Identification of a Conserved Gene Signature Associated with an Exacerbated Inflammatory Environment in the Hippocampus of Aging Rats

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ABSTRACT: There have been a few descriptive studies in aged rodents about transcriptome changes in the hippocampus, most of them in males. Here, we assessed the age changes in spatial memory performance and hippocampal morphology in female rats and compared those changes with changes in the hippocampal transcriptome. Old rats displayed significant deficits in spatial memory. In both age groups, hole exploration frequency showed a clear peak at hole 0 (escape hole), but the amplitude of the peak was significantly higher in the young than in the old animals. In the hippocampus, there was a dramatic reduction in neurogenesis, whereas reactive microglial infiltrates revealed an inflammatory hippocampal state in the senile rats. Hippocampal RNA-sequencing showed that 210 genes are differentially expressed in the senile rats, most of them being downregulated. Our RNA-Seq data showed that various genes involved in the immune response, including TYROBP, CD11b, C3, CD18, CD4, and CD74, are overexpressed in the hippocampus of aged female rats. Enrichment analysis showed that the pathways overrepresented in the senile rats matched those of an exacerbated inflammatory environment, reinforcing our morphologic findings. After correlating our results with public data of human and mouse hippocampal gene expression, we found an 11-gene signature of overexpressed genes related to inflammatory processes that was conserved across species. We conclude that age-related hippocampal deficits in female rats share commonalities between human and rodents. Interestingly, the 11-gene signature that we identified may represent a cluster of immune and regulatory genes that are deregulated in the hippocampus and possibly other brain regions during aging as well as in some neurodegenerative diseases and low-grade brain tumors. Our study further supports neuroinflammation as a promising target to treat cognitive dysfunction in old individuals and some brain tumors. © 2017 Wiley Periodicals, Inc.

Key Words: aging; spatial memory; hippocampal transcriptome; immune pathways; Barnes maze

INTRODUCTION

In humans and rats, aging is associated with a progressive deterioration of spatial learning and memory. These functional alterations are correlated with morphological changes in the brain, particularly in the hippocampus, a key brain region for the formation and consolidation of spatial memory (O’Keefe et al., 1976; Squire et al., 1993). Two of the most conspicuous changes associated with aging in the rat hippocampus are a sharp decline of neurogenesis in the hippocampal dentate gyrus (Morel et al., 2015) and a progressive activation of the microglia, the immune effector of inflammation and antigen recognition in the brain (Streit et al., 2004). There is clear evidence that a constellation of gene expression changes underlie hippocampal phenotype aging. Thus, gene expression studies in aging rodents have documented significant changes in hippocampal genes related to cholesterol synthesis, inflammation, transcription factors, neurogenesis, and synaptic plasticity (Blalock et al., 2003; Verbitsky et al., 2004; Burger et al., 2007, 2008; Rowe et al., 2007). While those studies revealed that aging itself is associated with the majority of gene expression changes, a smaller portion of the transcriptional differences in the hippocampus are related to changes in learning and spatial memory performance. To our knowledge, most studies assessing age-related transcription changes in the hippocampus of rodents have been performed in males. In the only study we are aware of, electrophoretic gel shift assays were used to document that the binding activity of nuclear factor kappa B, activator protein-1, peroxisome proliferator-activated receptor, and liver X receptor in the cortex and hippocampus from young (3-month old) and old (18-month old) male and female...
rats, display a complex pattern of changes in their brain-binding activity, depending on the anatomical origin of the samples (cortex or hippocampus), and the sex of the animals studied (Sanguino et al., 2006). Therefore, there still is a significant information gap concerning possible sex-related differences in the impact of aging on the hippocampal transcriptome in rodents. In humans, gene expression profiles were assessed in the hippocampus, entorhinal cortex, superior-frontal gyrus, and postcentral gyrus across the lifespan of 55 cognitively intact individuals aged 20–99 years (Berchtold et al., 2008). The results revealed clear gender differences in brain aging, suggesting that the human brain undergoes sexually dimorphic changes in gene expression not only in development but also in later life. More gene expression changes were detected in males than in females across all brain regions. Therefore, it is likely that aging in rodents is also associated with sex-related differences in gene expression in the brain. Since we have characterized the impact of aging on cognitive performance and hippocampal morphology in female rats, it was of interest to correlate these changes with transcriptome changes in the hippocampus of aging female rats. To this end, we performed RNA-seq analysis of the whole female rat hippocampus and correlated our results with public databases recording age-related changes in gene expression in the hippocampus of male rodents and humans of both sexes.

