exposure, peritraumatic reactions) as well as protective factors (social support, psychological resilience) that may influence these mental health outcomes. To ensure confidentiality, the obtained data within SAM were encrypted before they were sent from the participant’s device to a secured database. SAM did not provide feedback to the participants about their mental health status or symptom severity. The domains within SAM are measured with
freely available questionnaires. Each questionnaire was presented on a separate screen and was introduced by using the official introduction text of the original scale. Participants were provided with the contact details of the researchers in case of any (technical) problems.

As a first step in validating SAM, this study addresses the accuracy of SAM in assessing PTSD and depressive symptoms after trauma exposure. For the assessment of PTSD symptoms, SAM contains the Dutch version of the PCL-5 without the criterion A component (Boeschoten, Bakker, Jongedijk, & Olff, 2014; Weathers, Litz, et al., 2013). In this version, criterion A is not specifically assessed and the PCL-5 scores are not related to a specific event. The PCL-5 is a widely used self-report questionnaire assessing the 20 DSM-5 PTSD symptoms in the past month. The four measured PTSD symptom clusters are: cluster B re-experiencing (5 items); cluster C avoidance (2 items); cluster D negative alterations in cognitions and mood (7 items) and cluster E hyperarousal (6 items). Each item is rated on a 5-point Likert scale, ranging from ‘0 = not at all’ to ‘4 = extremely’, resulting in a total score ranging from 0 to 80. A certain symptom is present when the PCL-5 score is two (‘moderately’) or higher. An indication for possible PTSD based on PCL-5 scores may be derived in two ways: 1) by applying the DSM-5 diagnostic rule, which requires the presence of ≥ 1 symptom in cluster B and C and ≥ 2 symptoms in cluster D and E, or 2) by applying the currently suggested cut-off point of ≥ 33 (http://www.ptsd.va.gov/professional/assessment/adult-sr/ptsd-checklist.asp). Several studies have found good psychometric properties for the PCL-5, with good internal consistency (ranging from 0.56 to 0.96 for the total scale), test-retest reliability, convergent validity, discriminant validity and diagnostic utility (Armour, Contractor, Shea, Elhai, & Pietrzak, 2016; Ashbaugh, Houle-Johnson, Herbert, El-Hage, & Brunet, 2016; Blevins, Weathers, Davis, Witte, & Domino, 2015; Bovin et al., 2016; Wortmann et al., 2016). The internal consistency in the current study was excellent for the PCL-5 total severity score, α = .93, and good for all subscales: re-experiencing, α = .87; avoidance, α = .84; negative alterations in cognitions and mood, α = .83; hyperarousal, α = .76.

To evaluate depressive symptoms, SAM contains the short Dutch version of the well-established Depression Anxiety and Stress Scale (DASS-21) (de Beurs, Van Dyck, Marquenie, Lange, & Blonk, 2001). Participants indicated to what extent the statements applied to them on a 4-point Likert scale, ranging from ‘0 = not at all’ to ‘3 = very often’. The depression subscale consists of seven items, and each score has to be multiplied by two to calculate the final score. A score between 21 and 27 on the depression subscale indicates severe depressive symptoms. The instrument has good reliability and good test-retest reliability (de Beurs et al., 2001). Internal consistency for the depression subscale in the current sample was excellent, α = .91.
Concerning demographics and psychiatric history, SAM asks for gender, age, education and marital status. Furthermore, participants indicated if they were treated (currently) for psychological problems, and if so, for what kind of psychological problems.

**Diagnostic interview**

The clinical diagnostician of the police-outpatient clinic conducted an unstructured interview with the participant about the current psychological symptoms and the potential causes and event(s) that led to the development of these symptoms. Thereafter, the clinical diagnostician determined the presence of a traumatic event according to criterion A. Criterion A requires an exposure to actual or threatened death, serious injury, or sexual violence by either experiencing it, personally witnessing it, learning that the event(s) occurred to a close family member or close friend; and/or experiencing repeated or extreme exposure to aversive details of the event(s) (American Psychiatric Association, 2013). If more than one traumatic event was experienced, the clinical diagnostician asked which of the experienced events currently had the most negative impact on the participant’s life (index trauma). Each index trauma was categorized by the researchers in work-related vs. private events.

During the diagnostic interview, the golden standard Clinician-Administered PTSD Scale for DSM-5 (CAPS-5) was used to measure PTSD symptoms in the past month and to determine DSM-5 diagnosis for PTSD (Boeschoten, Bakker, Jongedijk, van Minnen, et al., 2014; Weathers, Blake, et al., 2013) . The clinical diagnostician only administered the CAPS-5 when the A criterion was met. The CAPS-5 is a 30-item structured clinical interview. Items are rated with a single severity score, ranging from ‘0 = absent’ to ‘4 = extreme/incapacitating’. A symptom is considered endorsed if the severity score is two (‘moderate/threshold’) or higher. The total symptom severity score (range 0-80) is determined by summing up the severity scores of all 20 DSM-5 symptoms. A PTSD diagnosis is established by following the diagnostic rule for DSM-5 symptom criteria. Furthermore, the disturbances have to last at least one month (criterion F) and should cause either clinically significant distress or functional impairment (criterion G). In this study, the internal consistency was good for the CAPS-5 total severity score, \( \alpha = .90 \), and all subscales: *re-experiencing*, \( \alpha = .82 \); *avoidance*, \( \alpha = .66 \); *negative alterations in cognitions and mood*, \( \alpha = .76 \); *hyperarousal*, \( \alpha = .76 \).

To determine Axis I diagnoses other than PTSD, the Dutch version (Van Groenestijn, Akkerhuis, Kupka, Schneider, & Nolen, 1999) of the Structured Clinical Interview for DSM-IV (SCID-I/P) (Spitzer, Gibbon, Janet, & Janet, 1996) was administered. The SCID-I/P measures several Axis I disorders, including mood disorders.
Statistical analyses

The internal consistency was determined by calculating Cronbach’s alpha for PCL-5 and CAPS-5 total and subscale scores and the DASS-21 depression subscale. Further, we calculated descriptive statistics for age, gender, educational level, current psychological treatment, traumatic events, PTSD symptoms (PCL-5 and CAPS-5) and PTSD diagnosis (CAPS-5).

To examine the agreement between the indication for possible PTSD in SAM (PCL-5) and the clinician derived PTSD diagnosis (CAPS-5) the following statistics were calculated between the two measurements: the number of true positives (TP; individuals with PTSD as assessed with the CAPS-5 correctly identified by the PCL-5 in SAM), true negatives (TN; individuals without PTSD correctly identified as non-PTSD cases in SAM); false positives (FP; individuals without PTSD incorrectly identified as PTSD cases in SAM), false negatives (FN; individuals with PTSD incorrectly identified as non-PTSD cases in SAM), the observed agreement ($\rho$; percentage of agreement between the PCL-5 and CAPS-5) and kappa ($\kappa$; measure of the magnitude of the observed agreement). In addition, convergent validity was investigated by calculating Spearman’s rho correlations (non-normally distributed variables) between the PCL-5 and CAPS-5 total and subscale scores.

To investigate the diagnostic accuracy for PTSD symptoms of the PCL-5 in SAM, a Receiver Operating Characteristics (ROC) curve was calculated and the sensitivity (the proportion of people with PTSD according to the CAPS-5 correctly identified in SAM), specificity (the proportion of people without PTSD correctly identified in SAM), optimal cut-off point and the Area Under the Curve (AUC) were determined. Also, the Diagnostic Odds Ratio (DOR; ratio between true positives and true negatives) for the optimal cut-off point on the PCL-5 was calculated.

Finally, to examine the comparability between depressive symptoms in SAM (DASS-21 depression subscale) and the clinician derived depression diagnosis (SCID-I/P), the observed agreement, kappa and DOR values were calculated. All analyses were performed in IBM SPSS Statistics version 23. P-values < .05 were considered significant.

Results

PTSD symptoms

Fifty-two (58.4%) participants were diagnosed with PTSD according to the CAPS-5, and had an average CAPS-5 total severity score of 34.88 ($SD = 9.36$, range 18-64). Within SAM, in 56 (62.9%) and in 54 (60.7%) participants the PCL-5 scores indicated possible PTSD, according to
the diagnostic rule and the suggested cut-off of 33, respectively. The observed agreement between the CAPS-5 and the PCL-5 in SAM was 77.5% (κ = .530, diagnostic rule) and 75.3% (κ = .487, suggested cut-off of 33), indicating a moderate inter-rater agreement. The DOR of the PCL-5 in SAM was 11.46 (diagnostic rule), and 8.750 (cut-off of 33). Participants with a clinician-rated PTSD diagnosis had significantly higher scores on the PCL-5 (M = 43.10, SD = 13.55) than participants without a PTSD diagnosis (M = 25.73, SD = 11.01), t(87) = -6.43, p < .001.

Convergent validity for PTSD symptoms
The PCL-5 total score showed a significant positive correlation with the total CAPS-5 score, r = .768, p < .001. This applied as well to the separate subscales; re-experiencing, r = .718, p < .001; avoidance, r = .537, p < .001; negative alterations in cognitions and mood, r = .683, p < .001 and hyperarousal, r = .620, p < .001.

Diagnostic accuracy for PTSD symptoms
The ROC curve for the PCL-5 is shown in Figure 1. The AUC of the PCL-5 was .845 (95%, CI .765 - .925). The sensitivity of the PCL-5 was 81% and the specificity was 68% at the cut-off of 33. The optimal balance between sensitivity and specificity for this study population was found at a cut-off of 31, with a sensitivity of 89%, a specificity of 68%, and an observed agreement of 79.8% (κ = .574). The DOR of the PCL-5 at the cut-off of 31 was 15.97, indicating that participants who score > 31 have a 15.97 higher chance to have PTSD than participants who score < 31 on the PCL-5. Table 1 shows an overview of the TP, TN, FP and FN values for the PCL-5.

<table>
<thead>
<tr>
<th>PCL-5 diagnostic criteria</th>
<th>TP</th>
<th>TN</th>
<th>FP</th>
<th>FN</th>
<th>Total correctly identified cases (n, %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCL-5 diagnostic criteria</td>
<td>44</td>
<td>25</td>
<td>12+</td>
<td>8</td>
<td>69 (77.5%)</td>
</tr>
<tr>
<td>PCL-5 ≥ 33</td>
<td>42</td>
<td>25</td>
<td>12+</td>
<td>10</td>
<td>67 (75.3%)</td>
</tr>
<tr>
<td>PCL-5 ≥ 31</td>
<td>46</td>
<td>25</td>
<td>12+</td>
<td>6</td>
<td>71 (79.8%)</td>
</tr>
</tbody>
</table>

Note. TP = true positives, TN = true negatives, FP = false positives, FN = false negatives, total correctly identified cases = TP + TN

+In two of these 12 participants, the experienced event did not meet criterion A (as determined during the diagnostic interview), therefore the CAPS-5 was not administered beyond criterion A and PTSD was not diagnosed.
Depressive symptoms
According to the SCID at intake, 19 (21.3%) participants were diagnosed with depression. Regarding the DASS-21 depression subscale within SAM, in 20 (22.5%) participants the scores indicated severe depressive symptoms. The agreement between the SCID and the DASS-21 depression subscale in SAM was 76.4% (κ = .328), indicating fair inter-rater agreement. The DOR of the DASS-21 depression subscale was 5.31. Participants with a clinician-rated depressive disorder had significantly higher scores on the DASS-21 depression subscale ($M = 10.84, SD = 4.02$) than participants without a depressive disorder ($M = 6.14, SD = 4.46$), $t(86) = -4.15, p < .001$.

Discussion
The current study is one of the first studies to report on the validity of a mobile screener in the aftermath of trauma. The results show that SAM was able to accurately screen for PTSD and depressive symptoms, and therefore can be considered a valid and reliable (diagnostic) screening tool in the investigated population.

Overall, our results revealed a substantial degree of agreement between SAM and the diagnostic interview in the assessment of PTSD symptoms and depressive symptoms. There was a strong, positive association between the PCL-5 in SAM and CAPS-5 interview
total scores. Similarly, the separate symptom clusters *re-experiencing*, *avoidance*, *negative alterations in cognitions and mood* and *hyperarousal* (as measured by both instruments) showed significant positive correlations, but the association between the PCL-5 and CAPS-5 on the cluster *avoidance* was relatively low. This could be explained by the fact that the subscale only consists of two items, while the other subscales include more items.

An optimal cut-off score of 31 (with a sensitivity of 89% and a specificity of 68%) was found for the PCL-5 in SAM for possible PTSD in our sample of referred police officers. This cut-off is slightly lower than the currently suggested cut-off of 33 (http://www.ptsd.va.gov/professional/assessment/adult-sr/ptsd-checklist.asp), but in line with the results of a recent study that validated the PCL-5 against the CAPS-5 in veterans from a large VA Health Care database (Bovin et al., 2016). Also, a study in undergraduate students showed an optimal cut off point of 31 and 32 for the French and English versions of the PCL-5 respectively (however, this study did not validate the PCL-5 against a DSM-5 golden standard like the CAPS-5) (Ashbaugh et al., 2016). In addition, the sensitivity level of 89% in our current study clearly exceeds the recommended minimal sensitivity level of 80% for PTSD screening instruments (Mouthaan, Sijbrandij, Reitsma, Gersons, & Olff, 2014; O’Donnell, Bryant, Creamer, & Carty, 2008). Moreover, our AUC value of .845 confirmed that the PCL-5 in SAM adequately differentiates between individuals with and without PTSD.

Besides PTSD symptoms, SAM was also able to correctly identify depressive symptoms. Results showed a satisfactory agreement between the self-reported DASS-21 depression subscale within SAM and the SCID during the diagnostic interview. The overall agreement, inter-rater agreement and DOR values were slightly lower for depressive symptoms compared to PTSD symptoms.

Interestingly, a remarkably high response rate (83.2%) and even higher completion rate (94.7%) were found in our study. This may imply that the willingness to use SAM was high, and the app was easy to use. Regarding the willingness to use mobile screeners for mental health, one of the few studies on a mobile screener for mental health found that active-duty soldiers highly preferred their smartphone over computer and paper methods to complete screening measures (Bush et al., 2013). Also, a study on a screening app for depression showed that a large number of people from different countries downloaded the app, and the majority (73.9%) of these individuals completed the depression questionnaire within the app (BinDhim et al., 2015). Although speculative, these findings suggest that the potential uptake and usage of mobile screeners for mental health could be high.
Mobile screeners like SAM may reach a diverse range of trauma-exposed individuals, both in the general population and in high-risk professions where employees are continuously exposed to PTE’s (van der Meer et al., 2016). These tools may be incorporated as a first step in a (cost-)effective stepped care model to help identify individuals who need further diagnostic examination and care (Price et al., 2016). Further developments may enhance the added value and uptake of mobile screeners in the field of trauma. Incorporating personalized feedback to the end-user is recommended to increase the usability and sustained use of post-trauma mobile screeners (Price et al., 2016). In addition, these tools would be even more valuable if they provide contact details of professional health care institutions (Olff, 2015; Price et al., 2016) and are linked to other evidence based mobile apps, such as the PTSD Coach that offers self-help to individuals with PTSD symptoms (Kuhn et al., 2014; Miner et al., 2016; Owen et al., 2015; Possemato, Kuhn, Johnson, Hoffman, & Brooks, 2016). In this manner, the needs of trauma survivors can be (timely) met and the appropriate post-trauma care can be delivered and received (Olff, 2015; Owen et al., 2015; Price, Ruggiero, et al., 2014; Price et al., 2016; Price, Yuen, et al., 2014).

In a next stage, SAM could provide immediate feedback to the end-user and may be investigated as a monitoring tool that informs both the end-user and the clinician about the patient’s mental health. SAM is currently implemented at the police-outpatient clinic prior to the diagnostic interview to guide clinicians in addressing specific mental health domains that require attention during the diagnostic interview. In the current study, SAM was examined as a diagnostic screener for PTSD, therefore, the period between the experienced traumatic event and usage of SAM was at least one month. Future studies should investigate the validity of SAM in detecting more acute trauma-related symptoms, such as acute stress disorder (ASD). ASD symptoms are highly similar to PTSD symptoms, however, ASD is only diagnosed within the first month after trauma (American Psychiatric Association, 2013). Prospective studies are needed to examine the use of SAM as a predictive mobile screener in a large sample of individuals who use SAM directly after trauma. Also, further studies should be conducted in a counterbalanced design, and study the test-retest reliability of SAM and the sensitivity to clinical change. Lastly, the cost-effectiveness of SAM and the potential negative consequences of screening need to be addressed in future research endeavours (e.g. the risk of potentially stigmatizing false positive cases).

Some limitations of the current study must be considered. Our sample consists of referred (male) police officers, allowing limited generalization to other trauma-exposed individuals. A proportion of our sample received treatment for trauma-related issues, and were, presumably, familiar with PTSD symptoms. Therefore, these participants were potentially better able to recognize their PTSD symptoms than participants who did not receive
treatment for trauma-related symptoms. This is the first Dutch study to report on PCL-5 and CAPS-5 comparison, further studies are necessary to confirm the current findings, also with other (offline) methods, study designs and in other trauma-exposed populations.

Our study has several important strengths. The response and completion rates in our study were very high, reducing potential biases and the possibility of including a selective subsample. Furthermore, we used well-established instruments in our study, such as the PCL-5 and CAPS-5. Moreover, the diagnostic interview was performed by highly experienced clinical diagnosticians at the police-outpatient clinic.

To conclude, this study is a crucial first step in the evaluation of the mobile app SAM and showed that SAM is a valid mobile screener for PTSD and depressive symptoms. Mobile screeners such as SAM can be incorporated as a first step in a stepped care model to identify trauma-survivors in need for further diagnostics and care. Moreover, SAM may contribute to (timely) referral to appropriate and professional care if needed. It is of utmost importance to conduct high quality research on the validity and efficacy of mHealth tools before releasing them into the market (Donker et al., 2013; Olff, 2015). In this manner, we may realize the great opportunities that mHealth offers, and provide evidence based tools that truly contribute to the improvement of post-trauma care.

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**Declaration of interest statement**

The authors have no conflicts of interest to declare.
References


Chapter 2


