Cognitive dysfunction occurs frequently in patients with heart failure (HF), but early detection remains challenging. Serum glial fibrillary acidic protein (GFAP) is an emerging biomarker of cognitive decline in disorders of primary neurodegeneration such as Alzheimer’s disease. We evaluated the utility of serum GFAP as a biomarker for cognitive dysfunction and structural brain damage in patients with stable chronic HF.

Methods and results
Using bead-based single molecule immunoassays, we quantified serum levels of GFAP in patients with HF participating in the prospective Cognition.Matters-HF study. Participants were extensively phenotyped, including cognitive testing of five separate domains and magnetic resonance imaging (MRI) of the brain. Univariable and multivariable models, also accounting for multiple testing, were run. One hundred and forty-six chronic HF patients with a mean age of 63.8 ± 10.8 years were included (15.1% women). Serum GFAP levels (median 246 pg/mL, quartiles 165, 384 pg/mL; range 66 to 1512 pg/mL) did not differ between sexes. In the multivariable adjusted model, independent predictors of GFAP levels were age ($T = 5.5; P < 0.001$), smoking ($T = 3.2; P = 0.002$), estimated glomerular filtration rate ($T = −4.7; P < 0.001$), alanine aminotransferase ($T = −2.1; P = 0.036$), and the left atrial end-systolic volume index ($T = 3.4; P = 0.004$). NT-proBNP but not serum GFAP explained global cerebral atrophy beyond ageing. However, serum GFAP levels were associated with the cognitive domain visual/verbal memory ($T = −3.0; P = 0.003$) along with focal hippocampal atrophy ($T = 2.3; P = 0.025$).

Conclusions
Serum GFAP levels are affected by age, smoking, and surrogates of the severity of HF. The association of GFAP with memory dysfunction suggests that astroglial pathologies, which evade detection by conventional MRI, may contribute to memory loss beyond ageing in patients with chronic HF.

Keywords
Glial fibrillary acidic protein; GFAP; Chronic heart failure; Cognitive decline; Memory dysfunction; Brain atrophy

Introduction
Heart failure (HF) has become one of the most prevalent chronic diseases in industrialized countries. The extent and severity of secondary organ affections significantly contribute to HF-related health care costs and long-term outcome. About every second patient with HF exhibits cognitive deficits. Within the last years, there has been increasing interest in serum markers related to cognitive deficits and predicting cognitive decline. The intermediate filament glial fibrillary acidic protein (GFAP) holds promise in this regard. It is expressed mainly in mature astrocytes and its up-regulation is a hallmark of disease-related reactive astrogliosis in the central nervous system. A growing body of evidence suggests that blood GFAP levels can be used to detect even subtle injury to the central nervous system. In a recent
population-based sample \((n = 1327)\), a diagnosis of clinical Alzheimer’s disease was 62% or 149% more likely, if GFAP levels were in the third quartile \((>232 \text{ pg/mL})\) or the fourth quartile \((>337 \text{ pg/mL})\), compared with lower serum levels of GFAP within the first quartile \((<160 \text{ pg/mL})\). Accordingly, serum GFAP above median predicted a 130% faster cognitive decline over time compared with those in the lowest quartile. Serum GFAP has been studied in numerous neurological disorders such as Alzheimer’s disease,\(^9,10\) frontotemporal dementia,\(^10\) and neuromyelitis optica,\(^11\) but its utility to detect and predict secondary cognitive impairment in cardiac diseases remains unknown. While one recent study demonstrated that GFAP predicted neurological outcome after out-of-hospital cardiac arrest,\(^12\) the role of GFAP in chronic HF remains unknown.

We hypothesized that serum GFAP levels relate to cognitive function in patients with HF but may depend on comorbidities. To address these issues in a cross-sectional approach, we made use of the deep clinical phenotyping approach adopted in the context of the Cognition.Matters-HF study, which investigated patients with chronic stable HF without focal neurological impairment.

**Methods**

**Study design**

The design and applied methodology of the Cognition.Matters-HF study have been reported in detail previously.\(^{13,14}\) In brief, the Cognition.Matters-HF study was a prospective, monocentric cohort study conducted in compliance with the Declaration of Helsinki and approved by the local ethics committee (\#245/10).\(^{14}\) The study included adult patients with confirmed chronic HF according to the then-current guidelines of the European Society of Cardiology,\(^1\) whereas patients with newly diagnosed or decompensated HF were not eligible. The selection criteria are summarized in Supporting Information Table S1. Of note, patients exhibiting apparent neurological or psychiatric disease, history of clinical stroke, or carotid artery stenosis over 50% were not eligible.

