Corticotropin-releasing hormone and copeptin as acute stress markers in serial cerebrospinal fluid – first evidence for non-response during insulin hypoglycemia test

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Abstract

**Background:** During stress, arginine vasopressin (AVP) and corticotropin-releasing hormone (CRH) can act as potent hypothalamic stimulators of the hypothalamic pituitary adrenocortical (HPA) axis. Recently, plasma CT-proAVP, also termed copeptin, was found to be a stable and sensitive surrogate marker for AVP release. A valid assessment of both CRH and copeptin in cerebrospinal fluid (CSF) might directly quantify an individual’s current stress level. Here we investigated how concentrations of CRH and copeptin in CSF alter during insulin-induced hypoglycemia test (IHT) - which has been shown to activate both hypothalamic AVP and CRH - and hypothesized an increase of both.

**Methods:** Five healthy young men were studied from 08:00 until 14:00 after over-night fasting. They received an i.v. injection of human insulin (0.1 IE/kg body weight) at 11:02. CSF samples were drawn from a subarachnoidal catheter every 20 minutes from 10:40 to 14:00 for the measurement of CSF CRH and CSF copeptin. Plasma adrenocorticotropic hormone (ACTH) and cortisol were analyzed in parallel.

**Results:** Data could be assessed in three of five subjects, of which two responded to IHT by showing glucose levels of < 40 mg/dl and clinical symptoms (sweating). Despite a 17-fold, respectively 11-fold increase of plasma ACTH, neither our hypothesized increase of CSF CRH nor of CSF copeptin was seen in these two responders.

**Conclusions:** This is the first study investigating CSF CRH and CSF copeptin in man during acute stress. Copeptin had not been measured in CSF before. Our results support recent speculations that CSF CRH is not a measure of hypothalamic but rather extrahypothalamic CRH (non-HPA CRH). Further research on copeptin in CSF is needed.

**Keywords:** corticotropin releasing hormone (CRH), copeptin, serial cerebrospinal fluid, hypothalamic-pituitary-adrenal axis, stress, insulin-induced hypoglycemia test.
1. Background

In humans, stress response is mediated by the hypothalamic-pituitary-adrenal (HPA) axis and the sympathetic nervous system (1). Their dysfunctions have been hypothesized to be associated with the pathophysiology of stress-related psychiatric diseases such as mood and anxiety disorders (2). Corticotropin-releasing hormone (CRH) and its cofactor arginine vasopressin (AVP), also named antidiuretic hormone (ADH), serve as the most important hypothalamic hormones of the HPA axis to stimulate adrenocorticotropic hormone (ACTH) and consecutively cortisol. Therefore, a valid assessment of both CRH and AVP might directly quantify an individual’s current stress level.

Because CRH in plasma mostly stems from intestinal origin, its measurement in cerebrospinal fluid (CSF) as a putative marker of CNS endocrine function has been established and its serial assessment via a lumbar subarachnoidal catheter offers possibilities for dynamic measurement (3). CSF CSH and CSF AVP concentrations (single-point measurements) were reported to predict a major amount of the variation in the ACTH response to exogenously administered CRH in man (4). However, so far there are only few human studies investigating the response of serial CSF CRH to pharmacological or psychological stimulation (5, 6) and CSF AVP has not been measured in this context. Insulin hypoglycemia test offers a powerful stimulus for the hypothalamic release of both secretagogues, as shown in preclinical studies (7). As a reliably assessed and stable surrogate parameter for AVP in plasma, CT-proAVP, also termed copeptin, has recently been established (8). Our group detected copeptin also in human CSF (Wiedemann, personal communication). Here, we investigated in a pilot study whether serial CSF concentrations of CRH and copeptin alter during an insulin hypoglycemia test in healthy man and we hypothesized an increase of both.

2. Methods

2.1. Participants

Five young men (mean age 24.6 years, range 24 – 26; mean body mass index 25.7 kg/m², range 24.2 – 27.2) were studied. Neither of them smoked tobacco since it is reported to be associated with lower CSF CRH (5). All participants were healthy according to medical history including epileptic seizures and endocrine dysfunctions, physical examination, standard blood tests and urinary drug screen. They did not take any prescription and non-prescription drugs. Current or prior psychiatric disorders were excluded on the basis of the Structured Clinical Interview for the DSM-IV (SCID), axis I. With the help of the SCID post-traumatic stress disorder modules trauma screen a history of major psychological trauma was tested negative in each subject. None had a history of shift work or transcontinental flights during the past three months. Written informed consent was obtained from all after full oral and written explanation of the purpose and procedures of the investigation. This study had been approved by the Ethical Committee of the Medical Board Hamburg afore.

2.2. Procedure

Subjects were studied from 08:00 until 14:00 after over-night fasting. They lay in a supine position in a sound proof private room and were not allowed to sleep, eat or drink. An intravenous cannula was placed into a forearm vein at 08:00 kept open by 50 ml/h normal saline. Subsequently, after
applying local anesthesia, a subarachnoidal catheter (Spinocath®, B. Braun Melsungen AG, Melsungen, Germany) was inserted through the lumbar interspace (3-4; 4-5) by a trained anesthesiologist. An intravenous injection of human insulin (0.1 IE/kilograms body weight) in 10 ml of 0.9 % sodium chloride was injected at 11:02 in a single-blind manner. One physician was attendant at all times. We defined responding to the insulin hypoglycemia test (IHT) as a drop of blood sugar below 40 mg/dl or below 50 % of the initial value of blood sugar after insulin injection, accompanied by at least sweating as a clinical symptom. From 10:40 to 14:00 every 20 minutes 2 ml of CSF were asseted through the subarachnoidal catheter (after discarding 0.5 ml of dead space CSF volume) in order to determine CRH and copeptin concentrations in CSF. In the same time manner, 10 ml of blood were drawn every 20 minutes for the measurement of blood sugar, ACTH and cortisol (and stored at -80 °C). Both CSF and blood samples were immediately filled into prechilled tubes and stored at -80 °C until analysis. Probands were regularly asked for clinical symptoms of hypoglycemia such as sweating, increased appetite, tachycardia, neurologic symptoms like confusion, dizziness and somnolence.