Interestingly, we identified an 11-gene signature of overexpressed genes related to inflammatory processes conserved across species, providing novel insight into aging mechanisms and age-associated disorders in the brain.

## METHODS

### Animals

Two-month old (young) and 28-months old female Sprague-Dawley rats were used. The young and old animals (8 per group) weighed 199 ± 1 and 267 ± 5 g, respectively.

Rats were housed in a temperature-controlled room (22 ± 2°C) on a 12:12 h light/dark cycle. Food and water were available ad libitum. All experiments with animals were performed in accordance to the Animal Welfare Guidelines of NIH (INIBIOLP’s Animal Welfare Assurance No A5647-01). The ethical acceptability of the animal protocols used here have been approved by our institutional IACUC (Protocol # T09-01-2013).

### Spatial Memory Assessment

The modified Barnes maze protocol used here has been previously documented (Morel et al., 2015). It consists of an elevated (108 cm to the floor) black acrylic circular platform, 122 cm in diameter, containing 20 holes around the periphery. The holes are of uniform diameter (10 cm) and appearance, but only one hole is connected to a black escape box (tunnel). The escape box is 38.7 cm long × 12.1 cm wide × 14.2 cm in depth and it is removable. A white cylindrical starting chamber (an opaque, 25 cm in diameter and 20 cm high, open-ended chamber) is used to place the rats on the platform with a random orientation of their bodies.

Four proximal visual cues are placed in the room, 50 cm away from the circular platform. The escape hole is numbered as Hole 0 for graphical normalized representation purposes, the remaining holes being numbered 1–10 clockwise, and −1 to −9 counterclockwise. During the whole experiment, Hole 0 remained in a fixed position, relative to the cues in order to avoid randomization of the relative position of the escape box. A 90-dB white-noise generator and a white-light 500-W bulb provided the escape stimulus from the platform. At the beginning of the experiment, rats were habituated to the task. The habituation routine consists of placing the animals in every compartment during 180 s (starting chamber and escape box). An acquisition trial (AT) consists of placing a rat in the starting chamber, located at the center of the platform, for 30 s; the chamber is then raised, the aversive stimuli (bright light and high pitch noise) are switched on, and the rat is allowed to freely explore the maze for 120 s. The purpose of ATs is to train the rats on finding the escape hole. A probe trial (PT) is similar to an AT except that the escape box has been removed, its purpose being to assess recent spatial memory retention. During PTs, the rats explore the maze for 90 s. The behavioral performances were recorded using a computer-linked video camera mounted 110 cm above the platform. The performance of the subjects was determined using the Kinovea v0.7.6 (http://www.kinovea.org) software. The behavioral parameters assessed were as follows.

a. Escape box latency: Time (in s) spent by an animal since its release from the start chamber until it enters the escape box (during an AT) or until the first exploration of the escape hole (during a PT).

b. Nongoal hole exploration: Number of explorations of holes different from the escape one. Each exploration of an incorrect hole is counted as an error, provided that the rat lowers its nose below the plane of the table surface.

c. Exploration frequency: The times the rat explores every hole of the maze.

### Experimental Design

The day for the first AT was termed experimental day 1. In a first stage, rats were submitted to the Barnes maze protocol for 9 days, each day involving four ATs. On the last day of this series, rats were submitted to a PT (PT0), 15 min after the last AT.

In order to follow the cognitive performance further, 1 month after PT0, the rats were submitted to a short reinforcement protocol of 3 days, of four ATs each. On the last day (Day 41), the subjects were submitted to another PT, the PT1 (Fig. 1B). The rationale for choosing this design was that replicating the test after 1 month would provide a more reliable estimate of spatial memory at each age than a single testing. At
the young and old ages chosen, average spatial memory was expected to show little variation in a 30-day interval.

**Brain Processing**

At the end of the experiment, all rats were euthanized. Four rats from each group were sacrificed by rapid decapitation and their brains processed for transcriptonal analysis as described below. The remainder of the rats were placed under deep anesthesia and perfused with phosphate buffered para-formaldehyde 4%, (pH 7.4) fixative. The brains were removed and stored in para-formaldehyde 4%, (pH 7.4) overnight at 4°C. Brains were kept in cryopreservative solution at −20°C until use. For immunohistochemical assessment, brains were cut coronally in 40-μm-thick sections with a vibratome (Leica).

