INTRODUCTION

Post-traumatic stress disorder (PTSD) develops after being exposed to or having witnessed a life-threatening event. Symptoms of PTSD are re-experiencing the event or parts of it in flashbacks or nightmares, hyperarousal and avoiding aspects related to the traumatic event [1]. Somatically, PTSD patients often show various diseases such as cardiovascular, respiratory, gastrointestinal and autoimmune diseases [2]. Even when controlling for risk factors for somatic diseases such as smoke status or somatic conditions such as hypertension, PTSD patients still show an elevated risk for somatic diseases [2, 3].

Based on these findings, a direct link between psychological disorders and somatic diseases is hypothesized. It is assumed that an alteration of the hypothalamic pituitary adrenal-axis (HPA-axis) and the immune system might promote the development of somatic diseases via cytokines [4]. Cytokines, a family of proteins, mediate immune responses to injury, infection or other organismal stress and are associated with inflammatory diseases, e. g. cardiovascular diseases, cancer, or autoimmune disorders [4, 5]. As described in past studies, an activation of the HPA-axis because of stress triggers a cascade resulting in an immunoregulatory function by inhibiting inflammatory cytokines such as Interleukin (IL-) 6 and increasing production of anti-inflammatory cytokines such as IL-10 [6]. Different studies investigated these mechanisms and found alterations of the HPA-axis and the immune system in PTSD patients showing a blunted HPA-axis stress reactivity compared with healthy controls [7, 8]. Regarding the immune system, mainly a so-called ‘low-grade inflammation’ was found in PTSD patients showing increased inflammatory cytokine levels compared with healthy controls [4]. These elevated levels of IL-6 were found to be associated with worse cognitive functioning of PTSD patients [9]. Therefore, it is supposed that these altered mechanisms of the HPA-axis and the immune system might lead to inflammatory body states in PTSD patients promoting the development of the mentioned diseases.

What remains unclear is, if cytokine levels also influence therapy outcome. 30% of patients with psychological disorders during clinical trials and 65% of patients in routine care do not respond to psychotherapy [10]. 30–50% of PTSD patients do not benefit from trauma-focused therapy sufficiently but show substantial residual symptoms after the end of the therapy [11, 12]. For this reason, past studies investigated aspects influencing therapy outcome.
and found various predictive factors [13, 14]. Regarding therapy outcome of PTSD patients, neuroimaging data and data of pre-treatment biomarkers among others were used for predicting therapy outcome [12, 15]. Higher bedtime salivary cortisol and lower urinary cortisol before treatment predicted greater symptom reduction [15, 16]. Studies examining the predictive character of cytokine levels on treatment outcome are rather rare. Rhein et al. (2021) assessed IL-6 levels of PTSD patients after a psychosocial stress test before therapy and the severity of depressive, trauma-related, and somatic symptoms 8 weeks after treatment. It was shown that patients with a high reactivity to the stress test, reflected by high IL-6 levels, showed a more negative therapy outcome. This study first showed an association between cytokine levels and therapy outcome [17].

The mentioned studies show that different biomarkers such as cortisol and cytokines influence therapy outcome of PTSD patients [14–16]. Nevertheless, the role of anti-inflammatory cytokines remains unclear and studies addressing the influence of cytokines on therapy outcome are very rare in general. Therefore, in this study, we assess the predictive character of pro-inflammatory IL-6 and anti-inflammatory IL-10 levels after a psychosocial stress test at the beginning of psychotherapeutical treatment regarding therapy outcome in female PTSD patients.