**Clinical evaluation**

Physical examination, electrocardiography, echocardiography, and 6 min walk test were performed according to standard operating procedures (for details, refer to the supporting information, Supplemental Methods).\(^{13,14}\) Neurological evaluation included extensive clinical examination. Psychological testing was performed between 9 AM and 11 AM using a comprehensive test battery based on taxonomy of attention dimensions and is summarized in Table S2. T-standardized output values accounting for the modifying effect of age, gender, and educational level were reported with a mean of 50 and standard deviation of 10.

**Cerebral magnetic resonance imaging**

Brain magnetic resonance imaging (MRI) was performed at a 3-Tesla scanner (Siemens MAGNETOM Trio, Siemens Healthcare, Erlangen, Germany) as described previously.\(^4\) Briefly, the applied MRI protocol enabled the estimation of global and regional measures of brain structure degeneration by visual rating (Table S3). Visual rating of cerebral atrophy ranged on a scale from 1 to 8. For the assessment of medial temporal lobe atrophy, Scheltens score ranging from 0 (normal) to 4 (severe atrophy) was applied, and the mean scores of both sides (left and right) were reported.

**Laboratory analysis**

We collected non-fasting venous blood samples from all patients for routine clinical chemistry investigations at the certified facility of the University Hospital Würzburg. Participants were positioned seated for at least 5 min before puncture. Serum samples were processed immediately, that is, remained at room temperature for 30 min and were centrifuged for 10 min at 2000g. Serum was aliquoted in dedicated fluid tissue tubes (Micronic, Lelystad, Netherlands) and stored in the standardized interdisciplinary biomaterial bank at \(-80^\circ\text{C}\) until analysis.\(^{15}\) Serum GFAP was measured using the Simoa GFAP-kit (102336; Quanterix\(^{\text{TM}}\), Billerica, MA, USA) on a Simoa HD-1 analyser instrument (Quanterix\(^{\text{TM}}\), Billerica, MA, USA) according to the manufacturer’s instructions. These measurements were performed blinded to the patients’ other results at the Department for Neurology of the University Hospital Ulm.\(^{16}\)

**Data analysis**

Statistical analysis was performed using the statistical software SPSS (Version 26). To test for normal distribution, we used the Shapiro–Wilk test. Variables were natural log normalized if required. The few missing values (less than 1% missing) were imputed by the mean value of respective variable. For nominal and ordinal data, \(\chi^2\) test, or Fisher’s exact test were used, according to the nature of the data. For correlation analysis, the Spearman rho coefficient \((\rho)\) was computed. To identify metric correlates of GFAP levels, univariate linear regression analysis was used, and a trend test across quartiles was reported. All tests were performed two-sided.

When identifying determinants of GFAP, to reduce the optimism introduced by multiple testing, a Bonferroni correction of \(P\) values was used for the 87 clinical parameters that were used for analysis. Thus, identified correlates were then included into a multivariable regression model. First, we analysed correlates using an ‘enter’ approach. We reduced

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multicollinearity by showing variance inflation factors of < 10 in each model including Durbin–Watson statistics. Second, we fed the remaining correlates into another regression model, which then was reduced to the final model through backward elimination, and reproduced by forward entry again. The explained variance of a model was indicated by the coefficients of determination ($R^2$).

**Results**

**Patient characteristics**

Because serum for GFAP determination was not available in 2 out of the 148 patients of the Cognition.Matters-HF cohort, all analyses refer to 146 patients. Their mean age was 63.8 (standard deviation 10.8) years, and 15.1% were female. The characteristics of the study sample are shown in Tables **1** and **2** and have partially been published before.\(^1\) Mean left ventricular ejection fraction was 43 (8) %, and 72% of patients were in New York Heart Association functional class II or III. Coronary artery disease was the predominant cause for HF in 65%, and 84% of all patients received optimal HF pharmacotherapy according to established guidelines at study inclusion.\(^3\) Serum concentrations of GFAP ranged from 66 to 1512 pg/mL, with mean of 297 (190) pg/mL and median of 246 pg/mL. For histograms, refer to Figure **S1**.

