Endogenous calcitonin gene-related peptide in cerebrospinal fluid and early quality of life and mental health after good-grade spontaneous subarachnoid hemorrhage—a feasibility series

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Abstract
The vasodilatory calcitonin gene-related peptide (CGRP) is excessively released after spontaneous subarachnoid hemorrhage (sSAH) and modulates psycho-behavioral function. In this pilot study, we prospectively analyzed the treatment-specific differences in the secretion of endogenous CGRP into cerebrospinal fluid (CSF) during the acute stage after good-grade sSAH and its impact on self-reported health-related quality of life (hrQoL). Twenty-six consecutive patients (±m = 13.8; mean age 50.6 years) with good-grade sSAH were enrolled (drop out 19% (n = 5): 35% (n = 9) underwent endovascular aneurysm occlusion, 23% (n = 6) microsurgery, and 23% (n = 6) of the patients with perimesencephalic SAH received standardized intensive medical care. An external ventricular drain was inserted within 72 h after the onset of bleeding. CSF was drawn daily from day 1–10. CGRP levels were determined via competitive enzyme immunoassay and calculated as “area under the curve” (AUC). All patients underwent a hrQoL self-report assessment (36-Item Short Form Health Survey (SF-36), ICD-10-Symptom-Rating questionnaire (ISR)) after the onset of sSAH (t1: day 1–35) and at the 6-month follow-up (t2). AUC CGRP (total mean ± SD, 5.7 ± 1.8 ng/ml/24 h) was excessively released into CSF after sSAH. AUC CGRP levels did not differ significantly when dichotomizing the aSAH (5.63 ± 1.77) and pSAH group (5.68 ± 2.08). aSAH patients revealed a higher symptom burden in the ISR supplementary item score (p = 0.021). Multiple logistic regression analyses corroborated increased mean levels of AUC CGRP in CSF at t1 as an independent prognostic factor for a significantly higher symptom burden in most ISR scores (compulsive-obsessive syndrome (OR 5.741, p = 0.018), anxiety (OR 7.748, p = 0.021), depression (OR 2.740, p = 0.005), the supplementary items (OR 2.392, p = 0.004)) and for a poorer performance in the SF-36 physical component summary score (OR 0.177, p = 0.001). In contrast, at t2, CSF AUC CGRP concentrations no longer correlated with hrQoL. To the best of our knowledge, this study is the first to correlate the levels of endogenous CSF CGRP with hrQoL outcome in good-grade sSAH patients. Excessive CGRP release into CSF may have a negative short-term impact on hrQoL and emotional health like anxiety and depression. While subacutely after sSAH, higher CSF levels of the vasodilator CGRP are supposed to be protective against vasospasm-associated cerebral ischemia.
from a psychopathological point of view, our results suggest an involvement of CSF CGRP in the dysregulation of higher integrated behavior.

**Keywords** Calcitonin gene-related peptide · CGRP · Health-related quality of life · Impairment · Neuropsychological outcome · Spontaneous subarachnoid hemorrhage

**Introduction**

Spontaneous subarachnoid hemorrhage (sSAH) represents a complex and still devastating neurovascular disease, associated with substantial morbidity and mortality. Over the past 30 years, advances in neurovascular treatment strategies and specialized neurocritical care have led to decreasing case fatality rates [1, 2] with an absolute annual reduction in 30-day mortality of 0.9% over the past decades [1]. However, the mortality in sSAH patients is still as high as 50% [3]. The in-hospital mortality is estimated 18% [4]. About 85% of the non-traumatic spontaneous events comprise aneurysmal subarachnoid hemorrhages (aSAH) and 10% are non-aneurysmal perimesencephalic subarachnoid hemorrhages (pSAH) [5]. Epidemiologically, aSAH and pSAH are two diseases with different evolutions [6]. Compared with aSAH, pSAH represents a subarachnoid hemorrhage (SAH) entity with a very distinct and usually more benign clinical course [6–8]. The survivors harbor serious risks of neurological dysfunction, functional disability, and cognitive impairment [9, 10], even months to years after ictus [11, 12]. There is a marked disparity between reattained functional independence in up to 70% [13] of the sSAH patients and considerable long-term neuropsychological deficits [9, 10, 14] in up to 94.6% [15] with a reduced health-related quality of life (hrQoL) in 35% of the patients 1 year after sSAH [12, 16], anxiety (in up to 54%) [11, 17], depression (approaching 61.7%) [17, 18], and, in up to two-thirds [9, 19], the inability to reassume one’s previous occupation [19, 20] [9, 14].

The underlying pathomechanisms are poorly understood and deemed to be multifactorially mediated [9, 21, 22]. A combination of focal and diffuse brain injury is assumed, probably due to the primary insult, determined by the severity of the bleeding, or subsequent profound secondary complications, most notably the arterial cerebral vasospasm (CV) and the delayed cerebral ischemia [23]. In light of this, the early identification of reliable predictive outcome parameters in sSAH is as tempting as challenging. When evaluating the potential pathophysiological role of vasoactive endogenous neuropeptides in sSAH-related cerebral hemodynamic changes, CV-induced cerebral ischemia, and outcome after sSAH, neuropeptide Y (NPY) [24–27] and calcitonin gene-related peptide (CGRP) [28–33] have gained paramount interest.

The 37-amino acid neuropeptide CGRP, firstly described in 1982 [34], is a highly potent microvascular vasodilator [35] and neuromodulator [36], which is widely expressed and stored in the central and peripheral nervous system [37]. In the cerebral circulation, CGRP is released from presynaptic vesicles in sensory perivascular fibers that almost exclusively originate from the gasserian ganglion [38, 39]. Physiologically, together with other neuropeptides, CGRP restores the cerebrovascular tone in response to vasoconstriction via a remarkable relaxation of the smooth muscle layer, hereby dilating the arteries and, consecutively, increasing the cerebral blood flow (CBF) [29]. Through this dynamic reflex, termed the “trigemino-vascular response”, CGRP opposes excessive vasoconstriction [39]. In aSAH, CGRP has been demonstrated to be excessively released into cerebrospinal fluid (CSF) [31, 33] with a potential neuroprotective effect by preventing CV and cerebral ischemia [33].

