

CHAPTER 24

# The HPA-axis and immune function in burnout

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**Abstract:** Burnout results from chronic work stress. Its complaints may be related to HPA-axis disturbances or changes in immune function. In our studies the salivary cortisol awakening response, day-curve, and the suppressed level after dexamethasone intake were not different in a burned-out group compared to a control group. Nor was there a change in cortisol after a treatment period. Higher levels of DHEAS and the monocyte released anti-inflammatory cytokine IL-10 were observed, however T-cell stimulated and dexamethasone inhibited cytokine release were not affected. The increased IL-10 level may be related to an increased sensitivity for infections.

**Keywords:** burnout; chronic stress; cortisol; cytokines; dexamethasone suppression test; DHEAS; follow-up

## Introduction

Burnout is the ultimate outcome of a chronic process in which work stress is supposed to play a decisive role. People with burnout feel extremely fatigued, have become alienated from their work, experience reduced professional competence, and report a whole range of complaints such as depressed mood, increased irritability, inability to relax, disrupted sleep, somatic complaints such as aching muscles, headaches, gastro-intestinal problems, and concentration and memory problems (Maslach et al., 2001). When we assume that burnout is a stress-related syndrome, one may expect to find a disturbance in hypothalamus pituitary adrenal (HPA)-axis functioning. Inadequate

glucocorticoid signaling has been suggested for other stress-related syndromes like post-traumatic stress disorder (PTSD), chronic fatigue syndrome (CFS), and major depression disorder (MDD). Reviewing the literature on burnout and related stress-syndromes has led to the hypothesis that the fatigue symptoms in burnout are related to a state of hypocortisolism, and increased feedback sensitivity of the HPA-axis (Heim et al., 2000). On the other hand, the depressive symptoms would suggest a hypercortisolemic state, and a relative non-suppression in response to dexamethasone (DEX) (Raison and Miller, 2003). Assuming a disturbance of the HPA-axis in burnout, we expected a reduction in burnout complaints to be related to a recovery of this disturbance. A longitudinal study was set up to correlate changes in complaints with changes in salivary cortisol parameters.

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1 Glucocorticoids play a decisive role in immune  
 2 functioning. Cortisol inhibits pro-inflammatory  
 3 cytokine release, e.g., TNF- $\alpha$ , IFN- $\gamma$ , interleukin  
 4 (IL)-6 and IL-1, and stimulates anti-inflammatory  
 5 IL-10 and IL-4 release (Elenkov and Chrousos,  
 6 2002). Chronic psychosocial stress has been related  
 7 to impaired immune functioning leading to physical  
 8 illness. This process may be mediated by  
 9 glucocorticoids through affecting the balance between  
 10 pro- and anti-inflammatory cytokines (Kiecolt-Glaser  
 11 et al., 2002).

13 **Results**

15 The major finding of our study was the absence of  
 16 a disturbance in salivary cortisol parameters in  
 17 burnout. A burnout group ( $n = 74$ ) was compared  
 18 to a healthy control group ( $n = 38$ ). The burnout  
 19 persons were on sick leave, and had received a  
 20 clinical diagnosis for work-related neurasthenia  
 21 according to International Statistical Classification  
 22 of Diseases and Related Health Problems (ICD-  
 23 10) criteria. Primary Diagnostic and Statistical  
 24 Manual of Mental Disorders Edition IV (DSM-  
 25 IV) disorders such as MDD or anxiety disorder  
 26 were excluded. The cortisol awakening response

(CAR) was measured on 2 days at 0, 15, and 30  
 2 min after awakening, and at noon, 18:00 h and  
 3 22:30 h to assess the diurnal cortisol course. A low-  
 4 dose (0.5 mg) DEX was taken to test the feedback  
 5 sensitivity of the HPA-axis. The suppressed cortisol  
 6 level after DEX intake was measured at 0, 15,  
 7 and 30 min after awakening. The cortisol CAR,  
 8 day-curve and suppressed DEX level were not  
 9 different between the burnout and control group  
 10 (Mommersteeg et al., 2006a–c) (Fig. 1). Cortisol  
 11 was not related to fatigue or depression complaints  
 12 within the burnout group, thus showing no indication  
 13 of an opposing hypo- or hyperfunction of  
 14 the HPA-axis, potentially masking the effect in  
 15 burnout.