**Clinical correlates of glial fibrillary acidic protein levels**

To identify clinical correlates of GFAP, we defined four equally sized patient groups according to the quartiles 165, 246, and 384 pg/mL (**Table** **1**). After correction for multiple testing, we found positive associations of GFAP with age, smoking (ever), NT-proBNP, urea, and left atrial volume index (LAVI), estimated glomerular filtration rate (eGFR), and alanine aminotransferase (ALAT). In line, thus identified correlates significantly related to ln (GFAP) in Spearman’s correlation (**Figure** **2**) and univariable regression analysis (**Table** **2**). Although there were minor differences in unadjusted GFAP levels depending on the type of HF-related medication and drugs for important comorbidities (**Table** **S2**), none of these substance classes emerged as a major predictor in multivariable analysis (**Table** **2**).

**Determinants of glial fibrillary acidic protein in multivariable analyses**

To compare the relative effect of the above identified variables, we performed multivariable regression (**Table** **2**). Both in the ‘enter’ and the ‘backward elimination’ approach, five independent predictors of ln (GFAP) emerged, hereafter sorted by their relative weight in the model: age, eGFR, smoking status (ever), LAVI, and ALAT. Those models exhibited a $R^2$ of 0.55 as an indicator for the explained variance. These associations yield the following approximation formula for GFAP levels (ever smoking = 0; never smoking = 1):

$$GFAP \left( \frac{pg}{mL} \right) = 87.4 \ast 1.019^{age} \ast 0.991^{eGFR} \ast 1.238^{smoking status} \ast 1.0059^{LAVI} \ast 0.9948^{ALAT}$$

**Gliarial fibrillary acidic protein levels and brain morphologic alterations**

As displayed in **Table 1** and **Figure 2**, serum levels of GFAP were associated with cerebral atrophy score. When GFAP was added to age-adjusted multivariable models of the above mentioned parameters, the explained variance ($R^2$) of cerebral atrophy was not increased (**Table** **3**). In this model, NT-proBNP, but not GFAP, significantly predicted global brain atrophy beyond ageing.

**Gliarial fibrillary acidic protein levels and cognitive functioning**

Aiming to evaluate the value of GFAP as a biomarker of cognitive decline, we found that out of the five tested cognitive domains, only visual/verbal memory was significantly associated with GFAP after correction for multiple testing (**Table** **1**). In line, serum GFAP showed high (negative) correlations coefficients ($\rho$) regarding memory $T$ values. Of note, the $T$ values of cognitive domains are already adjusted for age, and partially for gender and educational level (**Figure** **2**).

Using multivariable models, we addressed, whether GFAP could improve models of visual/verbal memory (**Table** **4**). We found that brain MRI parameters were related to memory impairment, for example, the hippocampal atrophy score. When GFAP was added to these models, the explained variance was increased. There, GFAP emerged as an independent predictor of memory dysfunction along with hippocampal atrophy.

**Discussion**

The current study provides for the first time a detailed characterization of serum GFAP levels in extensively phenotyped patients free from stroke or other focal neurological deficits yet suffering from chronic HF.\(^4\) In this cohort, 68% of participants exhibited cognitive impairment. We quantified serum levels of GFAP to explore, whether it has a mediating role be-
Table 1  Descriptive clinical characteristics of study participants according to serum glial fibrillary acidic protein (GFAP) quartiles

| Table 1: Descriptive clinical characteristics of study participants according to serum glial fibrillary acidic protein (GFAP) quartiles |
|---------------------------------|
| **n**  | **Total** | **<165 pg/mL (n = 36)** | **165 to 244.9 pg/mL (n = 36)** | **245 to 383.9 pg/mL (n = 38)** | **≥384 pg/mL (n = 36)** | **T** | **P** | **P(adjusted)** |
|------|----------|------------------------|------------------|---------------------|---------------------|------|------|-----------------|
| **Age (years)** | 146 | 63.8 (10.8) | 53.6 (8.7) | 63.2 (9.7) | 67.6 (8.1) | 70.4 (8.8) | 8.26 | <0.001 | <0.001 |
| **Female sex** | 146 | 22 (15.1%) | 3 (8.3%) | 8 (22.2%) | 3 (7.9%) | 8 (22.2%) | 0.128 | ns |
| **Body mass index (kg/m²)** | 145 | 29.1 (5.2) | 30.5 (7.5) | 29.8 (4.1) | 28.5 (4.5) | 27.8 (3.6) | -2.50 | 0.014 | ns |
| **Parameters of heart failure (HF)** | | | | | | | |
| **Duration of HF (years)** | 145 | 6.31 (6.03) | 4.6 (4.9) | 6.5 (6.3) | 6.1 (6.3) | 8.1 (6.3) | 2.25 | 0.026 | ns |
| **Previous hospitalizations due to HF** | 146 | 0.6 (0.9) | 0.7 (1.0) | 0.5 (0.7) | 0.8 (0.9) | 0.5 (0.9) | 0.40 | 0.688 | ns |
| **New York Heart Association functional class** | | | | | | | |
| **I** | 146 | 39 (26.7%) | 13 (36.1%) | 12 (33.3%) | 7 (18.4%) | 7 (19.4%) | 8.26 | <0.001 | <0.001 |
| **II** | 146 | 88 (60.3%) | 20 (55.6%) | 20 (55.6%) | 26 (68.4%) | 22 (61.1%) | 0.128 | ns |
| **III** | 146 | 19 (13.0%) | 3 (8.3%) | 4 (11.1%) | 4 (11.1%) | 7 (19.4%) | 0.422 | ns |
| **6 min walk test distance (m)** | 138 | 391.5 (99.3) | 425.1 (94.0) | 376.5 (91.8) | 393.1 (120.4) | 369.7 (79.9) | 2.00 | 0.048 | ns |
| **Pre-existent conditions** | | | | | | | |
| **Atrial fibrillation/atrial flutter** | 146 | 33 (22.6%) | 5 (13.9%) | 7 (19.4%) | 10 (26.3%) | 11 (30.6%) | 0.338 | ns |
| **Coronary artery disease** | 146 | 100 (68.5%) | 22 (61.1%) | 30 (83.3%) | 26 (68.4%) | 22 (61.1%) | 0.139 | ns |
| **History of myocardial infarction** | 146 | 80 (54.8%) | 20 (55.6%) | 27 (75.0%) | 28 (73.7%) | 25 (69.4%) | 0.026 | ns |
| **Cardiovascular risk factors** | | | | | | | |
| **Diabetes mellitus type IIa** | 146 | 42 (28.8%) | 9 (25.0%) | 15 (41.7%) | 12 (31.6%) | 6 (16.7%) | 0.338 | ns |
| **Arterial hypertensionb** | 146 | 116 (79.5%) | 22 (61.1%) | 31 (86.1%) | 30 (78.9%) | 33 (91.7%) | 0.009 | ns |
| **Hyperlipidaemicc** | 146 | 105 (71.9%) | 25 (69.4%) | 27 (75.0%) | 28 (73.7%) | 25 (69.4%) | 0.931 | ns |
| **(Former) smoking** | 146 | 88 (60.3%) | 32 (88.9%) | 21 (58.3%) | 22 (57.9%) | 13 (36.1%) | <0.001 | 0.008 |
| **Laboratory results** | | | | | | | |
| **Alanine aminotransferase (U/L)** | 145 | 28.1 (14.1) | 34.6 (16.3) | 31.5 (17.7) | 23.4 (8.3) | 23.3 (8.2) | 4.25 | <0.001 | <0.001 |
| **Haemoglobin (g/dL)** | 145 | 14.3 (1.4) | 15.0 (1.2) | 14.2 (1.3) | 14.3 (1.3) | 13.9 (1.7) | 3.05 | 0.003 | ns |
| **Urea (mg/dL)** | 146 | 42.6 (16.4) | 43.9 (10.7) | 39.2 (11.7) | 45.0 (17.3) | 51.1 (19.7) | 4.79 | <0.001 | <0.001 |
| **N terminal pro brain natriuretic peptide (pg/mL)** | 146 | 1330 (2041) | 497 (438) | 1.084 (1.587) | 1.220 (1.744) | 2.492 (3.002) | 4.14 | <0.001 | 0.005 |
| **Estimated GFR (mL/min/1.73 m²)** | 140 | 42.0 (17.5) | 35.4 (13.2) | 39.2 (15.5) | 40.2 (14.3) | 52.1 (20.4) | 4.21 | <0.001 | 0.004 |
| **Cognitive domains** | | | | | | | |
| **Intensity of attention (T score)** | 145 | 41.8 (7.5) | 43.2 (6.3) | 40.8 (7.8) | 39.9 (7.9) | 43.5 (7.6) | 0.02 | 0.983 | ns |
| **Selectivity of attention (T score)** | 145 | 45.3 (6.2) | 47.7 (4.3) | 46.4 (6.0) | 42.7 (6.8) | 44.4 (6.3) | -2.05 | 0.014 | ns |
| **Visual/verbal memory (T score)** | 145 | 45.3 (7.9) | 48.4 (7.7) | 44.4 (9.0) | 45.9 (8.0) | 46.3 (9.2) | -0.74 | 0.462 | ns |
| **Hippocampal atrophy/Scel1ens score (0–4)** | 146 | 44.8 (7.3) | 45.4 (7.3) | 44.5 (7.2) | 44.6 (7.3) | 44.9 (7.1) | -0.24 | 0.813 | ns |