Besides its eminent vasoactive role, peptidergic psychoactive implications of CGRP have been repeatedly described in humans and, translationally, in various animal models, with a crucial involvement in multifaceted neurobehavioral processes [40] such as depression [41–47], anxiety [48], learning and memory [49], possibly in dementia [50], in the pathophysiology of inflammatory and neuropathic pain [51–53], and, by unalterable cerebral vasodilation, in migraine [54–56]. To date, the behavioral profile of the action of supraspinal CGRP has insufficiently been elucidated, though.

To the best of our knowledge, no data are yet available on the relevance of CGRP in supratentorial CSF on hrQoL outcome after sSAH in humans. We hypothesize that the excessive release of endogenous CGRP in the subacute phase after sSAH might impact quality of life, even in good-grade patients.

**Patients and methods**

**Ethical approval**

All procedures performed in studies involving human participants (i.e. the clinical database, the prospective liquid biobanking, and the study protocol) were in accordance with the ethical standards of the institutional research committee (Ethikkommission des Universitätsklinikums Regensburg, Ethikvotum 06-179) and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.
Patient population

The cohort has been reported previously [21, 24]. Twenty-six consecutive patients with acute non-traumatic, angiographically confirmed aneurysmal or non-aneurysmal sSAH in prognostically favorable neurological condition were prospectively enrolled in this single-center trial at our University Medical Center between February 2013 and May 2016.

Study selection criteria

After obtaining written informed consent, we selectively included native German speakers, aged 18 to 75 years, with non-traumatic sSAH. The patients either underwent microsurgical aneurysm occlusion (MS group) or endovascular aneurysm occlusion (EV group) for an intracranial aneurysm in the anterior or posterior circulation (aSAH). Patients with a non-aneurysmal pSAH received standardized treatment in the intensive care unit (ICU) (pSAH group). Each patient had been admitted to hospital within 48 h of ictus in prognostically favorable, good to moderate neurological condition, that means with a Hunt and Hess (HH) score [57] of 1 to 4 and an initial Glasgow Coma Scale (GCS) of ≥ 9. Within the first 72 h after the onset of sSAH, all patients received an external ventricular drain because of a radiologically confirmed acute oclusive hydrocephalus. Exclusion criteria were (1) preceding neurosurgical or neurovascular procedures, (2) a previous history of intracranial disorders, (3) a previous history of psychiatric or neurodegenerative diseases, (4) severe autoimmune or systemic diseases, (5) (giant) aneurysm causing mass effect, and (6) severe postprocedural complications, such as intracranial bleeding after treatment of aneurysm or clinically symptomatic cerebral ischemia.

The clinical database comprised all demographic, neurological, and radiological variables, comorbidities, non-invasive procedures, complications, outcome grading (Glasgow Outcome Scale [GOS] [59] and modified Ranking Scale [mRS] [60]), and comprehensive pharmacological screening (at discharge and at the 6-month FU). All patients were examined by means of cerebral computed tomography (CT) and digital subtraction angiography (DSA) and treated according to our ICU standard operating protocol [61]. Transcranial Doppler ultrasound examinations [62] were conducted daily. Our neuroradiologists individually decided on the timing and number of DSA controls on a patient-to-patient basis, depending on the initial DSA findings.

Therapeutic procedures

For the aSAH patients, neurosurgeons and neuroradiologists decided on the treatment modality after interdisciplinary consent. Our standardized surgical and endovascular procedure protocols have been described elsewhere [63].

Self-reported assessment of hrQoL and mental health

Outcome evaluation was conducted in a single session in a noisefree setting by having the participants complete both surveys, as an inpatient at t1 and as an outpatient at t2, respectively. No effects of fatigue were apparent. FU assessment additionally comprised a neurological examination and a non-standardized semi-structured interview, including the patient’s subjective health status, the current medication, and the employment status. All patients completed the German version of the 36-Item Short Form Health Survey (SF-36) [64] (a performance score) and the ICD-10-Symptom-Rating questionnaire (ISR) [65] (a score for symptom burden) in the subacute phase after the onset of bleeding (between day 11 and 35 after sSAH; t1) and in the short term (chronic phase) at the 6-month FU (t2).

ISR The ISR aims at a comprehensive evaluation of the severity of psychological disorders. The ISR 2.0 comprises 29 items and six syndrome scales: depression, anxiety, obsessive/compulsive disorders, somatoform disorders, eating disorders, and a supplementary scale, which covers a variety of syndromes (including concentration, suicidality, sleep, appetite, obliviousness, flash backs, problems with activities of daily living, feelings of displacement and alienation, non-organic sexual dysfunction), as well as a total score. Each syndrome scale ranges from a minimum of 0 (best performance) to a maximum of 4 points with higher scores indicating a more severe symptom burden. Cutoff values for each syndrome scale grade the degree of severity of symptoms in “suspected”, “mild”, “moderate”, and “severe” [65].