METHODS

Study sample

Patients were recruited consecutively at the University Medical Center Carl Gustav Carus of the Technische Universität Dresden during the beginning of their inpatient treatment. 17 female PTSD patients between 28 and 58 years old (M = 47.12, SD = 8.52) with fluent German language skills were included in this sample. Exclusion criteria were a lifetime history of substance use disorder, psychotic or bipolar disorder, psychopharmacological or glucocorticoid-containing medication intake (e.g. asthma inhaler, topical cortisone creams), severe medical illnesses (e.g. cancer, autoimmune diseases, diabetes) and pregnancy. The Structured Clinical Interview (SCID-IV) [18] was conducted by trained interviewers for assessing mental disorders based on the DSM-IV-TR. All of the participants did at least show one comorbidity, n = 1 showed the maximum of 5 additional diagnosis to PTSD, most of the patients (n = 9) showed 3 comorbidities. N = 16 PTSD patients did additionally suffer from a major depressive disorder, n = 6 from panic disorder with/ without agoraphobia, n = 9 from social anxiety disorder, n = 8 from somatoform disorder. None of the participants did use contraceptives, n = 14 took medication and n = 10 of them were taking psychotropic drugs. N = 8 patients were married, 3 were divorced, 4 were single and 2 were widowed. 3 of the patients were incapable for work and 6 of them did smoke. All of the participants provided written informed consent and the study procedure was conducted in accordance with the Declaration of Helsinki and approved by the local Ethics Committee of the Medical Faculty of the Technische Universität Dresden, Germany.

Psychosocial stress induction and hormone sampling

At the beginning of the inpatient treatment, patients participated in a standardized stress test, the Trier social stress test (TSST). The TSST reliably induces acute moderate psychosocial stress under laboratory conditions [19, 20]. It requires a mock job interview (5 min) and a mental arithmetic task (5 min) of the participants, which are performed in front of a mock selection committee. The TSST took place after 2 pm in the afternoon to account for the circadian rhythm of cytokine secretion. The female participants were tested in the luteal phase of their menstrual cycle. Blood samples were collected via a venous catheter 15 min and 1 min prior to the TSST as well as 1 min, 10 min, 20 min, 30 min, 45 min and 60 min after the TSST and set to coagulate for 30 min at room temperature. Afterwards, the blood samples were centrifuged at 20 °C for 10 minutes at 2500xg RCF. Blood samples were stored at −80 °C before being assayed for cytokines. Serum cytokine concentrations (IL-6 and IL-10) were determined using highly-sensitive ELISA enzyme-linked immunosorbent assays (IBL International GmbH, Germany).

Therapy procedure

PTSD patients received a multimodal trauma-focused psychotherapy, the mean duration of treatment was 9 weeks (SD = 3.44). Each week, patients participated in two sessions of individual therapy, which included trauma-focused elements such as Eye Movement Desensitization and Reprocessing or trauma-focused cognitive behavioural therapy. Individual therapy sessions were either offered by board-certified psychotherapists or experienced psychologists and were all supervised by an experienced psychotherapist. Furthermore, each week patients received two sessions of group therapy, two sessions of communicative movement therapy, two sessions of art therapy, one session of progressive muscle relaxation, one session of psychoeducation group, one session of training in social competences group and one session of trauma-stabilization group or skills training. If indicated, daily brief sessions with a nurse took place.

Clinical assessment

Two German language self-report questionnaires were handed out to the study participants for a clinical characterization in the beginning of the therapy and as a measure of therapy outcome at the end of therapy. In both, higher questionnaire scores indicate higher symptom burden. The Symptom Checklist (SCL-90-R) [21, 22] assessed the general psychological symptom burden within the last seven days and consists of 90 items. The global severity index (GSI) reflects the degree of general impairment. The SCL-90-R meets high internal consistencies with Cronbach’s α = 0.97 for the global index. Depressiveness was evaluated with the Beck Depression Inventory II (BDI-II) [23] including 21 items that match the DSM-IV-TR major depression criteria. Internal consistencies (Cronbach’s alpha) are high with α = 0.90 to 0.93.

![Fig. 1 Mean (±SE) IL-6 and IL-10 levels across measurement points of the TSST in PTSD patients. IL-6 Interleukin 6, IL-10 Interleukin 10, TSST Trier social stress test, PTSD posttraumatic stress disorder, time points in minutes.](Image 140x71 to 464x253)
Statistical analyses
A power analysis was conducted with StataIC 16 V5 to calculate an appropriate sample size. Therefore, based on a power of 80%, α = 0.05 and R² = 0.50 were assumed. Based on this calculation, a sample size N = 18 was proposed and implemented. One patient had to be excluded because of missing data in the clinical assessment resulting in N = 17. As IL-6 and IL-10 values were skewed, they were subjected to In-transformations [24]. For IL-10 values, there were in total 5.88% of missing data for different patents and time points. Therefore, missing cytokine values were imputed using predictive means matching over 10 iterations using R with the package mice [25]. Pre-/post-differences in subjective symptom burden (SCL-90_R GSI, BDI) were calculated using paired t-tests. Results were considered significant when p < 0.05. Area under the curve with respect to increase (AUCG) and decrease (AUCI) were calculated for cytokine levels (IL-6, IL-10) during the TSST using StataIC 16 V5. To check for multicollinearity before regression analyses, AUC scores and pre-treatment scores were correlated using Pearson correlations. No correlations exceeded 0.7, so no evidence of multicollinearity was found. At first, in regression analyses pre-treatment scores were used as predictors for therapy outcome. Then, values for AUCG and AUCI were additionally added as predictors for therapy outcome (SCL-90 GSI, BDI). R² greater than |0.26| indicate a high goodness-of-fit [27]. ΔR² was computed to assess additional benefit from including AUC scores in the regression analyses.