**Immunohistochemistry**

All immunohistochemistry techniques were performed on free-floating sections. For each animal, separate sets of sections were processed using either an anti-doublecortin goat polyclonal antibody (dilution 1:1000; Santa Cruz Biotechnology Cat# sc-8066, RRID:AB_2088494) or an anti-Ilba-1 rabbit polyclonal antibody (dilution 1:1000; Wako Cat# 016-20001, RRID:AB_839506). Briefly, after overnight incubation at 4°C with the primary antibody, sections were incubated with biotinylated horse anti-goat serum (dilution 1:300; Vector Laboratories Cat# BA-9500, RRID:AB_2336123), or a goat anti-rabbit serum (dilution 1:300; Vector Laboratories Cat# BA-1000, RRID:AB_2313606), as appropriate, for 120 min, rinsed and incubated with avidin-biotin-peroxidase complex (dilution 1:500; Vector Laboratories Cat# PK-6100, RRID:AB_2313606), as appropriate, for 120 min, rinsed and incubated with avidin-biotin-peroxidase complex (dilution 1:500; Vector Laboratories Cat# PK-6100, RRID:AB_2336819) for 90 min and then incubated with 3, 3-diamino benzidine-tetrahydro-chloride (DAB). Sections were counterstained with cresyl violet (Nissl staining) as described elsewhere (Morel et al., 2015). Finally, they were dehydrated and mounted with Vectamount (Vector) and used for image analysis.
Image Analysis

In each hippocampal block, one every six serial sections was selected in order to obtain a set of noncontiguous serial sections spanning the dorsal hippocampus. Typically, a whole dorsal hippocampus comprises about 48 coronal sections, thus yielding eight sets of noncontiguous serial sections (240 µm apart). The number of cells was assessed in the dorsal hippocampus, which is typically located between coordinates −2.8 to −4.5 mm relative to the bregma (Paxinos et al., 1998). For this task, we used an Olympus BX-51 microscope attached to an Olympus DP70 CCD video camera (Tokyo, Japan). All morphological parameters were assessed bilaterally.

Within the hippocampus, we focused our stereological assessment on immature neurons and microglial cells in the Dentate Gyrus (DG) and in the Stratum Radiatum (SR), respectively, two regions vulnerable to degeneration in AD patients (West et al., 1994).

Neuroblast Analysis

A stereological assessment of migrating neuroblasts, immunoreactive to doublecortin antibody (DCXir), was performed in the DG, including the subgranular zone (SGZ) and granular cell layer (GCL). DCXir cell number was assessed using a modified optical fractionator technique as previously described (Morel et al., 2015). To this end, the entire DG sections were used as the counting frame, making the area sampling fraction (asf) = 1. The section sampling fraction (ssf) was 1/6. Estimates were based on counting DCXir cell bodies as they came into focus. N= 3 animals per group.

Microglial Cell Analysis

Microglial cells were identified as Iba-1 immunoreactive cells (Iba1ir) and counted in the hippocampal SR, whose upper limit is the CA1 pyramidal layer, lower limit is Stratum Lacunosum Moleculare, and the lateral limit is the Stratum Lucidum of the dorsal hippocampus (Paxinos et al., 1998). To this end, a random grid consisting of squared probes (area = 5,625 µm²) was superimposed over calibrated images taken at ×400 magnification and cells inside the probe area were counted. Cells making contact with boundary inclusion lines of the probes were included in the count, whereas cells in contact with boundary exclusion lines were not counted. Iba-1ir cells were morphologically classified as Type I, II, III, IV, and V on the basis of previously documented criteria (Diz-Chaves et al., 2012). Types I, II, and III were categorized as nonreactive glia whereas Types IV and V were taken as reactive glia. N= 3 animals per group.

RNA Extraction, Library Preparation, and Sequencing

Four rats per group were euthanized and their right hippocampus dissected for transcriptome analysis. The hippocampus was stored at −80°C until RNA extraction. Tissues were homogenized in TRIzol Reagent (Life Technologies). The quality of the isolated RNA was assessed by measuring the RIN (RNA Integrity Number) using the Fragment Analyzer. Library preparation for RNA-Seq was performed using the truSeq RNA Sample Preparation Kit starting from 500 ng of total RNA. Accurate quantitation of cDNA libraries was performed using the QuantiFluor TM dsDNA System (Promega). The size range of final cDNA libraries was 280–320 bp and was determined applying the DNA Chip for NGS Libraries using the Fragment Analyzer (Advanced Analytical). cDNA libraries were amplified and sequenced by using the cBot and HiSeq2000 from Illumina (SR: 50 bp; ca. 30–35 million reads per sample). Raw datasets have been submitted to NCBI GEO database.