SF-36 The SF-36 is a 36-item generic general health questionnaire that yields scores on eight health subscales relating to physical health (physical functioning (Pfi), role limitations due to physical health problems (Rolph), bodily pain (Pain), general health perceptions (Ghp)) and psychological health (vitality (Vital), social functioning (Social), role limitations because of emotional problems (Rolem), and general mental health (Mhi)). These eight subscales can be summarized in a corresponding physical component summary (PCS) and a mental component summary (MCS). The SF-36 also includes a single item that provides an indication of perceived change in health (health transition item, Rawhtran). Each item is scored on a 0 to 100 range and a high score defines a more favorable health state. Items in the same scale are averaged together to create the 8 scale scores [64, 66].
Laboratory procedures

CSF was drawn directly from the external ventricular drain and collected daily over the 10-day period after the onset of sSAH. Immediately after sampling, the samples were centrifuged at 1200 rpm for 10 min, and the supernatants were aliquoted and stored at −80 °C until further use. The samples were thawed, aliquoted (1 ml), evaporated on a vacuum concentrator (Christ RVC 2-25 CD plus; Osterode am Harz, Germany), and dissolved in 250 μl of buffer resulting in a fourfold concentration. CGRP levels were measured in duplicate CSF samples using competitive enzyme immunoassay (EIA; Phoenix Pharmaceuticals Inc., Burlingame, CA). According to pharmacological studies, cerebral exposure to the released endogenous CSF CGRP over time was measured as area under the curve (AUC) and expressed as ng/ml × 24 h.

Statistical analysis

Continuous data and test results on hrQoL and mental health are presented as mean ± standard deviation (SD) and range (minimum to maximum) and categorical data as frequency counts.

hrQoL assessment: Changes over time within each group were analyzed with a paired t test. Differences between groups at postinterventional assessment were analyzed with an analysis of variance (ANOVA) followed by Fisher’s LSD post hoc pairwise comparisons.

Correlation of CGRP with hrQoL assessment: Univariate and multiple logistic regression analyses were conducted for correlations of AUC CGRP with hrQoL test scores and/or clinical variables. Changes over time within each group were analyzed with a paired t test. Intragroup variances (correlations between hrQoL test scores and clinical variables) were analyzed using an analysis of variance (Bartlett’s test for equal variances). Statistical analysis was conducted according to Stata procedures (Stata Version 14.2; Stata Corp. College Station, TX, USA).

A p value of < 0.05 was considered statistically significant.

Results

Demographics and descriptive statistics

During February 2013 and May 2016, a total of 160 sSAH patients of all HH grades was admitted and treated in our medium volume neurovascular center. Among these, 109 patients presented with an acute hydrocephalus requiring CSF drainage via external ventricular drain. Applying our strictly defined selection criteria, only 26 good-grade sSAH patients could initially be enrolled. Another five patients had to be excluded from analysis during FU (lost to FU: n = 3; incompletely answered questionnaires: n = 1; postsurgical bihemispheric chronic subdural hematoma requiring revision and epilepsy: n = 1). Accordingly, 21 consecutive patients with good-grade sSAH (8 men, 13 women) were included as depicted in the flowchart (see Fig. 1). Mean age was 50.6 years (range 27 to 72 years). Our cohort encompassed three patients (14%) with HH grade III and even one patient (5%) with HH grade IV. As all of these four aSAH patients neurologically improved immediately after insertion of an external ventricular drain following hospital admission (i.e., HH I or HH II), the poorer HH score was obviously related to acute occlusive hydrocephalus. With this qualification, we consider the term “good-grade” sSAH patients as appropriate for our cohort. Surprisingly, our good-grade sSAH population exclusively encompassed patients with severe
radiological grades of sSAH (Fisher score 3 or 4) with concomitant hydrocephalus. Consecutively, the presented cohort is imbalanced in terms of the overestimation of the true hydrocephalus rate after sSAH. Statistical intergroup comparisons yielded no significant differences except for a higher number of middle cerebral artery aneurysms in the MS group (n = 3, EV n = 0, pSAH n = 1 (unruptured aneurysm); p = 0.022), a higher intake of antiplatelets in the EV group (n = 5, MS and pSAH n = 0 each; p = 0.016) at t1, an unsurprisingly longer duration (mean ± SD) of MS vs. EV (MS 242.33 ± 24.75 min (range 205–279) vs. EV 136.67 ± 46.87 min (range 55–200); p = 0.004), and a longer mean time (mean ± SD) spent on mechanical ventilatory support in the EV group than in the pSAH group (EV 117.99 ± 264.14 h (range 6.92–808.83); pSAH 5.82 ± 6.34 h (range 0–14.83); MS 116.03 ± 244.14 h (range 2.17–613.63); EV vs. pSAH p = 0.0496; EV vs. MS p = 0.864, MS vs. pSAH p = 0.065). The baseline data including the aneurysm site, GCS, HH, WFNS and Fisher score, procedure variables, medication, and outcome grading has previously been reported [21, 24].

Neuropsychological assessment

The self-reported performance in hrQoL and mental health has been reported previously [21, 24]. We conducted intragroup comparisons and variance analyses: (1) Within 6 months (from t1 to t2), sSAH patients had significantly improved with regard to depression, anxiety, Pfi, Pain, Ghp, Social, and in the PCS. (2) When dichotomizing sSAH versus pSAH patients, no significant differences in terms of SF-36 and ISR scores were detected either except for a significantly higher symptom burden for aSAH patients in the ISR supplementary items score (p = 0.021). (3) Poor self-reported hrQoL performance (ISR scores: total, depression, compulsive-obsessive; physical SF-36 items: Rolph, Pain, Ghp) in the subacute phase correlated with worse outcome on the GOS at discharge. (4) Neurological status at hospital admission in terms of the HH score correlated positively with all psychological SF-36 item scores (compulsive-obsessive syndrome, anxiety, depression, somatoform syndrome, and in the supplementary items score) and with poorer performance in two physical SF-36 items (Ghp and the PCS). The respective regression analyses are depicted by means of scatterplots in Figs. 2a and b and 3a and b. In contrast, at the 6-month FU, CSF AUC CGRP concentrations over the first 10 days no longer showed any significant correlations with hrQoL test performance.

Regression analyses did not reveal any significant correlation between AUC CGRP levels and patient variables like age, gender, treatment modality (MS vs. EV vs. pSAH), site of the aneurysm, HH grade, WFNS grade, Fisher grade, initial GCS, GOS at follow-up, and CV.