RESULTS
PTSD patients did show significant lower symptom burden on the BDI after therapy (M = 21.94, SD = 14.87) compared with BDI scores at the beginning (M = 29.81, SD = 11.72; t(15) = 2.50, p = 0.024, d = 1.16). Regarding the SCL-GSI score, there were no significant differences (beginning: M = 1.85, SD = 0.72, end: M = 1.46, SD = 0.97; t(15) = 1.84, p = 0.086). Figure 1 shows cytokine levels (IL-6, IL-10) in PTSD patients for different time points before and after the TSST, Table 1 includes means and standard deviations of IL-6 and IL-10 levels for each time point before and after the TSST. Further analyses of these patterns in comparison with depressive patients and healthy controls can be derived from a former article [28]. Results of Pearson correlation of AUC scores and pre-treatment scores can be derived from Table 2.

In regression analyses, when predicting therapy outcome by pre-treatment scores, SCL-GSI pre-treatment scores significantly predict SCL post-treatment scores (F(1,15) = 5.13, p = 0.039, R² = 0.255) and also BDI pre-treatment scores significantly predict BDI scores after treatment (F(1,15) = 8.05, p = 0.013, R² = 0.349). Results from regression analyses including pre-treatment scores and AUC scores as predictors for therapy outcome can be derived from Table 3. Model fits are consistently higher than R² = 0.26, reflecting good model fits. 69.9% of the variance of SCL-GSI pre-treatment scores is explained by IL-6 AUC scores and pre-treatment SCL-GSI scores. In this model, AUCG (β = 0.853, p = 0.001) and AUCI (β = −0.554, p = 0.024) are significant predictors for SCL-GSI pre-treatment scores whereas SCL-GSI pre-treatment scores are not (β = 0.204, p = 0.287). 74.4% of the variance of BDI post-treatment scores is explained by IL-6 AUC scores and pre-treatment BDI scores. Also in this model, BDI pre-treatment scores do not significantly predict therapy outcome (β = 0.302, p = 0.104) whereas AUCG (β = 0.814, p = 0.001) and AUCI (β = −0.449, p = 0.044) coefficients show significant predictions. For IL-10, explained variances are smaller but the models still show a good model fit (SCL-GSI: R² = 0.571, BDI: R² = 0.545). Regarding SCL-GSI, AUCG_IL-10 (β = −0.509, p = 0.018) and SCL-GSI pre-treatment scores (β = 0.686, p = 0.005) show significant regression coefficients, whereas AUCG_IL-10 does not significantly predict SCL-GSI post-treatment scores (β = 0.686, p = 0.319). The same pattern was found in the regression model predicting BDI post-treatment scores (AUCG-IL10 β = −0.449, p = 0.036; AUCI-IL10 β = −0.158, p = 0.425; BDI pre-treatment scores β = 0.572, p = 0.009). ΔR² were between 0.196 for predicting BDI post-treatment scores with IL-10 AUC scores and pre-treatment scores as predictors and 0.444 for predicting SCL_GSI post-treatment scores with IL-6 AUC scores and pre-treatment scores.