Real-Time Quantitative PCR

Total RNA was treated with gDNA wipeout and cDNA was synthesized with the Qiagen QuantiTect Reverse transcription kit (#205310). qPCR was performed with the MESA BLUE qPCR MasterMix Plus for SYBR Assay Low ROX on a Stratagene Mx3000P qPCR system. The primers used are listed in Supporting Information Table S1.

RNA-Seq Data Analysis

Illumina HiSeq 2000 fluorescence images were transformed to BCL files with the Illumina BaseCaller software and samples were demultiplexed to FASTQ files with CASAVA (version 1.8.2). Sequencing quality was checked and approved via the FastQC software. Sequences were aligned to the genome reference sequence of Rattus norvegicus (RGSC assembly v5.0) using the STAR alignment software (Dobin et al., 2013; version 2.3.0e) allowing for two mismatches within 50 bases. Subsequently, resulting SAM files were converted to sorted BAM files, filtering of unique hits and counting was conducted with SAMtools (Li et al., 2009, version 0.1.18) and HTSeq (Anders et al., 2015; version 0.6.1p1).

To identify differentially expressed genes between hippocampal samples from young and old rats, we utilized the DESeq2 algorithm based on the normalized number of counts mapped to each gene (Love et al., 2014). Functional enrichment analyses were performed using the database for annotation, visualization and integrated discovery (DAVID, http://david.abcc.ncifcrf.gov/), Enrichr (http://amp.pharm.mssm.edu/Enrichr/), and InnateDB resource (http://www.innatedb.com/) resources based on the list of deregulated transcripts between hippocampal groups (P-adj. < 0.05; fold changes > 2). Data integration and heatmap visualization of differentially expressed transcripts were done with R and the Multi Experiment Viewer software (MeV v4.9) (Saeed et al., 2014). In addition, we used the ESTIMATE algorithm (Estimation of Stromal and Immune cells in Malignant Tumors using Expression data) to infer the immune component from each RNA-Seq Hippocampal sample (Yoshihara et al., 2013).
Comparative Analysis of the Hippocampal Signature between Aging Rats and Human Counterparts

Gene expression profile of deregulated genes (n = 210) obtained by RNAseq in the hippocampus of young and aging rats was assessed in a human context. Microarray data (Affymetrix) from normal human brain in aging individuals was obtained from the study GSE11882; CEL files were downloaded from GEO (http://www.ncbi.nlm.nih.gov/geo/), processed and normalized with R/Bioconductor by using the fRMA package. The original dataset includes microarrays from four brain regions (hippocampus, entorhinal cortex, superior frontal gyrus, and postcentral gyrus) of cognitively intact humans (age 20–99 years). For the purpose of this study only hippocampal samples were considered (n = 43). From the 210 deregulated genes, a total of 170 orthologous genes were found in the human Affymetrix Platform (HG-U-133 Plus_2), thus obtaining a final matrix of 43 samples vs. 170 genes. Samples were then categorized into two groups: young (age 20–50;
FIGURE 3. Age-related gene expression changes in the hippocampus. (A) Unsupervised hierarchical clustering of rat hippocampal samples. (B) Heat map of the 210 differentially expressed genes between young and old hippocampal samples. (C) Gene ontology (GO) analysis of deregulated transcripts revealed by functional enrichment analysis. (D) Set of genes commonly deregulated across mice and rat models strongly associated with the aging process. (E) Box-plot depicting the immune score difference between young and old rat, as assessed with ESTIMATE. (F) Box-plot of eight representative genes validated by quantitative RT-PCR. The results confirm the RNA-seq data. [Color figure can be viewed at wileyonlinelibrary.com]
RESULTS

Latency to Escape Box

To assess hippocampal function in rats, we used a modified Barnes maze test. After Day 1, escape hole latency fell rapidly in young and old rats, reaching a plateau around experimental day 4 [two-way repeated measures (RM) ANOVA day factor degrees of freedom (DF)= 11; F = 49.057; P < 0.001]. The plateau remained relatively constant for both age groups from Day 4 until Day 41 (the last day tested), a time span where latency remained consistently lower in the young as compared to the old rats (Fig. 1A, upper panel) (two-way RM ANOVA day factor F = 4.922; P = 0.044; Holm-Sidak test P < 0.05 for Days 7, 8, 9, 39, 40, and 41). As expected, the young group had a lower latency in both PT0 and PT1 (Mann–Whitney test P = 0.015 and P = 0.003 for PT0 and PT1, respectively) (inset in Fig. 1A, upper panel).