AUC CGRP (total 5.7 ± 1.8 ng/ml/24 h) was excessively released into CSF after sSAH. Mean CGRP levels ranged highest in the EV group (6.1 ± 2.1 ng/ml/24 h) followed by the pSAH group (5.7 ± 2.1 ng/ml/24 h), and the MS group (5.0 ± 1.1 ng/ml/24 h). The mean total AUC CGRP level in CSF averaged 5.7 ± 1.8 ng/ml/24 h. The AUC CGRP levels were separately calculated for aSAH (5.63 ± 1.77 ng/ml/24 h) and pSAH (5.68 ± 2.08 ng/ml/24 h) and did not differ significantly (p = 0.521). The analysis of the distinct AUC curves over the first 10 days revealed a highly interindividual pattern without consistency of the AUC dynamics.

Since AUC CGRP concentrations did not significantly differ between the subgroups, subsequent analyses were conducted for the whole study cohort (mean CSF AUC CGRP total level 5.65 ± 1.81 ng/ml × 24 h; range 3.07 to 8.81 ng/ml × 24 h) without differentiation between the sSAH groups. The analysis of cognitive test performances and correlation with the AUC CGRP levels in CSF are summarized in Table 1. Increased mean values of AUC CGRP in CSF at t1 significantly correlated with a higher symptom burden in most ISR scores (compulsive-obsessive syndrome, anxiety, depression, somatoform syndrome, and in the supplementary items score) and with poorer performance in two physical SF-36 items (Ghp and the PCS). The respective regression analyses are depicted by means of scatterplots in Figs. 2a and b and 3a and b. In contrast, at the 6-month FU, CSF AUC CGRP concentrations over the first 10 days no longer showed any significant correlations with hrQoL test performance.

Regression analyses did not reveal any significant correlation between AUC CGRP levels and patient variables like age, gender, treatment modality (MS vs. EV vs. pSAH), site of the aneurysm, HH grade, WFNS grade, Fisher grade, initial GCS, GOS at follow-up, and CV.

After having detected significant correlations between increased AUC CGRP levels and various hrQoL domains in the univariate analysis, we developed a multivariate model including AUC CGRP, age, gender, treatment modality (MS vs. EV vs. pSAH), site of the aneurysm, HH grade, WFNS grade, Fisher grade, initial GCS, and CV to analyze their impact on the respective hrQoL domains. As for the ISR, all factors that significantly correlated with the AUC CGRP in the univariate analysis were also found to be significant in the multivariate model except for somatoform syndrome (OR
As for the SF-36, the PCS remained statistically significant, whereas the Ghp (OR 0.759, \( p = 0.282 \)) were not (cf. Table 2). Conclusively, multiple logistic regression analyses corroborated AUC CGRP as an independent prognostic factor for outcome in terms of compulsive-obsessive syndrome (OR 5.741, \( p = 0.018 \)), anxiety (OR 7.748, \( p = 0.021 \)), depression (OR 2.740, \( p = 0.005 \)), the supplementary items (OR 2.392, \( p = 0.004 \)), and the PCS score (OR 0.177, \( p = 0.001 \)). For the past few decades, in sSAH research, neuroscientists and clinicians have increasingly focused on targeting novel molecular genetic, vascular, inflammatory, oxidative stress and protein biomarkers, especially in the CSF, to reliably predict functional, cognitive, and hrQoL outcomes for individualized treatment strategies [27, 67, 68]. Only one previous study [67] has addressed neurodegenerative ventricular CSF biomarkers in the context of poor hrQoL outcome after aSAH. Our research group [24, 27, 33] and others [31, 32, 68] have corroborated the predictive capacity of endogenous neuropeptides after sSAH, in particular, the vasoactive CGRP.

**Table 1** Health-related quality of life and emotional health of the cohort (\( n = 21 \)) and correlation with the calcitonin gene-related peptide concentrations in supratentorial cerebrospinal fluid in the subacute interval after the onset of spontaneous subarachnoid hemorrhage (\( t_1 \)) and at 6-month follow-up (\( t_2 \))