16 Because there is considerable variation in cortisol  
 17 levels between and within persons, it is quite  
 18 well possible that within a group burnout persons  
 19 the reduction of the burnout complaints will covary  
 20 with the cortisol parameters after a treatment  
 21 and a follow-up period. This possibility was studied  
 22 in the longitudinal part of the previous study  
 23 (Mommersteeg et al., 2006). Burnout complaints  
 24 were significantly reduced after a treatment period,  
 25 without a further reduction at follow-up. Complaints  
 26 remained substantially higher than norm  
 27 scores for a healthy population. Cortisol after

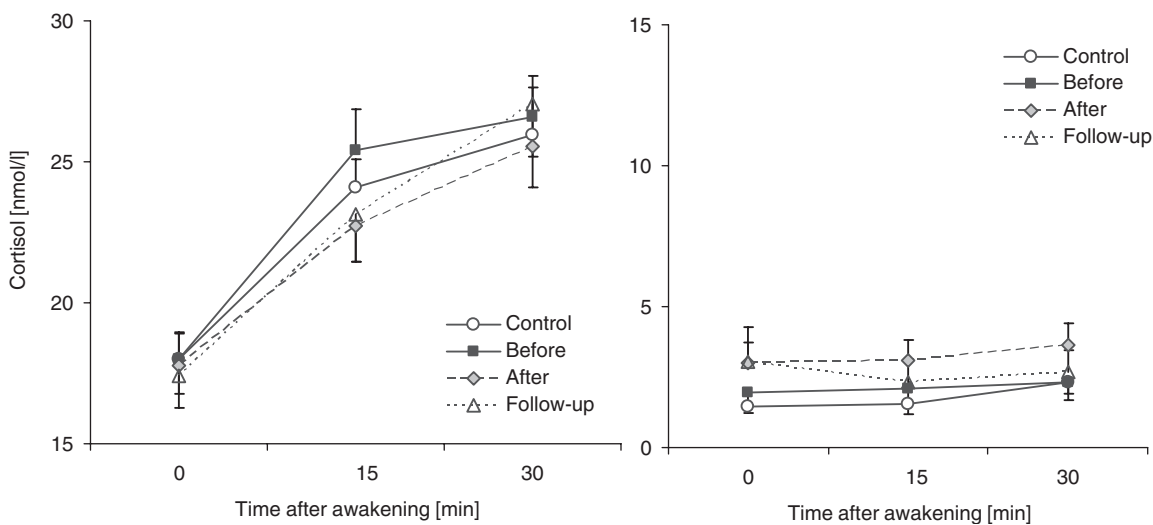


Fig. 1. Cortisol awakening response (CAR, left) and the suppressed CAR after dexamethasone intake (right) in the burnout group before treatment, after treatment and at follow-up, and in the control group. There are no differences between the groups or within the burnout group at consecutive measurements. Means and SEM are shown.

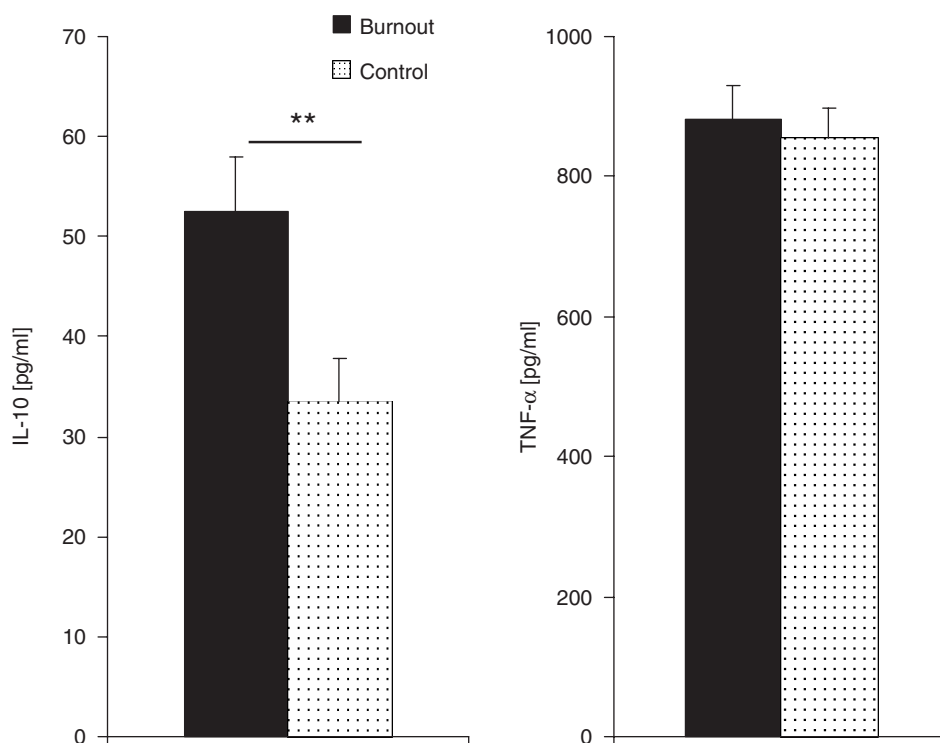


Fig. 2. Anti-inflammatory IL-10 (left) and pro-inflammatory TNF- $\alpha$  release (right) of LPS stimulated monocytes in the burnout and control group. The burnout group had significantly higher levels of stimulated IL-10 ( $F_{(1,83)} = 9.01, p = 0.004$ ). Means and SEM are shown.

awakening and after DEX intake (Fig. 1) showed, however, no parallel changes with complaint reduction. Some isolated associations emerged; the CAR (averaged over the three measurements) was significantly correlated with initial exhaustion level. A decrease in depressive symptoms correlated with an *increased* CAR, whereas the decrease in fatigue in time correlated with a *decrease* of the CAR over the three measurements (Mommersteeg et al., 2006). The latter findings are in contradiction to the supposed hyper- and hypoactive state of the HPA-axis in MDD and CFS, respectively, and moreover explained only a minor part of the variance in complaints within (3%) and between (4%) the burnout individuals.