Errors to Escape Box

During the training series, the error number fell sharply in both age groups until reaching a plateau at Day 3–4 (two-way RM ANOVA day factor DF = 11; F = 20.570; P < 0.001). The young animals made significantly fewer errors than the old group on every experimental day, except for Day 4 (Fig. 1A, lower panel) (two-way RM ANOVA group factor DF = 1; F = 32.740; P < 0.001) (Holm-Sidak test for Day 4 P = 0.223). The young group made fewer errors than the old group in both PT0 and PT1 (Student’s t-test P < 0.001 and Mann–Whitney test P < 0.001 for PT0 and PT1, respectively) (inset in Fig. 1A, lower panel).

Hole Exploration Frequency

In both age groups, hole exploration frequency showed a clear peak at hole 0 in both PT0 and PT1 (Fig. 1C). The amplitude of the peak was significantly higher in the young than in the old animals in both PTs (Student’s t-test P = 0.015 and Mann–Whitney test P < 0.001 for PT0 and PT1, respectively).

Age-Associated Morphological Changes in the Dorsal Hippocampus

In order to determine whether the behavioral alterations were associated with morphological changes in the brain, a stereological assessment of immature neurons and microglial cells was performed in the dorsal hippocampus of the young and aged rats. As expected, there was a sharp reduction in the number of DCXir [strike_start] [strike_end] cells, (immature neurons) in the DG of old rats as compared with their young counterparts (Student’s t-test P < 0.001) (Figs. 2A–C).

Hippocampal Iba1-ir microglial cells were classified as non-reactive and reactive, and were counted in the hippocampal SR. Although we found no significant differences in the number of non-reactive microglial cells (ANOVA factor group F = 1.710; P = 0.262) we observed a significant increase in reactive microglial cells in the old versus the young group (ANOVA group factor F = 9.200; P = 0.0402) (LSD test 0.0321). The total number of microglial cells (non-reactive and reactive) in the SR showed a trend towards an increase in the old versus young animals (ANOVA group factor F = 5.175; P = 0.0624) (Figs. 2D–F).

Age-Related Gene Expression Changes in the Hippocampus

Next, we interrogated the transcriptome in order to identify gene expression differences that could underlie the morphological differences and explain the functional alterations observed. Unsupervised hierarchical clustering analysis of RNA-Seq profiles demonstrates a clear segregation of young and old hippocampal tissue (Fig. 3A). Statistical analysis of RNA-Seq data revealed 210 transcripts differentially expressed between young and old hippocampal samples (p-adjust < 0.05; Fold changes > 2). Among the deregulated genes, 81 were up-regulated and 129 were down-regulated transcripts in old rats (Fig. 3B). The top 20 up- and down-regulated genes are shown in Table 1. This Table also shows the cell type in which the genes enlisted are expressed genes between the two groups, a t-test method was applied (P < 0.05).

Statistical Analysis

BM parameters and DCXir count estimates were analyzed with the Sigma Plot v.11 software (Systat Software, San Jose, CA). For BM latency and error analysis, the data from each day’s ATs were averaged. These parameters were analyzed by two-way repeated measures ANOVA, considering group and day factors. When this ANOVA was significant, comparisons between means were performed with the Holm-Sidak post hoc test.

Statistics for Iba1-ir cell count was conducted with the Statistica 8 software (Tulsa, USA). Thus, we performed an analysis of variance (ANOVA) with a nested mixed model, in which variance was hierarchically decomposed into three categories, namely “Age group-Rat-Brain section.” Age group was set as the fixed grouping variable, whereas rat and brain section were set as random variables. Iba1-ir count data were taken as the dependent variable. When ANOVA was significant, post hoc LSD test for differences between means was performed. P < 0.05 was set as significant.

In all cases, plots were performed with the Sigmaplot v.11 Software and show mean ± SEM.

n = 17) and old (age 51–99; n = 26). To get differentially expressed genes between the two groups, a t-test method was applied (P < 0.05).
analysis of deregulated transcripts between the two groups of samples revealed specific GO terms related to antigen processing via MHC II-immune response, cell adhesion - integrin-mediated pathway and response to corticosteroid - steroid hormones (Fig. 3C).