E, early diastolic mitral valve inflow velocity; e', early diastolic mitral annular relaxation velocity; GFR, glomerular filtration rate (MDRD formula); n, valid values; P, two-sided P value; P(adjusted), adjusted P value after Bonferroni correction.

Metric data are displayed as mean (standard deviation), for detection of trends, univariable regression with quartiles as independent variable was applied. Chi-squared test was used to find differences between expected and observed frequencies; if the expected value was below five, we used Fischer’s exact test.

aHistory of diabetes mellitus type II or HbA1c > 6.5%.

bSitting blood pressure > 140/90 mmHg or history of hypertension before onset of heart failure.

cHyperlipidaemia or statin treatment.

Serum GFAP indicates memory impairment in chronic HF patients.
between HF, cognitive impairment, and associated changes in brain morphology. Three key findings emerged: first, we revealed that serum GFAP levels were independently affected by age, smoking, and surrogates of the severity of chronic HF. Second, GFAP correlated to global brain atrophy, but not beyond ageing. Third, higher serum GFAP levels indepen-

Table 2 Univariable and multivariable correlates of ln (GFAP)

|                  | Univariable analysis | Multivariable analysis—enter approach | Multivariable analysis—backward elimination |
|------------------|----------------------|---------------------------------------|---------------------------------------------|
|                  | n                    | T | P          | R² | VIF | T | P      | R² | VIF | T | P      |
| Age (years)      | 146                  | 9.38 | <0.001    | 1.41 | 5.51 | <0.001 | 1.40 | 5.50 | <0.001 |
| (Former) smoking (y/n) | 146 | 3.96 | <0.001    | 1.06 | 3.10 | 0.002 | 1.05 | 3.17 | 0.002 |
| Left atrial volume index (mL/m²) | 143 | 4.36 | <0.001    | 1.22 | 2.75 | 0.007 | 1.07 | 2.95 | 0.004 |
| Alanine aminotransferase (U/L) | 146 | -4.65 | <0.001  | 1.21 | -2.10 | 0.038 | 1.16 | -2.12 | 0.036 |
| Urea (mg/dL)    | 146                  | 4.91 | <0.001    | 1.87 | -0.67 | 0.506 |
| NT-proBNP (pg/mL) | 140 | 3.70 | <0.001    | 1.34 | 0.35 | 0.725 |
| Estimated GFR (mL/min/1.73 m²) | 146 | -7.47 | <0.001  | 2.01 | -3.94 | <0.001 |

The T value indicates the direction of association and the relative weight of a variable in a model; R² indicates the variance explained by the model.

DWS, Durbin–Watson statistics; GFR, glomerular filtration rate (MDRD formula); P = two-sided P value; R², coefficient of determination; VIF, variance inflation factor.
dently predicted worse performance in the cognitive domain of visual/verbal memory.