| Neurropsychological assessment | Test scores (mean ± SD) | Difference \( t_1 \) vs. \( t_2 \) (mean ± SD) | Paired \( t \) test \( t_1 \) vs. \( t_2 \) (\( p \) value) | AUC CGRP (\( p \) value) |
|------------------------------|--------------------------|-----------------------------------------------|-------------------------------------------------|--------------------------|
| ISR scores                   |                          |                                               |                                                 |                          |
| Depression                   | 1.5 ± 1.2                | 1.1 ± 0.9                                     | 0.4 ± 1.1                                       | 0.046*                   | 0.011*                   | 0.431                      |
| Anxiety                      | 1.3 ± 1.1                | 0.7 ± 0.9                                     | 0.7 ± 0.3                                       | 0.019*                   | 0.035*                   | 0.668                      |
| Compulsive-obsessive         | 0.8 ± 1.0                | 0.8 ± 1.0                                     | 0.0 ± 1.4                                       | 0.480                    | 0.022*                   | 0.678                      |
| Somatoform                   | 0.6 ± 0.7                | 0.4 ± 0.6                                     | 0.1 ± 1.0                                       | 0.252                    | 0.013*                   | 0.438                      |
| Nutrition disorder           | 0.6 ± 1.0                | 0.6 ± 0.7                                     | −0.0 ± 1.2                                      | 0.523                    | 0.421                    | 0.898                      |
| Supplementary items          | 0.7 ± 0.5                | 0.5 ± 0.6                                     | 0.2 ± 0.6                                       | 0.064                    | 0.015*                   | 0.114                      |
| Total                        | 0.9 ± 0.7                | 0.7 ± 0.6                                     | 0.2 ± 0.8                                       | 0.098                    | 0.086                    | 0.737                      |
| SF-36 scores                 |                          |                                               |                                                 |                          |
| Physical items               |                          |                                               |                                                 |                          |
| Rawhtran                     | 4.5 ± 0.6                | 2.9 ± 1.4                                     | 1.6 ± 1.4                                       | 1.000                    | 0.066                    | 0.499                      |
| Pfi                          | 19.5 ± 28.9              | 72.4 ± 26.3                                   | −52.8 ± 35.5                                    | 0.001*                   | 0.057                    | 0.080                      |
| Rolph                        | 39.3 ± 43.7              | 47.6 ± 45.3                                   | −8.3 ± 64.4                                     | 0.280                    | 0.090                    | 0.166                      |
| Pain                         | 48.5 ± 34.6              | 67.2 ± 29.9                                   | −18.7 ± 37.9                                    | 0.018*                   | 0.251                    | 0.229                      |
| Ghp                          | 56.3 ± 16.8              | 74.1 ± 18.2                                   | −17.8 ± 19.0                                    | 0.001*                   | 0.043*                   | 0.283                      |
| Psychological items          |                          |                                               |                                                 |                          |
| Vital                        | 51.0 ± 19.7              | 51.2 ± 20.3                                   | −0.2 ± 24.4                                     | 0.482                    | 0.536                    | 0.162                      |
| Social                       | 66.1 ± 26.0              | 80.4 ± 25.2                                   | −14.3 ± 28.6                                    | 0.017*                   | 0.453                    | 0.653                      |
| Rolem                        | 61.7 ± 46.2              | 61.7 ± 45.0                                   | −0.0 ± 58.8                                     | 0.500                    | 0.835                    | 0.102                      |
| Mhi                          | 61.1 ± 20.6              | 68.2 ± 21.7                                   | −7.0 ± 28.6                                     | 0.137                    | 0.621                    | 0.808                      |
| PCS                          | 31.0 ± 11.0              | 46.0 ± 10.0                                   | −13.3 ± 12.7                                    | 0.001*                   | 0.006*                   | 0.083                      |
| MCS                          | 49.1 ± 12.5              | 47.5 ± 10.7                                   | 1.4 ± 12.9                                      | 0.677                    | 0.582                    | 0.549                      |

\( SD \), standard deviation; \( test t_1 \), test in the subacute phase after the onset of bleeding (between day 11 to 35 after subarachnoid hemorrhage); \( test t_2 \), test in the short-term (chronic phase) after treatment at 6-month follow-up; AUC, area under the curve; CGRP, calcitonin gene-related peptide; ISR, ICD-10-Symptom-Rating questionnaire; SF-36, German version of the 36-Item Short Form Health Survey; Rawhtran, health transition item; Pfi, physical functioning; Rolph, role limitations because of physical health problems; Pain, bodily pain; Ghp, general health perceptions; Vital, vitality; Social, social functioning; Rolem, role limitations because of emotional problems; Mhi, general mental health; PCS, physical component summary; MCS, mental component summary

*Statistical significance \( p < 0.05 \)

Discussion

Our prospective study advocated an association of increased CSF CGRP concentrations in the acute phase of sSAH with unfavorable short-term hrQoL.

Calcitonin gene-related peptide propaedeutics

For the past few decades, in sSAH research, neuroscientists and clinicians have increasingly focused on targeting novel molecular genetic, vascular, inflammatory, oxidative stress and protein biomarkers, especially in the CSF, to reliably predict functional, cognitive, and hrQoL outcomes for individualized treatment strategies [27, 67, 68]. Only one previous study [67] has addressed neurodegenerative ventricular CSF biomarkers in the context of poor hrQoL outcome after aSAH. Our research group [24, 27, 33] and others [31, 32, 68] have corroborated the predictive capacity of endogenous neuropeptides after sSAH, in particular, the vaso- and psychoactive CGRP.
isolated from thyroid tissue in 1982 [34] and is derived from the calcitonin gene. CGRP exists in two isoforms, αCGRP and βCGRP. αCGRP, which is synthesized by alternative splicing of the calcitonin gene, represents the predominant form of CGRP within the body and is mainly expressed in the central and peripheral nervous system [34]. CGRP

Fig. 2 Correlation of elevated area under the curve (AUC) values of calcitonin gene-related peptide (CGRP) in cerebrospinal fluid (CSF) with a higher symptom burden in the ICD-10-Symptom-Rating questionnaire (ISR) scores within the first 10 days after the onset of spontaneous subarachnoid hemorrhage. Area under the curve (AUC) values of endogenous calcitonin gene-related peptide (CGRP) in supratentorial cerebrospinal fluid (CSF) within the first 10 days after the onset of spontaneous subarachnoid hemorrhage plotted versus the ICD-10-Symptom-Rating questionnaire (ISR) scores [65] in a depression and b anxiety. The ISR with 29 items and 6 syndrome scales aims at comprehensively evaluating the severity of psychological disorders. Each syndrome scale ranges from a minimum of 0 (best performance) to a maximum of 4 points with higher scores indicating a more severe symptom burden. Each dot represents the mean level of CSF CGRP in [ng/ml × 240 h] for each patient, indicating a significant linear correlation (compare regression line) with a higher symptom burden. *p < 0.05

Fig. 3 Correlation of the area under the curve (AUC) values of calcitonin gene-related peptide (CGRP) in cerebrospinal fluid (CSF) with reduced general health perception and an impaired physical component summary score in the 36-Item Short Form Health Survey (SF-36) within the first 10 days after the onset of spontaneous subarachnoid hemorrhage. Area under the curve (AUC) values of endogenous calcitonin gene-related peptide (CGRP) in supratentorial cerebrospinal fluid (CSF) within the first 10 days after the onset of spontaneous subarachnoid hemorrhage plotted versus the 36-Item Short Form Health Survey (SF-36) [64] scores in a the physical component summary score (PCS) and in b general health perceptions (Ghp). The SF-36 is a 36-item generic general health questionnaire yielding scores on 8 health subscales relating to physical and psychological health. These 8 subscales can be summarized in a corresponding physical component summary and an MCS. Each item is scored in the range 0 to 100, and a high score defines a more favorable state of health. Items in the same scale are averaged together to create the 8 scale scores. Each dot represents the mean level of CSF CGRP in [ng/ml × 240 h] for each patient, indicating a significant linear correlation (compare regression line) with poorer performance in hrQoL and emotional health. *p < 0.05
In accordance with the psycho-behavioral literature, during spinal anesthesia for minor orthopedic or urologic surgery, the concentration of calcitonin gene-related peptide (CGRP) in spinal fluid was studied. From a population of 29 non-neurosurgical patients (15 women and 14 men; mean age 52.8 years), spinal cerebrospinal fluid (CSF) was collected within the first 10 days after surgery. The CGRP concentration (mean 0.09 ng/ml) was found to be significantly lower compared to a historic control population of 29 non-neurosurgical patients (mean level of 0.6 ng/ml). This study provided insight into normal reference values of CSF CGRP.

Our study proves an excessive hypersecretion of CGRP into spinal fluid in the subacute interval after the onset of spontaneous subarachnoid hemorrhage (t1). CGRP constitutes one of the most potent endogenous vasodilators in humans [32, 38]. In the cerebral circulation, CGRP is primarily stored in presynaptic vesicles of sensory fibers that are closely associated with blood vessels [69] and that almost exclusively arise from the trigeminal ganglia [39]. In response to the intrinsic release of vasoconstrictive neuropeptides, CGRP physiologically restores the vascular tone mediating the “trigeminovascular reflex” [29]. Beyond, literature has repeatedly highlighted the abundantly expressed CGRP as a crucial psychoactive mediator in a variety of neurobehavioral and psychosomatic conditions [40]. Thus, establishing the contribution of endogenous CSF CGRP to neuropsychological outcome and health-related quality of life (hrQoL) in subarachnoid hemorrhage (SAH) is rather promising.

### Excessive release of CGRP after sSAH in a neurobehavioral context

Our study proves an excessive hypersecretion of CGRP into CSF (mean level of 0.6 ng/ml) within the first 10 days after SAH. A historic population [33] of 29 non-neurosurgical patients (15 women and 14 men; mean age 52.8 years) has provided insight into normal reference values of CSF CGRP. From these 29 patients, spinal CSF with a remarkably lower CGRP concentration (mean CGRP 0.09 ng/ml) was drawn during spinal anesthesia for minor orthopedic or urologic surgery [33]. In accordance with the psycho-behavioral literature, our study postulates a detrimental effect of CSF CGRP on psychological and physical health: Increased CSF AUC CGRP levels were significantly positively correlated to depression, anxiety, compulsive-obsessive syndrome, and the supplementary ISR items (which imply a variety of concomitant syndromes including problems with activities of daily living, sleep, concentration, flash backs, obliviousness, feelings of displacement and alienation, suicidality, appetite, and non-organic sexual dysfunction). A significantly negative correlation was established between high CSF AUC CGRP concentrations and the PCS score (covering Pfi, Rolph, Pain, and Ghp). At the supraspinal level, CGRP is broadly distributed like, for example, in the sensory and the trigeminal ganglia, the striatum, amygdala, pituitary gland, hypothalamus, medulla oblongata, and in the cortex [34, 37, 40]. The widespread presence of CGRP and its binding sites in the brain, eminently in limbic structures, indicates its potential involvement in a plethora of neurophysiological and neurobehavioral functions [40], like in depression [41, 42, 44], possibly in dementia [50], and in the pathophysiology of inflammatory and neuropathic pain [51–53]. Mathé and collaborators defined CGRP in lumbar CSF as a trait marker of major depression [44]. When administered intracerebroventricularly, intracerebrally, or intravenously in animal models, exogenous CGRP was found to potentiate fear-related behaviors [48], and it was attested a pivotal role in learning and consolidation of memory in passive avoidance tests [49], in locomotion, nociception, depression-like behaviors [43, 45–47], in anorexia [70], and in addiction [71–73]. Our multiple logistic regression analysis indicates a contribution of further clinical variables and, in addition, an interaction of the restrictions in most hrQoL domains.

Anxiety and depression [11, 17, 18] are among the most investigated realms of patient outcome after aSAH with a stable prevalence over the 18-month period after aSAH and an estimated frequency ranging from 27 to 54% and from 5 to 50% in aSAH survivors, respectively [9]. Likewise, somatoform disorders, especially pain syndromes like cephalgia, are a commonly reported and oftentimes a long-lasting symptom burden after SAH, plausibly affecting Ghp, poorer performance in several cognitive domains, and reduced hrQoL after SAH [9, 74]. In this context, the cerebral exposure to CGRP might reveal advanced insight and a potential therapeutic target in the future.

### Table 2

| hrQoL domains | Odds ratio | 95% CI | p value |
|---------------|------------|--------|---------|
| ISR           |            |        |         |
| Compulsive-obsessive syndrome | 5.741 | 1.341 | 24.581 | 0.018* |
| Anxiety       | 7.748      | 1.366 | 43.959 | 0.021* |
| Depression    | 2.740      | 1.360 | 5.519  | 0.005* |
| Somatoform syndrome | 83.991 | .548  | 12,865.310 | 0.084 |
| Supplementary items | 2.392 | 1.328 | 4.310  | 0.004* |
| SF-36         |            |        |         |
| General health perceptions (Ghp) | 0.759 | 0.459 | 1.255  | 0.282 |
| Physical component summary (PCS) | 0.177 | 0.065 | 0.481  | 0.001* |

hrQoL, health-related quality of life; test t1, test in the subacute phase after the onset of bleeding (between day 11 to 35 after subarachnoid hemorrhage); ISR, ICD-10-Symptom-Rating questionnaire; SF-36, German version of the 36-Item Short Form Health Survey

*Statistical significance p < 0.05
Release of CGRP after sSAH

The vasodilatatory CGRP [35] has been demonstrated to be excessively released into CSF [31, 33] and, to a even greater extent, into serum [31, 75] during the first 10 days [33, 75] after SAH. Endogenous CSF CGRP (upregulated on days 1 to 4 after sSAH) [33] and exogenously administered CGRP are postulated to be cerebroprotective and, thus, beneficial for functional outcome by preventing sSAH-induced CV and cerebral ischemia, respectively [28, 29]. The substantiated effects of CGRP on hemodynamics and, consecutively, on neurological outcome after sSAH contrast with the current findings, suggesting at least a contribution of CGRP to psychobehavioral dysregulation and reduced hrQoL.

The neuroanatomical circuitry involved in CGRP transmission and modulation remains to be clarified. Early observational studies on the cerebral circulation after experimental SAH and post mortem analyses after SAH confirmed a marked decrease in CGRP immunoreactivity in the perivascular nerve fibers (cf. references in [40]). Pathophysiologically, the neurotransmitter CGRP is supposed to be released from the perivascular nerve terminals, either induced by the blood in the subarachnoid space [76] or, possibly, caused by the direct affection or disruption of the neuropeptide-containing nerve fibers at the moment of aneurysm rupture and the subsequent inhibition of neuropeptide reuptake at the nerve-ending terminals. We propose that CSF CGRP may be dispersed with the circulating CSF from the basal cisterns into the ventricles where the neuropeptidergic concentrations are amenable to measurement. It is questionable whether CGRP in ventricular CSF reflects the proposed pathophysiological processes or whether it is rather mirroring other processes associated with critical illness.

It might further be speculated that the increase of endogenous CSF CGRP might be a consequence of altered CGRP synthesis and metabolism in certain brain regions and in CSF, respectively, anatomical localization of the ruptured aneurysm, or aneurysm treatment-induced mechanical manipulation of the parent vessel. In serum, peak concentrations of CGRP have been measured after rupture of aneurysms of the middle cerebral artery (MCA) [31, 75] and—regarding cerebrovascular manipulation—after endoluminal aneurysm treatment via coiling [75]. Our sSAH collective comprised a reference group with pSAH patients, characterized by conservative ICU management, to further illuminate the implication of the aneurysm-securing procedure on CGRP release and hrQoL outcome. Mean CGRP levels ranged lowest in the MS group. In line with subgroup analyses, MS patients demonstrated better short-term Pfi, experienced less pain, and more improvement in nutrition disorders than EV patients. As previously described, however, the treatment modality (MS vs. EV vs. pSAH) did not significantly affect CSF AUC CGRP levels, and overall hrQoL outcome did not differ between the MS and the EV group [21], either. We caveat the statement with the note that the small sample size of the current study is not capable for detecting any differences. Since 2002, the treatment modality-dependent outcome is controversially discussed [77]. The majority of authors disproved the hypothesis that clipped and coiled patients differ with respect to cognitive outcome, hrQoL, return to work, depression, anxiety, and sleep disturbances (cf. [9, 78] and references within), though.

It has to be highlighted that—in many ways—pSAH has to be considered a different disease than aSAH. Short-term complications are rare, and long-term outcome is excellent with respect to disability and death [6, 8]. aSAH implies a more aggressive clinical presentation, more diffuse distribution of subarachnoid blood, a higher probability of complications, and a longer inpatient period with higher economical costs for health care systems. Yet, both, aSAH and pSAH patients, suffer from neuropsychological deficits and hrQoL restrictions after ictus [7, 9, 79]. Contrary to former assumptions, which attested pSAH patients a favorable prognosis [80, 81], more recent findings [7, 82, 83] indicated that pSAH might not be as benign as previously believed. On average 39 months after pSAH, survivors continued suffering from cephalgia, dizziness, fatigue, irritability, depression, obliviousness, mild cognitive deficits, and incapacity to resume their previous occupations [82, 83]. Long-term studies on potential cognitive and hrQoL sequelae after pSAH are demanded [8] and should separately address the CGRP effect on outcomes for pSAH and aSAH.

Investigations into the temporary dynamics of sSAH-induced CGRP secretion are scarce and limited to the short-term [84]. As our sSAH patients significantly improved in multiple physical and emotional hrQoL items within 6 months, it might be reasoned that CGRP concentrations decrease or even normalize over time. Hypothetically, extra- and intraluminal CGRP receptors may finally be saturated non-competitively with the peptide, and CSF levels decrease due to the depletion of the releasing terminal nerve endings. In turn, lower CSF CGRP concentrations or reuptake of ventricular CGRP may be beneficial to hrQoL and mental health. In 2013, our research group detected peak concentrations of CSF CGRP during the first 4 days after onset of sSAH, followed by a gradual decrease [33]. Nozaki et al. [85] found the most marked suppression of CGRP immunoreactivity during the 7th to 14th day with a recovery to normal levels by the 42nd day after artificial SAH. Congruently, our results suggest that at least within the first 10 days after sSAH, elevated CSF CGRP levels account for the sSAH-related psychological traumatization and reduced hrQoL. At the 6-month FU, self-reported hrQoL did no longer correlate with the initial CGRP values. Neuropsychological deficits, predominantly in hrQoL, cognition, depression, anxiety, mood, and fatigue [9–11, 17, 21]
are—consistently with our findings—most common within the first 3 months [87] but may persist as long as 24.5 years after ictus or even longer [60, 71, 18, 55, 1].

The pathophysiology of impaired neurobehavioral processing following sSAH remains elusive because of its complex, multifactorial character [22]. Various predictors of unfavorable neuropsychological outcome after aSAH (like the HH score and the GOS in our results section) have been proposed [5, 21]. In cognitively impaired good-grade sSAH patients without morphological changes in neuroimaging, it seems conceivable that the initial insult of the bleeding may result in a widespread derangement of peptide neurotransmitter secretion in the brain [68]. We caveat our findings with the note that our experimental, hypotheses-generating pilot study was not designed to conclusively establish whether pathologically increased neurotransmitter secretion, in particular excessively elevated CGRP levels in supratentorial CSF, induce restrictions in hrQoL and emotional health. However, a contribution seems feasible, given the fact that such associations exist in a myriad of psychiatric and neurobehavioral disorders. Thus, we argue that, in sSAH, CGRP acts as an elementary psychoactive mediator in higher integrated behavior.

Methodological considerations

Bounded by our strict selection criteria and our institutional neurovascular volume, our pilot study is notably limited by the small sample size. It may be speculated that the five excluded patients were incapable to complete the hrQoL assessment due to severe neuropsychological impairment, resulting in an underestimation of the true impairment rate. Then, the inclusion of pSAH patients as a control group implica a certain collective heterogeneity because pSAH and sSAH have a different pathogenesis, clinical course, different rates of CV and delayed ischemic neurological deficits, and also different neuropsychological and hrQoL outcomes. Though pSAH patients are deemed equally burdened by neuropsychological deficits and reduced hrQoL [9], the outcome of aSAH patients is considered even worse. Therefore, future studies should separately address the CGRP effect on outcomes in both entities. Additionally, our study is severely biased by (1) the typical [78] overrepresentation of prognostically favorable good-grade sSAH patients, (2) the predominance of MCA aneurysms, which might have a significant confounding effect on outcomes, given very different potential perforator injuries and the extent of dissection required, and (3) the selection of patients with radiologically confirmed hydrocephalus. By nature, non-hydrocephalus patients are not amenable to CSF biomarker sampling. As a consequence, our data is not applicable to sSAH patients in general.

Multiple previous SAH investigations have used the SF-36, even in poor-grade SAH, showing an impact on all tested items of the SF-36 (cf. references in [12]). Both of our utilized measurement tools, the SF-36 and the ISR, were completed within 10 to 20 min and no effects of fatigue were apparent during the testing, neither at t1 nor at t2. However, we stress that in patients with a central nervous system disease, a confounding influence of fatigue [86] and/or cognitive impairment [88] cannot conclusively be excluded. Further research is necessary to develop assessments sensitive for the specific pattern of deficits in patients with intracranial aneurysms [5, 89]. Since statistically relevant correlations only occurred in the short term, a false-positive result has to be considered. Experiencing a traumatic and life-threatening event like the acute phase of an sSAH with consecutive ICU treatment expectedly bears the risk of low psychological and hrQoL scores. In concert with the supposed widespread derangement of neurotransmitters, CGRP, being part of an acute neurobiological response, is unsurprisingly high during this period. Conclusively, the established positive correlations have to be interpreted with caution. Our observational, correlative clinical pilot study prevents drawing final conclusions, establishing clear associations or implying causality.

Yet, our prospective, controlled study provides academically valuable data that offer new insight into the plausible interactions between CGRP in supratentorial CSF and psychopathology after good-grade sSAH as well as into the (to date still underreported) time course of hrQoL performance in the early stages of recovery. Standardized self-reported outcome measures are an important facet of a comprehensive hrQoL outcome assessment, the more since cognitive domain deficits are further complicated by reduced hrQoL, depression, anxiety, and sleep disturbances.

Conclusion

Our study reveals the first insight into the potential capacity of endogenous CGRP as a predictive psychoactive biomarker in the ventricular CSF subacutely after the onset of sSAH and its potential contribution to neurobehavioral impairment and reduced hrQoL. In line with preclinical data and the psychiatric literature, the present data suggests that, after sSAH, increased CSF CGRP concentrations significantly adversely affect short-term psychological and physical health with respect to depression, anxiety, somatoform syndrome, compulsive-obessive syndrome, the supplementary ISR items, general health perceptions, and the SF-36 physical component summary score. Of special note is the potential therapeutic dilemma as, on the one hand, it is highly conceivable that the psycho- and vasoactive neuropeptide CGRP is—at least in part—involved in the pathogenesis of reduced hrQoL after good-grade sSAH, whereas on the other hand, CGRP was certified a cerebroprotective role by counteracting sSAH-induced vasoconstriction and CV-related cerebral ischemia. Our interesting results justify further research on endogenous CGRP in CSF and plasma with a focus on the influencing
factors of its release, the temporary dynamics, and the pathophysiological interactions with higher integrated neurobehavior.

**Authors’ contributions** Authors’ contributions to the study and manuscript preparation include the following: Conception and design: Schebesch, Karl-Michael. Acquisition of data: Bründl, Elisabeth, Schödel, Petra, Bele, Sylvia, Höhne, Julius, Martin Proescholdt, Störr, Eva-Maria. Analysis and interpretation of data: Bründl, Schebesch, Proescholdt, Zeman, Florian. Drafting the article: Bründl. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Schebesch. Statistical analysis: Zeman, Proescholdt, Bründl, Schebesch. Study supervision: Schebesch, Bründl.

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**Data availability** The data that support the findings of this study are available from the corresponding author upon reasonable request.

**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

**Ethical approval** All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

**Informed consent** Informed consent was obtained from all individual participants included in the study.

**Health and safety** The authors confirm that all mandatory laboratory health and safety procedures have been complied with in the course of conducting any experimental work reported.

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Highlights

- Impairment in health-related quality of life (hrQoL) after treatment of spontaneous subarachnoid hemorrhage (sSAH) is common but underreported.
- Calcitonin gene-related peptide (CGRP) is a potent cerebroprotective vasodilator and psychoactive mediator.
- This study is the first to correlate endogenous CGRP with hrQoL outcome in good-grade sSAH.
- Excessive CGRP release into cerebrospinal fluid (CSF) may have a negative impact on hrQoL, anxiety, and depression.
- CSF CGRP is suggested to be involved in the pathogenesis of impaired higher integrated behavior after sSAH.

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