Table 1. List of the Top 40 Deregulated (Up and Down) Hippocampal Genes in Old Rats. Orange cells represent upregulated genes in old rats whereas green cells represent downregulated genes. RNA cell source has been obtained from public information on brain transcriptomics http://web.stanford.edu/group/barres_lab/brain_maseq.html. OPC, oligodendriag progenitor cell; NFO, newly formed oligodendrocyte; MO, myelinating oligodendrocyte. [Color table can be viewed at wileyonlinelibrary.com]

| Gene     | Description                                      | logFC     | Cell source                  |
|----------|--------------------------------------------------|-----------|-----------------------------|
| Slc9a3   | Solute carrier family 9, subfamily A             | 2.545671892 | N/S                         |
| Bpifb4   | BPI fold containing family B, member 4           | 2.176808781 | N/S                         |
| RT1-Ba   | RT1 class II, locus Ba                           | 2.12724301 | Not found                   |
| Cml3     | Probable N-acetyltransferase CML3                | 2.109236546 | Mostly astrocytes, also OPCs |
| C3       | Complement component 3                          | 1.98344423 | Microglia                   |
| Clec5a   | C-type lectin domain family 5, member A          | 1.967973654 | Mostly microglia, also OPC   |
| Fmo3     | Flavin containing monoxygenase 3                | 1.967494475 | N/S                         |
| Cd74     | Cd74 molecule, major histocompatibility complex  | 1.853031498 | Microglia                   |
| RT1-Db1  | RT1 class II, locus Db1                          | 1.839374172 | Not found                   |
| Clec12a  | C-type lectin domain family 12, member A         | 1.738702616 | Microglia                   |
| Lilrb3li | Leukocyte immunoglobulin-like receptor, subfamily B | 1.717229122 | Microglia                   |
| Cd4      | Cd4 molecule                                    | 1.699012324 | Microglia                   |
| Tbx4     | T-box 4                                         | 1.680920772 | Endothelial cells           |
| Card11   | Caspase recruitment domain family, member 11     | 1.674925125 | Mostly microglia, also OPC   |
| Pbd1     | Phenazine biosynthesis-like protein domain       | 1.620714722 | Astrocytes                  |
| LSP1     | Lymphocyte specific 1                           | 1.61610002 | Microglia                   |
| Aaad1    | Aminoadipate aminotransferase                   | 1.608310278 | Mostly neurons, also astrocytes and OPCs |
| Spp1     | Secreted phosphoprotein 1                       | 1.575763833 | Microglia                   |
| CTXN3    | Cortexin 3                                      | 1.574417002 | Mostly neurons, also MOs and OPCs |
| Gpr183   | G protein-coupled receptor 183                   | 1.543275371 | Microglia                   |
| Slc22a6  | Solute carrier family 22                         | -1.624224252 | OPCs                        |
| Abca4    | ATP-binding cassette, subfamily A, member 4      | -1.643218961 | Endothelial cells           |
| Tead2    | TEA domain family member 2                      | -1.648008854 | Mostly endothelial cells, also astrocytes |
| Aldh1a2  | Aldehyde dehydrogenase 1 family, member A2      | -1.660328345 | OPCs                        |
| Top2a    | Topoisomerase (DNA) II alpha                     | -1.669620965 | Mostly OPCs, also astrocytes and endothelial cells |
| Cdkn1c   | Cyclin-dependent kinase inhibitor 1C            | -1.681283549 | Mostly endothelial cells, also MOs, NFOs, neurons and OPCs |
| Sema3c   | Sema domain, immunoglobulin domain              | -1.685074175 | Endothelial cells and neurons |
| Thbs2    | Thrombospondin 2                                | -1.68850683 | Mostly neurons, also OPCs, endothelial cells and astrocytes |
| Cpz      | Carboxypeptidase Z                              | -1.701391957 | OPCs                        |
| Nid1     | Nidogen-1                                       | -1.712318577 | Endothelial cells           |
| Gstm2    | Glutathione S-transferase mu 2                  | -1.751363055 | Endothelial cells           |
| Mmp14    | Matrix metalloproteinase 14                     | -1.809848913 | Mostly astrocytes, also endothelial cells, OPCs and neurons |
| Pcolce   | