**Disarranged neuroplastin environment upon aging and chronic stress recovery in female Sprague Dawley rats**

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**Abstract**

Chronic stress produces long-term metabolic changes throughout the superfamily of nuclear receptors, potentially causing various pathologies. Sex hormones modulate the stress response and generate a sex-specific age-dependent metabolic imprint, especially distinct in the reproductive senescence of females. We monitored chronic stress recovery in two age groups of female Sprague Dawley rats to determine whether stress and/or aging structurally changed the glycolipid microenvironment, a milieu playing an important role in cognitive functions. Old females experienced memory impairment even at basal conditions, which was additionally amplified by stress. On the other hand, the memory of young females was not disrupted. Stress recovery was followed by a microglial decrease and an increase in astrocyte count in the hippocampal immune system. Since dysfunction of the brain immune system could contribute to disturbed synaptogenesis, we analyzed neuroplastin expression and the lipid environment. Neuroplastin microenvironments were explored by analyzing immunofluorescent stainings using a newly developed Python script method. Stress reorganized glycolipid microenvironment in the Cornu Ammonis 1 (CA1) and dentate gyrus (DG) hippocampal regions of old females but in a very different fashion, thus affecting neuroplasticity. The postulation of four possible neuroplastin environments pointed to the GD1a ganglioside enrichment during reproductive

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**Abbreviations**: AD, Alzheimer’s disease; AMPA-R, α-amino-3-hydroxy-5-methyl-4-isoaxolepropionic acid receptor; APP, amyloid precursor protein; Aβ, amyloid beta; BACE1, beta-secretase 1; CA1, Cornu Ammonis subfield 1; CA2, Cornu Ammonis subfield 2; DG, dentate gyrus; GFAP, glial fibrillary acidic protein; HPA, hypothalamic-pituitary-adrenal axis; Iba1, ionized calcium binding adaptor molecule 1; IDV, integrated density value; Np65, neuroplastin.

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An increasing body of data has demonstrated that chronic stress may underlie various events leading to cellular senescence and cognitive impairment (Russian et al., 2018; Kim & Diamond, 2002; Kuipers et al., 2003; Lyons & Bartolomucci, 2020; Sandi, 2004; Zhang et al., 2019), possibly contributing to Alzheimer’s disease (AD) development. This hypothesis on the contribution of chronic stress to AD development is supported by the following observations: (a) preferential glucocorticoid receptor accumulates in the brain areas that are most damaged in AD (Soontornniyomkij et al., 2010); (b) both of the proteins involved in amyloid beta processing, amyloid precursor protein (APP), and beta-secretase 1 (BACE1) contain a glucocorticoid response element in their sequences (Lahiri, 2004; Sambamurti et al., 2004; Li et al., 2011); and (c) prednisone therapy in AD patients accelerated cognitive deterioration (Aisen et al., 2000). Chronic stress effects in the brain are potentiated by aging through a decrease in the number of neurons, brain volume shrinkage, neuronal networks changes, circadian rhythms disruption, and lipid profile changes (Hung et al., 2010; Williamson & Lyons, 2018). On the other hand, stress response is attenuated by estrogen; hence, estrogen deficiency causes stress-induced damage during aging. This damage is particularly evident in women, suggesting the existence of a sex-specific mechanism of neurodegeneration (Dumas et al., 2012; Falcioni et al., 2016; Marin & Diaz, 2018; Zárate et al., 2017). In female Sprague Dawley rats, reproductive senescence starts at 8–10 months of age and is characterized by a decreased number of developing follicles and changes in estrous cyclicity (Cruz et al., 2017).

Another crucial factor contributing to brain preservation is related to cerebral lipid composition and homeostasis. Glycosphingolipids, especially gangliosides (sialic acid-containing glycosphingolipids) maintain the structure, function, and transport to target membranes of at least 10% of transmembrane proteins (Simons & Toomre, 2000). Ganglioside species are specifically abundant in the synaptic membranes (Blennow et al., 1991), a fact suggesting their involvement in synaptic processes. The four most common gangliosides in the adult brain are GD1a, GD1b, GT1b, and GM1 (Vajn et al., 2013).

Some of the proteins essential for synaptic plasticity regulation and memory formation are located specifically in lipid rafts (Beesley et al., 2014; Marin & Diaz, 2018; Prendergast et al., 2014). Neuroplastin is a raft-dependent protein belonging to the immunoglobulin superfamily. It is highly expressed in the hippocampus, where it regulates synaptic plasticity. The stability of this protein and its functioning within the neuronal membrane have been associated with GM1-enriched lipid rafts (Beesley et al., 2014; Marzban et al., 2003). Neuroplastin contributes to neurite growth, maintenance of excitatory, and inhibitory synapse ratio and mechanisms of long-term potentiation (Owczarek et al., 2011; Smalla et al., 2000). Neuroplastin knock-out mice (Np/- mice) have a significantly disturbed hypothalamic-pituitary-adrenal (HPA) axis, as reflected in the increased concentration of serum corticosterone under basal conditions. Interestingly, they also display decreased locomotion and contextual memory (Beesley et al., 2014; Bhattacharya et al., 2017). These physiological anomalies have been shown to be sex-related, as males have a doubled corticosterone increase compared with females. Another molecule associated with lipid rafts is AMPA-R which mediates most of the excitatory synaptic transmission (Carter et al., 2004; Chang et al., 2006; Zhang et al., 2018). AMPA-R-mediated excitatory synaptic transmission in the hippocampus is decreased by chronic stress (Carter et al., 2004; Chang et al., 2006; Kallarackal et al., 2013; Zhang et al., 2018), while AMPA-R exocytosis and expression on the cell surface are decreased by the disruption of lipid rafts (Hou et al., 2008; Prendergast et al., 2014).