DISCUSSION
In this study, the predictive value of IL-6 and IL-10 cytokine levels with regard on therapy outcome was assessed. It was found that IL-6 and IL-10 levels under acute stress in the beginning of the therapy predict therapy outcome regarding general symptom burden and depressive symptoms. Higher levels of IL-6 under acute stress before therapy are associated with higher general symptom burden and more severe depressive symptoms after therapy, whereas higher levels of IL-10 under acute stress before therapy are associated with reduced general symptom burden and less depressive symptoms after therapy. A high reactivity of IL-6 during acute stress before therapy predicts reduced general

| Table 1. Means and standard deviations of logarithmized IL-6 and IL-10 levels for all time points during the TSST for PTSD patients. |

|              | IL-6 Mean (SD) | IL-10 Mean (SD) |
|--------------|----------------|----------------|
| −15 min      | 0.76 (0.31)    | 0.86 (0.95)    |
| −1 min       | 0.76 (0.27)    | 0.76 (0.50)    |
| +1 min       | 0.85 (0.34)    | 1.05 (0.52)    |
| +10 min      | 0.82 (0.42)    | 0.75 (0.45)    |
| +20 min      | 0.94 (0.41)    | 0.79 (0.48)    |
| +30 min      | 0.99 (0.43)    | 0.97 (0.46)    |
| +45 min      | 1.02 (0.44)    | 1.04 (0.44)    |
| +60 min      | 0.98 (0.41)    | 1.08 (0.44)    |

Notes. SD standard deviations, min minutes.

| Table 2. Pearson correlation of AUC scores and pre-treatment questionnaires. |

|              | AUCG IL-6 | AUCG IL-10 | AUCI IL-6 | AUCI IL-10 | SCL-GSI_b | BDI_b |
|--------------|-----------|------------|-----------|------------|-----------|-------|
| AUCG IL-6    | 1         | 0.512*     | −0.640**  | 0.082      | 0.067     | 0.116 |
| AUCG IL-10   | 0.512*    | 1          | −0.510*   | 0.220      | −0.439    | 0.392 |
| AUCI IL-6    | −0.640**  | −0.510*    | 1         | −0.222     | 0.182     | −0.026 |
| AUCI IL-10   | 0.082     | 0.220      | −0.222    | 1          | −0.408    | 0.019 |
| SCL-GSI_b    | 0.067     | −0.439     | 0.182     | −0.408     | 1         | 0.497* |
| BDI_b        | 0.116     | −0.392     | −0.026    | 0.019      | 0.497*    | 1     |

Note. AUCG Area under the curve with respect to ground, AUCI Area under the curve with respect to increase, SCL_GSI Global severity index of the Symptom Checklist, BDI Beck Depression Inventory, b pre-treatment scores (beginning of therapy), r Pearson correlation coefficient.

*p < 0.05.

**p < 0.01.

Bold values denote significant p values.
In line with the rarely existing literature [17] and adds further symptom burden and less depressive symptoms after therapy. The patients.

predictive role of IL-10 regarding therapy outcome in female PTSD controls and this altered immune system also seems to in

system of PTSD patients shows alterations compared with healthy therapy outcome. Therefore, as past studies show, the immune IL-6 reactivity under acute stress predicts an advantageous 10 before therapy predict better therapy outcome. Additionally, to predict worse therapy outcome in this study, higher levels of IL-

might be the in

functioning of PTSD patients [29]. PTSD patients showed [30]. Therefore, CBT seems to have a positive long-term effect on associated with enhanced immunity six months after treatment [42]. Based on these findings, the question arises how the immune system can be influenced and if this ‘low-grade inflammation’ of PTSD patients can be reduced or normalized. In a meta-analysis it was found that especially cognitive behavioral therapy (CBT) lowers levels of pro-inflammatory cytokines, increase immune cells and increase natural killer cell activity. Moreover, CBT was still associated with enhanced immunity six months after treatment [30]. Therefore, CBT seems to have a positive long-term effect on inflammatory states. If this influence is also present in PTSD patients, needs to be addressed in future studies. Another possibility of influencing elevated cytokine levels of PTSD patients is psychopharmacological treatment. Here, several new approaches targeting cytokine levels as alternatives to the known serotonin-reuptake-inhibitor treatments are discussed. For example, glucocorticoids could inhibit the expression of cytokines such as IL-6 [31, 32]. Also, endocannabinoid signaling was shown to have anti-inflammatory effects, which is why an elevation of endocannabinoid signaling with synthetic cannabinoids is discussed to be effective. Studies already found positive effects of endocannabinoids on trauma-associated nightmares and general PTSD symptoms [33, 34].

This study shows predictive effects of elevated IL-6 and IL-10 levels under acute stress on therapy outcome of PTSD patients. Nevertheless, there are some limitations. The study is slightly underpowered as one patient needed to be excluded because of missing data. Therefore, interpretation of the results needs to be made with caution. In this study, only female participants were assessed. On one hand, the prevalence of PTSD is higher among women compared with men and more women seek for trauma-related treatment [35]. On the other hand, it is known that there are gender differences in the immune response of individuals under acute stress [36]. As the sample size is quite small, we tried to control for as many influencing factors as possible (e. g. differences in immune response between genders, menstruation) in advance to not include too many control variables in the calculations and therefore reduce power of our study. 16 out of 17 patients with multiple diagnosis. On the other hand, patients received a trauma-focused therapy addressing the traumatic

| Table 3. Predictors of SCL_GSI and BDI scores after therapy in regression analyses. |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| SCL_GSI_e | β            | SE | F   | p   | R²  |
| Model 1  | SCL_GSI_b | 0.505* | 0.309 | 5.13 | 0.039 | 0.255 |
| Model 2  | IL-6: AUCG | 0.853** | 0.007 | 10.07 | 0.001 | 0.699 |
|          | IL-6: AUCI | −0.354* | 0.015 | 0.256 |
|          | SCL_GSI_b | 0.204 | 0.207 |
| Model 3  | IL-10: AUCG | −0.509* | 0.007 | 5.76 | 0.010 | 0.571 |
|          | IL-10: AUCI | 0.209 | 0.007 | 0.278 |
|          | SCL_GSI_b | 0.656** | 0.543 |
| ΔR² | Model 2 – Model 1 | 0.444 | 0.316 |
|       | Model 3 – Model 1 | 0.007 |
| BDI_e | β            | SE | F   | p   | R²  |
| Model 1  | BDI_b | 0.591* | 0.275 | 8.05 | 0.013 | 0.349 |
| Model 2  | IL-6: AUCG | 0.814** | 0.102 | 12.57 | <0.001 | 0.744 |
|          | IL-6: AUCI | −0.449* | 0.219 | 0.228 |
|          | BDI_b | 0.302 | 0.110 |
| Model 3  | IL-10: AUCG | −0.449* | 0.107 | 5.19 | 0.014 | 0.545 |
|          | IL-10: AUCI | 0.158 | 0.247 |
|          | BDI_b | 0.572** | 0.204 |
| ΔR² | Model 2 – Model 1 | 0.395 |
|       | Model 3 – Model 1 | 0.196 |

Note. AUCG Area under the curve with respect to ground, AUCI Area under the curve with respect to increase, SCL_GSI Global severity index of the Symptom Checklist, BDI Beck Depression Inventory, b pre-treatment scores (beginning of therapy), e post-treatment scores (end of therapy).

*p < 0.05. **p < 0.01.
symptoms and making the patients more aware of their traumatic experiences. This often leads to a higher symptom burden at first, which does not necessarily decrease through therapy to a smaller symptom burden than before treatment. Additionally, in this study one representative for pro- and anti-inflammatory cytokines was assessed, which gives a first insight of predictive effects of these cytokines regarding therapy outcome but for more general conclusions and further insights, additional parameters of the immune system should be assessed.

To our knowledge this is the first study investigating predictive effects of pro- and anti-inflammatory cytokines on therapy outcome in female PTSD patients. The present study showed that pro-inflammatory IL-6 and anti-inflammatory IL-10 levels under acute stress before therapy predict therapy outcome of female PTSD patients regarding general symptom burden and depressive symptoms. These findings emphasize the importance of the often-shown inflammatory state of PTSD patients and its influence on therapy outcome. Future studies should address underlying mechanisms of the link between inflammation and therapy outcome and find ways to ensure a greater benefit from psychotherapy for PTSD patients with low-grade inflammations.

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AUTHOR CONTRIBUTIONS

JS, PJ and KP were involved in recruiting the sample and data collection. CK contributed his expertise regarding cytokines and analyzed cytokine levels. VR contributed to data processing, data analyzes and writing the manuscript. All of the authors contributed to finalizing the article.

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