Immune and endocrine variables were studied in another burnout group ( $n = 56$ ) and compared to 38 controls (Mommersteeg et al., 2006). Again no deviations in the cortisol CAR, or in the DEX suppression test (DST) were observed. The

dehydroepiandrosterone-sulphate (DHEAS) level (but not the cortisol/DHEAS-ratio) was significantly elevated in the burnout group. The burnout group had significantly higher levels of the anti-inflammatory cytokine IL-10 produced by LPS stimulated monocytes (Fig. 2). The IL-10 production of stimulated T-cells, however, was not different from the control group, and neither were there differences in the pro-inflammatory cytokine release of monocyte TNF- $\alpha$  (Fig. 2) or T-cell IFN- $\gamma$ . The capacity of DEX to modulate pro- and anti-inflammatory cytokine release in vitro did not differ between the burnout and the control group, nor was there a change in number of whole blood counts of T-cells, B-cells, and NK-cells.

## Discussion

The results show that there is no discernable disturbance of salivary cortisol in burnout. There is,

1 however, an increased production of IL-10 and  
 3 salivary DHEAS. These findings in a rather large  
 5 sample of clinical burnout persons raise doubts  
 7 about the existence of a relevant neuroendocrine  
 9 dysregulation in burnout as suggested by some  
 11 earlier studies. Still a variety of (neuroendocrine)  
 13 factors may show modest disturbances, altogether  
 leading to a state of ‘allostatic load’ in burnout  
 patients. Though studies in burnout and CFS that  
 included allostatic load parameters do not point in  
 that direction (Cleare, 2003; Grossi et al., 2003;  
 Schnorpfel et al., 2003), this type of approach may  
 be a viable option for further research.

Another option is that central mechanisms are  
 dysregulated in burnout. To test this possibility the  
 combined DEX/corticotrophin releasing hormone  
 (CRH) test, or CRH or adrenocorticotrop hormone  
 (ACTH) infusion are useful techniques. One  
 may doubt however whether these invasive techni-  
 ques are acceptable as a research tool for this  
 (relatively) mild syndrome. Our results point to-  
 ward an increased stimulated monocyte IL-10 re-  
 lease and increased DHEAS levels in burnout.  
 DHEAS has immunostimulatory effects, and at  
 the same time its non-sulphatized form DHEA has  
 been found to reduce susceptibility to viral, bac-  
 terial, and protozoan infections (Chen and Parker,  
 2004). Thus the relevance of the increased DHEAS  
 level in burnout for immune function remains to  
 be determined. Macrophage IL-10 release inhibits  
 T-cell proliferation and suppresses the release of  
 pro-inflammatory cytokines like the anti-viral  
 IFN- $\gamma$ . People with burnout report more common  
 cold and flu-like infections (Mohren et al., 2003).  
 Moreover, vital exhaustion is related to an in-  
 creased pathogen burden, with higher IL-10 serum  
 levels (van der Ven et al., 2003). Therefore an in-  
 creased IL-10 response in burnout may be related  
 to an increased sensitivity for viral infections. Fu-  
 ture studies should reveal the relevance of these  
 findings.

When we started this research project we hy-  
 pothesized that the HPA-axis should show distur-  
 bances in burnout. The results showed the absence  
 of any obvious peripheral deviation in salivary  
 cortisol, nor feedback by DEX in burnout. The  
 correlational effects observed in the longitudinal  
 study are too modest to represent any clinical or

diagnostic value. Overall we conclude that in this  
 study no obvious disturbance of the HPA-axis in  
 burnout was demonstrated. The possibility of  
 some disturbance in immune function and the  
 hormone DHEAS in burnout deserves further at-  
 tention, especially in relation to the sensitivity for  
 infections.

**Abbreviations**

ACTH	adrenocorticotrop hormone	11
CAR	cortisol awakening response	13
CFS	chronic fatigue syndrome	15
CRH	corticotrophin releasing hormone	17
DEX	dexamethasone	17
DHEAS	dehydroepiandrosterone-sul- phate	19
DSM-IV	Diagnostic and Statistical Man- ual of Mental Disorders Edition IV	21
DST	dexamethasone suppression test	23
HPA-axis	hypothalamus pituitary adrenal axis	25
ICD-10	International Statistical Classifi- cation of Diseases and Related Health Problems	27
IL	interleukin	29
MDD	major depression disorder	31
PTSD	post-traumatic stress disorder	33

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