Aging significantly changes lipid raft matrix stability, a process associated with the development of AD (Díaz et al., 2018; Mesa-Herrera et al., 2019). GM1 ganglioside accumulation and GD1a decrease, especially in the hippocampus, were detected in AD (Mesa-Herrera et al., 2019; Svennerholm, 1994). However, it remains unclear whether the long-term stress exposure could intensify gangliosides loss and whether this process is sex-specific.

The hypothesis of this study is that chronic stress during aging in females structurally changes the lipid microenvironment, a milieu playing an important role in cognitive functions. Since lipid microdomains affect the synaptic plasticity and cognitive processes, the impairment of lipid microenvironment upon stress may act as a basis for early senescence of stressed females, as well as its high dispersion in both regions and to GD1a and GM1 loss in the CA1 region. A specific lipid environment might influence neuroplastin functionality and underlie synaptic dysfunction triggered by a combination of aging and chronic stress.

**KEYWORDS**

GD1a, GM1, hippocampus, neurodegeneration, neuroinflammation, reproductive senescence
neurodegeneration in a sex-specific manner. Here, we explored the impact of chronic stress on memory and brain preservation after chronic stress recovery in young and old female rats. Since several factors affect luminescence intensity, pixel intensities of immunofluorescent stainings were analyzed using a newly developed Python script in order to explore neuroplastin microenvironments in situ.

2 | MATERIALS AND METHODS

2.1 | Animals

The study was performed in 2016. Forty female Sprague Dawley rats (Charles River, Milano, Italy) were divided into four groups of 10 animals. This number of animals per group ensured adequate statistical power and was in compliance with the EU Directive on Animal Welfare. The animals were assigned to two age groups using simple Excel software randomization— young (3.5 months old) and old (12 months old) females, with appropriate control groups. The experiments with young females were performed at the Faculty of Medicine, J. J. Strossmayer University of Osijek, Croatia, and those with old females were performed at the Faculty of Pharmacy, University of Szeged, Hungary. Both series of experiments were carried out with the same equipment and chemicals, so they can be considered as a single experiment. The animals were housed in self-ventilating cages (EHRET, Freiburg, Germany), with five air changes per minute, stable room temperature (21 ± 2°C) and humidity (40%–60%). Standardized rat chow (Mucedola, Milan, Italy; cat. no: 4RF21) and tap water were always provided, except when overnight fasting was used as a stressor. The day cycle was set to 7.00 a.m. to 7.00 p.m. All experiments (except lights-on or noise during the night) were conducted in the morning or early afternoon (9.00 a.m. to 2.00 p.m.). The study was approved by the Ethics Committee of the Faculty of Medicine Osijek and Croatian Ministry of Agriculture under class: 602-04/14-08/06, number: 2158-61-07-14-118 and the National Scientific Ethics Committee on Animal Experimentation, Hungary (permission number: IV./3796/2015). The 3R principles and EU directive on laboratory animal welfare were strictly followed.

2.2 | Chronic stress protocol

The chronic stress protocol consisted of three 10-day stress cycles with the application of eight stressors (overnight light or noise exposure, rotation of the animals in cages on a laboratory shaker, forced swimming in cold water, restraint in metal tubes, insulin injection, exposure to cold, and overnight fasting) followed by a 14-day recovery period between each stress cycle (Figure 1). The protocol lasted for 10 weeks. Two different stressors were applied each day to avoid stressor-specific responses, with some stressors being applied multiple times during each stress cycle. In the control groups, sham stress was performed by handling the animals in the same way as animals in experimental groups, but the stressors were not applied in order to compensate for the stress arising from standard handling. The anesthetics use was not necessary since no physical pain was caused and no surgical procedures were performed. The tissues were collected 2 weeks after the last stress cycle to allow us to determine only the extended metabolic changes.

2.3 | Passive avoidance test

Memory impairment was assessed with a step-through passive avoidance test performed 1 day after the end of the third chronic stress cycle. The apparatus (Ugo Basile, Varese, Italy) delivered an electric shock through the steel grid floor. The parameters used were as follows: door delay for 10 s, 0.5 mA, a foot-shock for 2 s, and total cut-off time of 180 s. The test lasted for 3 days. On the first day, the animals were allowed to explore the device for 180 s with no shock applied. On the second day, each rat upon entering the dark compartment

FIGURE 1 Chronic stress protocol scheme. Study included two experimental (stress) age groups (N = 10)—young (3.5 months old) and old (12 months old) female rats, with appropriate control groups (sham stress). The protocol lasted 10 weeks.
received a foot-shock. On the third day, memory acquisition was recorded in terms of the total time (seconds) necessary for the animals to enter the dark compartment.