Serum levels of glial fibrillary acidic protein in heart failure patients compared with neurological diseases

Median GFAP was 246 pg/mL in our sample, which is higher than in healthy adults\(^6\) and within the pathological range of neurological disorders such as neuromyelitis optica and Alzheimer’s disease. Neuromyelitis optica is caused by antibodies against aquaporin-4, which destroy astrocytes. In this condition, median GFAP levels in three previous reports were 208, 109, and 274 pg/mL compared with 97, 68, and 61 pg/mL in healthy controls.\(^{11,17,18}\) Patients with Alzheimer’s disease, exhibiting a more complex brain pathology, displayed median serum GFAP levels of 376 pg/mL, compared with 157 pg/mL in controls.\(^8\) The relatively high GFAP concentration in chronic HF suggests ongoing glial damage, which might be triggered by interleukin 6-induced neuroinflammation\(^{19}\) and/or neuronal and glial cell destruc-

Table 3 Univariable and multivariable correlates of cerebral atrophy

|                     | Univariable analysis | Multivariable analysis—enter approach | Multivariable analysis—backward elimination |
|---------------------|----------------------|---------------------------------------|-------------------------------------------|
|                     | n        | T      | P   | VIF | T      | P   | VIF | T      | P   | DWS = 1.576 | DWS = 1.552 |
| Age (years)         | 146      | 7.99   | <0.001 | 1.34 | 7.27 | <0.001 | 1.04 | 7.49 | <0.001 |
| NT-proBNP (pg/mL)   | 146      | 1.08   | 3.48 | 0.001 | 1.04 | 3.26 | 0.001 |
| +GFAP (pg/mL)       | 146      | 1.39   | −1.40 | 0.165 | 1.04 | 3.26 | 0.001 |

The \(T\) value indicates the direction of association and the relative weight of a variable in a model; \(R^2\) indicates the variance explained by the model.

DWS, Durbin–Watson statistics; \(P\), two-sided \(P\) value; \(R^2\), coefficient of determination; VIF, variance inflation factor.

Table 4 Univariable and multivariable correlates of age-adjusted visual/verbal memory \(T\) score

|                     | Univariable analysis | Multivariable analysis—enter approach | Multivariable analysis—backward elimination |
|---------------------|----------------------|---------------------------------------|-------------------------------------------|
|                     | n        | T      | P   | VIF | T      | P   | VIF | T      | P   | DWS = 2.249 | DWS = 2.254 |
| White matter hyperintensity volume (mm\(^3\)) | 146      | −1.04  | 0.302 | 1.09 | −0.12 | 0.908 | 1.03 | −2.26 | 0.025 |
| Cerebral atrophy score (0–8) | 146      | −1.59  | 0.115 | 1.20 | −0.18 | 0.860 | 1.03 | −3.02 | 0.003 |
| Hippocampal atrophy/Scheltens score (0–4) | 146      | −2.73  | 0.007 | 1.14 | −2.07 | 0.041 | 1.03 | −2.26 | 0.025 |
| +GFAP (pg/mL)       | 146      | −3.40  | 0.001 | 1.09 | −2.86 | 0.005 | 1.03 | −3.02 | 0.003 |

The \(T\) value indicates the direction of association and the relative weight of a variable in a model; \(R^2\) indicates the variance explained by the model.

DWS, Durbin–Watson statistics; \(P\), two-sided \(P\) value; \(R^2\), coefficient of determination; VIF, variance inflation factor.
tion upon hypoperfusion and hypoxia. Future studies are needed to advance these hypotheses and to disclose how GFAP gets access to the systemic circulation in the absence of an overt blood–brain barrier disruption as it is found in neuromyelitis optica. Because astrocyte feet are an integral component of the blood brain barrier, it is conceivable that astrocytic damage in HF patients concomitantly induces subtle leakage of the blood–brain barrier not strong enough to cause extravasation of the MRI contrast agent gadolinium-diethylenetriaminepentaacetate for detection but sufficient for release of GFAP.

**Effect of severity of heart failure**

To explain high concentrations of GFAP in HF comparable with those in severe neurological disorders, we analysed cardio-vascular correlates of GFAP. A higher symptom burden of HF (New York Heart Association functional class) and lesser physical performance (i.e. 6 min walk test distance) were not associated to GFAP. However, LAVI, which is an established echocardiographic parameter of diastolic dysfunction, was positively related to GFAP levels in our cohort, irrespective of the presence of atrial fibrillation. Increased LAVI has been shown to be independently associated with cognitive impairment and elevated brain amyloid in patients with sinus rhythm, which was even further aggravated by age. Other studies consistently reported an association between parameters of diastolic function and cognition. While causal factors like congestion, hypoperfusion, and HF-related chronic inflammation are being discussed, the cause-and-effect association of this potentially bilateral interaction as well as its prognostic utility must be detailed in future studies. Consistently, in univariable regression, serum levels of NT-proBNP, an indicator for left ventricular and left atrial stress, were positively correlated to GFAP. Higher NT-proBNP concentrations were also associated with incident cognitive impairment in a previous study independent of atherogenic and Alzheimer’s disease risk factors and in a study of the general population with mild cognitive impairment. Of note, we found no associations to GFAP for parameters of systolic dysfunction like the left ventricular ejection fraction.