2.3. Endocrine analyses

CSF concentrations of CRH were directly measured with a radioimmunoassay and hCRH was used as a standard and N-tyr-hCRH as tracer after labeling with I-125, as previously reported (9). Intra- and interassay coefficients of variation were below 9%, lower limit of detection was 20 pg/ml. Concentrations of copeptin were measured in 50 μl of CSF in a sandwich immunoluminometric assay (Brahms CT-proAVP, Thermo Scientific, Hennigsdorf, Germany), which uses one polyclonal and one monoclonal antibody. The analytical assay-sensitivity is 0,4 pmol/L, the functional assay-sensitivity is below 1 pmol/L. A single tube luminometer (Lumat® LB 9508, Berthold Technologies, Bad Wildbad, Germany) was used. Plasma cortisol concentrations were determined in 25 μl serum using a radioimmunoassay with the nucleotid I-125 (DRG, Marburg, Germany) and ACTH by an immunometric assay with the nucleotid I-125 (ACTH IRMA, DiaSorin Deutschland GmbH, Dietzenbach, Germany) as reported previously (10).

3. Results

3.1. Side effects and complications

Out of five subjects, data could be assessed in three. In two cases, no data were obtained due to a complete lack of CSF delivery in one subject and a missing CSF delivery after 11:00 in the other one. No complications (i.e. seizures, unconsciousness) were observed upon insulin injection. Three of five subjects developed mild postural headache within 12 hours of the catheter placement, which fully disappeared the following days. No epidural blood patch was needed for recovery.

3.2. Hypoglycemic response

Of the three subjects in which the lumbar catheter delivered CSF throughout, all displayed some typical clinical symptoms of insulin hypoglycemia such as increased appetite, sweating and fatigue within 40 min after intravenous insulin injection. However, only two subjects fully responded to IHT by showing a significant hypoglycemia in periphery blood tests accompanied by sweating, while the non-responder (subject 3) demonstrated a scarce drop of blood sugar by 44 % after 40 min and lacked sweating.
3.3. Peripheral HPA axis activation

Plasma ACTH showed an 11-fold increase in subject 1 and a 17-fold increase in subject 2 within 40 minutes after intravenous insulin injection whereas no change was observed in the IHT non-responder (subject 3). Plasma cortisol concentrations increased accordingly in the responders. Concentrations of plasma ACTH and plasma cortisol are both displayed in Fig. 1.

![Graph of Plasma ACTH](image1)

![Graph of Plasma Cortisol](image2)

**Figure 1.** Plasma ACTH and plasma cortisol in three subjects undergoing insulin hypoglycemia (0.1 IE/kilograms body weight of human insulin being injected at 11:02).
3.4. CRH and copeptin in serial CSF

Despite the clear-cut peripheral reaction, neither our hypothesized increase of CSF CRH nor of CSF copeptin was seen in the two responders (Fig. 2). The non-responder showed a similar time concentration curve. An unexplained isolated secretion pulse of both parameters was observed at 12:40 in subject 2. A slight trend towards higher concentrations of both substances with the course of time can be observed.

Figure 2. CRH and copeptin in CSF in three subjects undergoing insulin hypoglycemia (0.1 IE/kilograms body weight of human insulin being injected at 11:02).
4. Discussion

To our knowledge, this is the first study investigating CSF CRH and CSF copeptin in man during insulin-induced hypoglycemia. Contrary to our hypothesis, we report first evidence that CSF CRH and CSF copeptin do not respond to insulin hypoglycemia in healthy young men despite considerable peripheral HPA axis activation. CSF copeptin has not been measured before.

Recently, Geracioti at al. (11) measured CRH in serial CSF after presenting a stressful psychological suprahypophysial stimulus. Similar to us, they unpredictedly observed a missing increase in serial CSF CRH in patients with posttraumatic stress disorder during and after exposure to traumatic stimuli. They postulated that the unexpected nonappearance of a rise in CSF CRH may be explained by stress-induced uptake of CSF CRH into brain tissue, increased CRH utilization, increased CRH degradation or to an acute stress-related inhibition or suppression of CRH secretion – mechanisms that could also explain our findings after IHT. Analogously, CSF CRH has been measured after pharmacological suprahypophysial stimuli: Vytilingham and colleagues found a lack of CSF CRH response in man despite peripheral stimulation of the HPA axis by administration of the potent hypothalamic CRH-releaser naloxone (6). The same pattern was seen in rhesus monkeys, where both metyrapone and physostigmine did not lead to an increase of CRH in serial CSF despite a HPA axis activation (12).

In addition, we neither had seen acute suppressive effects of cortisol on CSF CRH in a previous study (13). All of these results further support that CSF CRH does not mirror acute HPA axis activity because it may not reflect neurons projecting from the PVN to the median eminence but rather non-HPA CRH. Altogether the sources and dynamics of CSF CRH still remain unclear, but it has been hypothesized that CRH neurons in limbic, cortical and brainstem regions altogether contribute to the CSF CRH pool (14, 5).

Our novel finding of detectable copeptin in CSF – although not modulated by IHT stimulus in our impressionistic data – encourages further study of its potential regulatory role in health and disease.

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