2.4 Tissue collection

The animals were euthanized with isoflurane overdose (Forane® isoflurane, Abbott Laboratories LTD; Lake Bluff, IL, USA), which was applied in isoflurane induction chamber until loss of consciousness and combined with 30-mg/kg intramuscular ketamine injection (Ketanest, Pfizer Corporation, New York, NY, USA). The brains were isolated at the ages of 6 and 14.5 months, fixed in 4% buffered paraformaldehyde solution (pH = 7.35), cryoprotected in a series of sucrose solutions for 24 hr (10%, 20%, and 30% solutions) and cut using a cryostat (CM3050S, Leica, Nussloch, Germany) into 35-μm sagittal sections.

2.5 Single immunohistochemical staining of AMPA-R, APP, microglia, and astrocytes

Single stainings were performed as previously described (Balog et al., 2015). The following primary and secondary antibodies were used: anti-ionized calcium-binding adapter molecule 1 (Iba1) (Wako, Neuss, Germany, cat. no: 019-19741) diluted to 1:1,000, antiglial fibrillary acidic protein (GFAP) (DAKO, Glostrup, Denmark, cat. no: M0761) diluted to 1:4,000, anti-AMPA-R subunit 2 (Neuro Mab, Davis, CA, USA, cat. no: L21/32) diluted to 1:500, biotinylated antimouse (Jackson Immunoresearch Laboratories, West Grove, PA, USA, cat. no: 115-065-166), biotinilated antiggoat (Jackson Immunoresearch Laboratories, West Grove, PA, USA, cat no: 705-065-147), and biotinilated antitrabbit (Jackson Immunoresearch Laboratories, West Grove, PA, USA, cat. no: 111-035-003) IgG antibodies combined with ABC kit (Vector Laboratories, CA, USA). All secondary antibodies were diluted to 1:500. The sections were mounted on glass slides and cover-slipped. In the case of fluorescent slides, the sections were stained with DAPI diluted to a concentration of 1 μg/ml for 10 min at +4°C with appropriate secondary antibodies: AF647 antimouse and CY3 antiggoat (AF647 antimouse and CY3 antiggoat; Jackson Immunoresearch Laboratories, West Grove, PA, USA, cat. no: 715-605-151 and 705-165-147, both diluted 1:500). After rinsing, the sections were additionally incubated (1.5 hr) with cholera toxin beta subunit conjugated with FITC fluorophore (Sigma, Saint Louis, MO, USA, cat. no: C1655, diluted 1:650) aimed to target GM1 ganglioside (referred to as GM1 later in the text to simplify the explanation of the results). Cholera toxin binds selectively to GM1. Additionally, we assumed that the overabundance of GM1 among other glycans recognized by cholera toxin in the brain justifies its use as a GM1 marker (Schnaar et al., 2014). After rinsing, the nuclei were stained with DAPI diluted to a concentration of 1 μg/ml for 10 min at +4°C and rinsed again in PBS. The microscope, magnification, and image resolution were the same as those used in the single fluorescent staining.

2.6 Triple immunofluorescent staining of neuroplastin, GD1α, and GM1 gangliosides

Free-floating sagittal brain sections were incubated in a blocking solution (1% BSA, 5% donkey serum, 5% goat serum) for 2 hr at +4°C followed by incubation for 48 hr at +4°C with two primary antibodies diluted in blocking solution: mouse anti-GD1α-1 (provided by prof. Schnaar. Johns Hopkins University, Baltimore, MA, USA, dilution 1:3,000) and goat antineuroplastin (Np65) (R&D Systems, Minneapolis, MN, USA, cat. no: AF5360, diluted 1:300). After four 10-min washes in PBS, the sections were incubated for 4 hr at +4°C with appropriate secondary antibodies: AF647 antimouse and CY3 antiggoat (AF647 antimouse and CY3 antiggoat; Jackson Immunoresearch Laboratories, West Grove, PA, USA, cat. no: 715-605-151 and 705-165-147, both diluted 1:500). After rinsing, the sections were additionally incubated (1.5 hr) with cholera toxin beta subunit conjugated with FITC fluorophore (Sigma, Saint Louis, MO, USA, cat. no: C1655, diluted 1:650) aimed to target GM1 ganglioside (referred to as GM1 later in the text to simplify the explanation of the results). Cholera toxin binds selectively to GM1. Additionally, we assumed that the overabundance of GM1 among other glycans recognized by cholera toxin in the brain justifies its use as a GM1 marker (Schnaar et al., 2014). After rinsing, the nuclei were stained with DAPI diluted to a concentration of 1 μg/ml for 10 min at +4°C and rinsed again in PBS. The microscope, magnification, and image resolution were the same as those used in the single fluorescent staining.

2.7 Immunohistochemical quantification and a newly developed Python script

All quantification analyses were performed by a researcher uninvolved in the experiments. The cell count of GFAP and Iba1 and the surface areas of APP-positive cells were analyzed in the CA1 and DG regions. The surface areas of AMPA-R-positive cells were analyzed in the CA1, CA2, and DG regions. All stainings were quantified with the Fiji software (Wayne Rasband, National Institutes of Health, USA). The total analyzed surface area in each hippocampal region was 0.01 mm². After adjusting the threshold according to the negative control, the integrated
density was measured. At least five biological replicates were used for microscopy. AMPA-R and APP images were analyzed with the integrated density value (IDV) plugin of the Fiji software (Schindelin et al., 2012). Fluorescent triple stainings were analyzed with a custom Python script. In the analysis of pixel intensities, a thresholding operation was used to reduce 16-bit images into binary ones. In this step, the pixel intensity value was replaced by a binary “1” or “0” if the molecule of interest was present or absent, respectively, at certain image coordinates. A threshold differentiating between pixel intensity values that should be treated as logical “1” or “0” was set for every sample and every imaging channel based on the control experiments. To calculate the colocalization between the channels, we applied logical AND operation for every pair of double or triple (GD1a/GM1/Np65) channel combinations. The resulting binary image represented the colocalized pixels, which were counted and used for further analysis as absolute values with respect to the total number of pixels that passed the thresholding. The triple stainings data were used to create pie charts illustrating the membrane disarrangement upon stress and aging. The pie charts included absolute pixel count of all channels. To illustrate the membrane composition and avoid the loss of pixel count when creating the pie chart, we applied the following formulas:

\[
Ch_1_{\text{tot}} = Ch_1_{\text{nCL}} + (dCL_1/2 - tCL) + (dCL_1/3 - tCL) + tCL,
\]

where Ch1 stands for Channel 1, tot is total pixel count, nCL are noncolocalized pixels, dCL1/2 or dCL1/3 are double colocalizations of Channel 1 and 2 or Channel 1 and 3, respectively, and tCL is triple colocalization. The same logic applies for Ch2_{tot} and Ch3_{tot}. Therefore, to obtain the absolute pixel count for certain colocalization, which will be presented by a pie chart, the following formula applies:

\[
\text{Absolute pixel count} = Ch_1_{\text{tot}} + Ch_2_{\text{tot}} + Ch_3_{\text{tot}}
\]

\[
= Ch_1_{\text{nCL}} + Ch_2_{\text{nCL}} + Ch_3_{\text{nCL}} + 2 \times (dCL_1/2 - tCL) + 2 \times (dCL_1/3 - tCL) + 2 \times (dCL_2/3 - tCL) + 3 \times tCL.
\]

The CA1 and DG hippocampal regions stained with Np65 and GM1 were 3D-reconstructed with confocal microscopy. Tissue reconstructions were described qualitatively. The colocalization of both markers in different z-sections was 3D-assembled by confocal microscopy at 40x magnification (oil immersion) using a Leica SP8 microscope and the matching LAS X Leica software.

2.8 | Statistical analysis

All statistical analyses were performed by a researcher uninvolved in the experiments. Statistical analysis of passive avoidance was performed in Python software with the Cox regression modeling and chi-square test in old females. Statistical analysis of immunostaining was performed with Statistica 12 software (TIBCO, Palo Alto, CA, USA). A p-value of <0.05 was considered significant. The two-way ANOVA was performed to determine the influence of stress, age, and their interaction on the analyzed parameters. Bonferroni correction was used for post hoc comparisons. The normality of data distribution was tested with the Shapiro–Wilk test.

3 | RESULTS

3.1 | Memory impairment

Memory impairment, tested by passive avoidance test, was present in 90% of stressed old females compared with 40% of the control group. Cox regression modeling revealed a greater hazard ratio for the effect of age. However, the effect did not reach significance, probably due to data uniformity among young animals (all young animals spent the maximum 180 s in the apparatus). Higher memory impairment in old stressed females was significant when compared to old control group of females (\( \chi^2 = 5.22, df = 1, p = 0.022 \)) (Figure 2).

3.2 | Chronic stress dysregulates immune response in the hippocampus

Microglia dynamics in response to aging and chronic stress recovery in the CA1 and DG regions was evaluated by determining the Iba1-positive cell count and IDV (Figure 3). Young and old stressed females had a reduced microglial cell number in the hippocampal regions, but no significant difference was observed. IDV in the DG followed the cell count, but in the CA1 the opposite effect was observed—while cell number decreased, IDV in both animal groups increased upon stress (CA1, age: \( F_{1,24} = 15.96, p = 0.001 \) and DG, stress: \( F_{1,24} = 12.52, p = 0.002 \); interaction of stress and aging: \( F_{1,24} = 16.25, p = 0.021 \)). We can speculate that in the CA1 region, increased IDV and decreased cell number represent the microglia activation upon stress.

Astrogial response to aging and stress was estimated by determining the GFAP-positive cell count (Figure 4). Both stressed animal groups had significantly increased astroglial cell number in the CA1 region (\( F_{1,24} = 8.67, p = 0.007 \)). However, in the DG, astroglial cell number decreased in
young stressed females and increased in old stressed females, with age and age–stress interaction reaching significance (age: $F_{1,24} = 7.99$, $p = 0.009$; interaction: $F_{1,24} = 11.26$, $p = 0.003$). The number of GFAP-positive cells significantly decreased in the old control group compared with the young control group (Bonferroni post hoc analysis $p < 0.001$). GFAP-positive cell count in the DG followed the trend of IDV (data not shown).

3.3 | Chronic stress conditions affect synaptic plasticity

AMPA-R expression in the CA1, CA2, and DG regions was assessed because of its involvement in long-term potentiation (Figure 5). Aging significantly affected AMPA-R expression in the CA2 region ($F_{1,19} = 5.034$, $p = 0.037$). Additionally, a post hoc between-group comparison revealed that age underlied AMPA-R decrease in the CA2 of old stressed females compared with young stressed females ($p = 0.006$). The interaction of aging and stress profoundly affected AMPA-R expression in both the CA2 and DG regions (CA2: $F_{1,19} = 5.083$, $p = 0.036$, DG: $F_{1,19} = 4.673$, $p = 0.044$). Stress significantly decreased AMPA-R expression in the DG region of old animals ($F_{1,19} = 5.464$, $p = 0.031$). Thus, AMPA-R expression was more profoundly reduced in the CA2 and DG than in the CA1 due to both aging and long-term consequences of stress and are reflected in memory test (Figure 5b).