**Effect of age and comorbidities**

In this investigation, patient age positively related to serum GFAP levels, which is in accordance with recent findings in depressive disorders, possible reflecting ‘normal’ neurodegenerative glial damage upon ageing. A recent study investigating human brains found age-related decline of synaptic transmission and increased expression of GFAP in both sexes. In mice, an age-related increase in GFAP RNA has been reported, potentially reflecting an increase in the size, number, and/or fibrous character of astrocytes. We additionally found higher GFAP levels in patients with impaired kidney and liver function, which in our cohort may be explained by a HF-induced typical cardio-hepato-renal compromise. These findings have important implications for the interpretation of serum GFAP in neurological disorders by pointing to age and renal and hepatic function as major factors confounding its measurement results.

**Relation of glial fibrillary acidic protein to brain morphology and cognitive impairment in heart failure**

In this cohort, single GFAP measurements did not add explained variance to models of brain atrophy. On the one hand, this might be due to the long-term emergence of these brain alterations. On the other hand, while GFAP reflects glial damage, neuronal biomarkers like neurofilament light chain are indeed related to brain degeneration in elderly. We were surprised that GFAP did not relate to white matter hyperintensity volume, as these lesions display areas of reduced glial density. As GFAP is a marker of glial damage with a short half-life, it might not be suitable to mark slowly progressive cerebral small vessel disease underlying white matter hyperintensities. Thus, future longitudinal assessment of GFAP and brain morphologic alterations might have the potential to increase the relevance of serum GFAP in this regard.

In this sample of HF patients, the glial biomarker GFAP independently predicted age-adjusted performance in the cognitive domain visual/verbal memory along with hippocampal atrophy. As previously published, cognitive deficits in the Cognition.Matters-HF cohort were associated with regional brain atrophy of the medial temporal lobe. While hippocampal atrophy, measured by Scheltens score, is known to foster memory deficits, our findings suggest that in chronic HF, memory impairment might partially also be due to ongoing glial activation and damage within the hippocampus, which evades conventional MRI. Accordingly, histological analysis revealed a decreased neurogenesis together with an increased number of reactive astrocytes in the ventral hippocampus in HF rats compared with sham rats. While similar correlations of GFAP to cognitive decline were found in Alzheimer’s disease, we here expand these finding to a cardiologically diseased cohort free of focal neurological deficits. This has potential implications for early identification and stratification of HF patients at risk for cognitive decline.

**Limitations and conclusion**

Limitations of the current study include its cross-sectional approach. Future studies have to confirm the utility of monitoring GFAP in order to predict changes in cognitive function in
HF over time. Strengths of this investigation are the extensive neurological and cardiological work-up applied, the comprehensive cognitive test battery including five different domains instead of global tests, and the detailed clinical and laboratory phenotyping.

In summary, this detailed evaluation of GFAP from sera of patients with chronic HF support its potential as biomarker for the detection of cognitive deficits in patients with HF, which are related to cardiac dysfunction. Furthermore, the associations of serum GFAP levels in our cohort indicate the important role of glial damage in cardiological disorders. When GFAP is determined in the context of neurological disorders, future analyses have to put more weight on the confounding effect of age and renal and cardiac function to avoid false conclusions, especially in elderly populations at risk for Alzheimer’s disease and other dementias.

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Conflict of interest

None to declare.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Table S1. Imaging protocol, sequence parameters and specifications.
Table S2. Extended descriptive clinical characteristics of study participants according to serum glial fibrillary acidic protein (GFAP) quartiles.
Table S3. Spearman’s correlation (ρ) to cognitive function and brain morphology.

Figure S1. Serum GFAP concentrations. Histograms and quantil-quantil (Q-Q) plots of absolute and ln-transformed GFAP measurements in the serum of chronic heart failure patients via ultrasensitive, bead-based single molecular immunosassay. Dots represent single patients (n=146).
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