3.4 | Chronic stress increases the expression of amyloid precursor protein

APP expression was analyzed by immunofluorescence in the CA1 and DG regions (Figure 6). APP expression in the CA1 increased in young stressed females, but was even higher in both groups of old females, with the main effects reaching significance (age, $F_{1,16} = 8.521$, $p = 0.01$; stress, $F_{1,16} = 89.925$, $p < 0.001$). In the DG, it was significantly affected by the interaction of stress and aging ($F_{1,16} = 5.056$, $p = 0.039$), with the increase observed only in old stressed females. Generally, the staining intensity for the DG region was lower compared with that for the CA1.

3.5 | Compromised neuroplasticity upon chronic stress recovery

We explored the colocalization of neuroplastin with GD1α and GM1 in the CA1 and DG regions. Pie charts representing the absolute pixel count for single, double, and triple combinations were used to illustrate the membrane disorganization as a result of stress and aging (Figures 7 and 8). In the CA1 region of the young female group, chronic
stress did not dramatically change the shares of the three membrane components or their colocalizations. However, in old stressed animals, the ratios of Np65 (main effect of age: $F_{1,25} = 94.052, p < 0.001$), GD1a (main effect of age: $F_{1,25} = 40.901, p < 0.001$, stress: $F_{1,25} = 38.439, p < 0.001$ and their interaction: $F_{1,25} = 39.328, p < 0.001$), and all double and triple staining combinations (main effect of stress: $F_{1,25} = 6.206, p = 0.02$) increased at the expense of GM1 (main effect of age: $F_{1,25} = 108.406, p < 0.001$ and stress: $F_{1,25} = 26.608, p < 0.001$) compared with old nonstressed females (Figure 7 and Table 1). Even though in young stressed females, free Np65 count (main effect

FIGURE 3 The quantitative and qualitative analysis of Iba1 positive cells in the CA1 and DG hippocampal regions in young and old female rats upon chronic stress. Iba1-positive staining (measured as integrated density value) is shown in (a). Microglia cell number decreased in both young and old stressed females in the CA1 and DG, but no statistical significance was reached (b). Image inserts are shown in (c). Significant main effects and their interaction are marked as follows: * $p < 0.05$, ** $p < 0.001$. Significant difference between the groups is marked by a § symbol, and the exact $P$-value is indicated. Abbreviations: C, negative control reaction; CA1, Cornu Ammonis subfield 1; DG, dentate gyrus; Iba1, ionized calcium binding adaptor molecule 1; NS, nonsignificant; OF-C, control group of old females; OF-S, stressed group of old females; YF-C, control group of young females; YF-S, stressed group of young females
FIGURE 4 The quantitative analysis of glial fibrillary acidic protein (GFAP)-positive cells in the CA1 and DG hippocampal regions of young and old female rats upon chronic stress. Cell count for the astrocyte positive cells was analyzed in two hippocampal regions, CA1 and DG (a). Image inserts are shown in (b). Statistical analysis was performed using a two-way analysis of variance (ANOVA). Bonferroni correction was used for post hoc comparisons. Significant main effects and their interaction are marked as *p < 0.05, **p < 0.001. Significant difference between groups in the DG region is marked by a § symbol, and the exact P-value is indicated. Abbreviations: C, negative control reaction; CA1, Cornu Ammonis subfield 1; DG, dentate gyrus; GFAP, glial fibrillary acidic protein; NS, nonsignificant; OF-C, control group of old females; OF-S, stressed group of old females; YF-C, control group of young females; YF-S, stressed group of young females.

of age: $F_{1,22} = 163.987, p < 0.001$, stress: $F_{1,22} = 4.923, p = 0.037$ and their interaction: $F_{1,22} = 6.590, p = 0.018$) in the DG increased, its bound form was retained in the same environment. The share of Np65 colocalization with GM1 (main effect of age: $F_{1,22} = 5.864, p = 0.024$ and interaction of age with stress: $F_{1,22} = 37.591, p < 0.001$)

FIGURE 5 AMPA-R immunoreactivity in the hippocampal CA1, CA2 and DG regions in female rats in response to chronic stress and aging. (a) AMPA-R immunoreactivity was analyzed in three hippocampal regions by free-floating immunohistochemistry. (b) Statistical analysis was performed using a two-way analysis of variance (ANOVA). Bonferroni correction was used for post hoc comparisons. Significant effects of age or stress and their interaction are marked as *p < 0.05. A significant difference between groups in the CA2 region is marked by a ‡ symbol, and the exact P-value is indicated. Abbreviations: AMPA-R, AMPA-receptor; C, negative control reaction; CA1, Cornu Ammonis subfield 1; CA2, Cornu Ammonis subfield 2; DG, dentate gyrus; NS, nonsignificant; OF-C, control group of old females; OF-S, stressed group of old females; YF-C, control group of young females; YF-S, stressed group of young females.
as well as in triple colocalization remained similar (main effect of age: $F_{1,22} = 55.026, p < 0.001$ and interaction of age with stress: $F_{1,22} = 20.542, p < 0.001$). In old control females, the share of colocalized Np65 was different than in young animals, with a significant reduction of free neuroplastin (Figure 8 and Table 2). In response to stress and aging, GM1 increased, while GD1a and neuroplastin decreased, and colocalizations remained at a similar level.

IDVs were analyzed as a measure of molecules dispersion for GD1a, Np65 and GM1 in the CA1 and DG (Figure S1).

3.6 | Tissue reconstruction

Three-dimensional tissue reconstruction showed a decrease in Np65 staining in the CA1 region in young and old stressed animals, while GM1 remained unchanged (Figure 9a). In the DG region of young control animals, Np65 intensity of staining and the colocalization of Np65 and GM1 were more obvious in old stressed females, with an increased scattered pattern (Figure 9b).

4 | DISCUSSION

In this study, aging and stress inducted a memory decline in female rats despite a recovery period after stress exposure. Stress caused GM1 and GD1a ganglioside disorganization, a phenomenon affecting the trafficking of molecules associated with synaptic plasticity, such as APP, AMPA-R, and neuroplastin. Our results showed a GM1 decrease and GD1a increased in old stressed females. APP expression increased in both analyzed regions followed by astrocyte number increase, while neuroplastin increased only in the DG. AMPA-R decreased in the CA1, CA2, and DG regions in old stressed females. These findings complement the growing literature demonstrating a deterioration of cognitive performance in stressful conditions (Conrad et al., 1999, 2017; Luine et al., 1994; Woo et al., 2018).

Chronic stress strongly affected neuroinflammation, neuroplasticity, amyloidogenesis, and neurodegeneration in older animals. Microglia activation has been reported in GM1 gangliosidosis mice, supporting the idea that alterations in glycosphingolipid content cause central nervous neuroinflammation (Belarbi et al., 2020; Jeyakumar et al., 2003). We observed significant changes in response to stress and aging in astrocyte cell count but not in microglia count. Nevertheless, a trend of microglia count reduction in stressed animals was clearly visible, which could support the role of stress in the modulation of microglial structure and function (Walker et al., 2013). We can hypothesize that in the CA1

**FIGURE 6** The quantitative analysis of APP expression in the CA1 and DG hippocampal regions of young and old stressed female rats. Image inserts are shown in (a). Immunofluorescent staining was analyzed using integrated density values for APP in the CA1 (B, on the left) and DG (b, on the right). Statistical analysis was performed using a two-way analysis of variance (ANOVA). Bonferroni correction was used for post hoc comparisons. Significant effects of age or stress and their interaction are marked as \(*p < 0.05, \***p < 0.001\). Abbreviations: APP, amyloid precursor protein; CA1, Cornu Ammonis subfield 1; DG, dentate gyrus; NS, nonsignificant; OF-C, control group of old females; OF-S, stressed group of old females; YF-C, control group of young females; YF-S, stressed group of young females.
region specifically, increased IDV and decreased cell number imply the microglia activation upon stress. In contrast, increased astrocyte number in both stressed groups in CA1 and in old females in DG suggested a glial inflammatory effect. According to the literature, reactive astrocytes can indicate glutamate homeostasis dysregulation (Palpagama et al., 2019). Moreover, this finding is in accordance with our results concerning AMPA-R–AMPA-R expression significantly decreased in old stressed animals in the CA2 and DG regions, while the number of astrocytes significantly increased in the DG region. Additionally, the expression pattern of astrocytes was followed by APP expression. The relationship between astrocyte number and APP expression observed in our study might imply an interplay of astrocytes and APP-expressing cells of the brain. Previously, astrocytes were found in a vicinity of amyloid plaques (Rossi & Volterra, 2009), suggesting their involvement in APP processing and consecutively amyloid-β formation and triggering of neurodegenerative processes.

Neuroplastin interacts with the HPA axis, directly influencing stress response and cognition (Beesley et al., 2014; Bhattacharya et al., 2017). It has been hypothesized that neuroplastin depends on the lipid environment (Ilic et al., 2019), so we analyzed it together with GM1 and GD1a gangliosides. The two gangliosides were studied together because GD1a acts as a reservoir of GM1–GM1 can be synthesized from GD1a apart from its classical synthesis (Miyagi & Yamaguchi, 2012; Sonnino

**FIGURE 7** Age-specific changes in GM1, GD1a, and neuroplastin expression upon chronic stress in the CA1 hippocampal region. Immunofluorescent staining was analyzed using the shares of single molecules: GD1a, GM1 and Np65; double colocalizations: GD1a/Np65, GM1/Np65, GD1a/GM1; and triple colocalization in absolute pixel counts for all four animal groups. The average ratio of pixel count is shown for each single or combination profile. The Venn diagram defines the expression of combination profiles. Statistical analysis was performed using a two-way analysis of variance (ANOVA) and is shown in Table 1. Abbreviations: CA1, Cornu Ammonis subfield 1; GD1a, ganglioside GD1a; GM1, ganglioside GM1 visualized by cholera toxin beta subunit; NS, nonsignificant; Np65, neuroplastin; OF-C, control group of old females; OF-S, stressed group of old females; YF-C, control group of young females; YF-S, stressed group of young females
et al., 2011) and it was previously found to be implicated in neurodegenerative processes (Chiricozzi et al., 2020; Kolyovska, 2016). GD1a concentration in the frontal brain decreases with aging (Svennerholm et al., 1989), and a similar decrease was observed in AD patients, especially in the hippocampus (Mesa-Herrera et al., 2019; Svennerholm, 1994).

The shares of all three markers (GM1, Np65, and GD1a) and their combinations (Figures 7 and 8) provided information of each epitope and its interactions in the total number of molecules observed at a given location. We calculated these shares to analyze the changes in the membrane architecture. When the total pixel number was divided in single, double, and triple staining, the shares of neuroplastin contributions (i.e. GM1/Np65, GD1/Np65) in young stressed animals were not changed, and neuroplastin environment in the CA1 remained constant. In old females, however, the shares of free GM1 and free neuroplastin changed—GM1 decreased and neuroplastin increased. If aging was observed without stress—a neuroplastin decrease was obvious, while its double colocalizations remained invariable and triple colocalization increased. Altogether, stress in young animals and stress-free aging in the CA1 region can be figuratively described as playing the same game with fewer players on the field. When aged animals are exposed to stress, there are too few players on the field to deal with additional challenges:
GM1 is consumed in the interaction with neuroplastin and GD1a, while GD1a enters the field (increased expression) as a back-up for GM1. These findings are summarized in Figure 10, where we propose a potential model of neuroplastin interaction with the two ganglioside species in the membrane lipid microenvironments. Each specific environment might influence neuroplastin functionality, and specific combinations could underlie synaptic dysfunction. The strongest cognitive decline was observed in old stressed females, who had a GD1a-rich environment in their CA1 and DG. The CA1 and DG perform different functions (Barth et al., 2018), which were reflected in our analysis of membrane architecture. In the DG of young stressed females, free neuroplastin share increased and the GM1 and GD1a count decreased. These changes consumed the available GD1a molecules alone or in interactions, extensively changing the

**TABLE 1** Statistical significance of the GD1a/GM1/Np65 staining in the CA1 region

| Molecule/overlap combinations | Statistical significance of the main effects or their interactions in CA1 region | Post hoc (Bonferroni) |
|-------------------------------|--------------------------------------------------------------------------------|----------------------|
|                               | Age | Stress | Interaction |                                      |                            |
| GD1a                          | $F_{1,25} = 40.901, p < 0.001$ | $F_{1,25} = 38.439, p < 0.001$ | $F_{1,25} = 39.328, p < 0.001$ | $YF-C$ versus $YF-S$, $p < 0.001$ |
| GM1                           | $F_{1,25} = 108.406, p < 0.001$ | $F_{1,25} = 26.608, p < 0.001$ | NS | - |
| Np65                          | $F_{1,25} = 94.052, p < 0.001$ | NS | NS | - |
| GD1a/GM1                      | $F_{1,25} = 20.028, p < 0.001$ | $F_{1,25} = 22.606, p < 0.001$ | $F_{1,25} = 5.917, p = 0.022$ | $YF-C$ versus $YF-S$, $p < 0.001$ |
| GD1a/Np65                     | $F_{1,25} = 13.411, p = 0.001$ | $F_{1,25} = 12.517, p = 0.002$ | NS | - |
| Np65/GM1                      | $F_{1,25} = 122.564, p < 0.001$ | $F_{1,25} = 29.788, p < 0.001$ | NS | - |
| GD1a/GM1/Np65                 | NS | $F_{1,25} = 6.206, p = 0.02$ | NS | - |

**Note:** Statistical analysis was performed using a two-way ANOVA. Bonferroni correction was used for post hoc comparisons. Significant effects of stress or age and their interaction are marked as exact $p$-values.

**Abbreviations:** ANOVA, analysis of variance; CA1, Cornu Ammonis subfield 1; GD1a, ganglioside; GM1, ganglioside GM1 visualized by cholera toxin beta subunit; Np65, neuroplastin; NS, nonsignificant; OF-C, control group of old females; OF-S, stressed group of old females; YF-C, control group of young females; YF-S, stressed group of young females.

**TABLE 2** Statistical significance of the GD1a/GM1/Np65 staining in the DG region

| Molecule/overlap combinations | Statistical significance of the main effects or their interactions in DG region | Post hoc (Bonferroni) |
|-------------------------------|--------------------------------------------------------------------------------|----------------------|
|                               | Age | Stress | Interaction |                                      |                            |
| GD1a                          | $F_{1,22} = 20.625, p < 0.001$ | NS | $F_{1,22} = 21.495, p < 0.001$ | $YF-S$ versus $OF-S$, $p < 0.001$ |
| GM1                           | NS | NS | $F_{1,22} = 32.177, p = 0.000$ | - |
| Np65                          | $F_{1,22} = 163.987, p < 0.001$ | $F_{1,22} = 4.923, p = 0.037$ | $F_{1,22} = 6.590, p = 0.018$ | $YF-C$ versus $YF-S$, $p = 0.002; YF-S$ versus $OF-S$, $p < 0.001$ |
| GD1a/GM1                      | $F_{1,22} = 7.173, p = 0.014$ | NS | $F_{1,22} = 24.994, p < 0.001$ | $YF-S$ versus $OF-S$, $p < 0.001$ |
| GD1a/Np65                     | NS | $F_{1,22} = 5.883, p = 0.024$ | $F_{1,22} = 37.591, p < 0.001$ | $YF-C$ versus $YF-S$, $p < 0.001; OF-C$ versus $OF-S$, $p = 0.019$ |
| Np65/GM1                      | $F_{1,22} = 5.864, p = 0.024$ | NS | $F_{1,22} = 12.712, p = 0.002$ | - |
| GD1a/GM1/Np65                 | $F_{1,22} = 55.026, p < 0.001$ | NS | $F_{1,22} = 20.542, p < 0.001$ | $YF-S$ versus $OF-S$, $p < 0.001$ |

**Note:** Statistical analysis was performed using a two-way ANOVA. Bonferroni correction was used for post hoc comparisons. Significant effects of stress or age and their interaction are marked as exact $p$-values.

**Abbreviations:** ANOVA, analysis of variance; DG, dentate gyrus; GD1a, ganglioside; GM1, ganglioside GM1 visualized by cholera toxin beta subunit; Np65, neuroplastin, OF-C, control group of old females; NS, nonsignificant; OF-S, stressed group of old females; YF-C, control group of young females; YF-S, stressed group of young females.
membrane architecture. In aged animals, free neuroplastin almost completely disappeared from the DG membranes, while elevated GD1a expression compensated for GM1 decrease. When we added stress to aging, the main effect was free GD1a increase compared with young animals, which probably means that neuroplastin in the DG does not use the available GD1a for interaction.

An additional stress mechanism in reproductive senescence was revealed by IDV analysis. Young stressed females experienced a decreased expression of both gangliosides, while old stressed females experienced a decreased GM1 expression and an increased GD1a expression in both hippocampal regions. Neuroplastin in the CA1 decreased upon stress independently from aging. However, in the DG of old stressed females, neuroplastin expression increased.

IDV can be considered as the membrane epitope scattering measure (Pavel et al., 2020). Based on IDV analysis, we propose that aging generally decreases the diameter of individual microdomains and seriously compromises their function, that is, in synaptogenesis.

Overall, the pie charts revealed the players in the CA1 and DG membrane architecture and their changes in response to stress and aging, while IDV supported the idea that in such conditions neuroplastin changes, as well as its environment.

A 3D tissue reconstruction presented the complexity of the CA1 and DG regions. Although the reconstructed images pointed to qualitative peculiarities of both regions, the method's limitations, such as rapid photobleaching and bleed-through between channels, prevented us from performing anything more than qualitative analysis. Due to the technical limitations, only two molecules, neuroplastin and GM1, were chosen for reconstruction. The expressions of both molecules decreased in response to stress and aging, except in the DG of old stressed females, where neuroplastin increased but became more scattered. This effect might be attributed to the association of neuroplastin with GD1a but not with GM1.

Future studies might further elucidate these processes by biochemical isolation of lipid rafts. However, hippocampal regions with different functions, such as the CA1 and DG, would be challenging to separate. In addition, all lipid rafts isolation protocols involve the use of detergents. In this study, we intentionally did not use detergents as they were shown in our previous study to lead to glycolipid transfer into the adjacent membranes even in fixed tissue (Heffer-Lauc et al., 2005). As our study’s focus was the glycolipid environment of neuroplastin in the CA1 and DG of old stressed females, where neuroplastin increased but became more scattered.

The interpretation of memory impairment process could be facilitated by performing additional behavioral tests or by performing the passive avoidance test at the beginning and end of the study. Additional behavioral data would have allowed us to better understand memory impairment fluctuation during the study. However, as the passive avoidance test strongly activates the amygdala, hippocampus, and entorhinal cortex (Izquierdo et al., 1997), it was performed only once.
Nonstressful tests, such as the object location memory test, that would be required to differentiate the effects of aging and stress on memory changes compared with the changes in the fear/stress response to the shock during the passive avoidance memory test were not available.

Memory is known to be affected by the hormonal status (Ali et al., 2018), but we did not determine the estrous stage for several reasons. If we had paired the animals according to the estrous stage, the number of animals per group would not have reached the expected sample size. Even if the animals entered the study with the same hormonal status, we expected desynchronization due to dysregulation of the gonadal axis by chronic stress (An et al., 2020; Lovick, 2012), especially considering the long study duration. Additional desynchronization was expected due to the (pre)menopausal age of the older animal group (Cruz et al., 2017).

Overall, this study suggests that chronic stress response did not affect the cognition of young female rats, even though some of the investigated molecules were expressed differently. However, old females exhibited diminished memory in response to both stress and aging. Cognitive dysfunction in reproductively senescent females might be underlied by expressional changes in the lipid environment of APP, neuroplastin, AMPA-R, glial cell number, as well as structural changes of the membrane lipids. The postulation of four possible neuroplastin environments pointed to GD1a enrichment in old stressed female rats, as well as its high dispersion in both hippocampal areas. The specific lipid environment might influence neuroplastin functionality, underlying synaptic dysfunction and amyloidogenesis triggered by the combination of aging and chronic stress.

In conclusion, our newly developed Python method proved to be a useful tool for the analysis of membrane
microenvironments in situ, which makes it suitable for use in similar studies.

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CONFLICTS OF INTEREST
The authors have no conflict of interest to declare.

AUTHOR CONTRIBUTIONS
MB, RG, SGV, and MH: conceptualized the study and designed methodology; MB, VI, SB, AB, ACA, DPP, and KFS: carried out investigation and acquired the formal analysis; BK, IL, RG, AB, SGV, RM, and MH: analyzed or interpreted the data, reviewed, and edited the manuscript; MB: writing–original draft. All authors critically revised the manuscript for important intellectual content; all authors gave final approval of the version to be submitted; all authors agree to be held accountable for all aspects of the work.

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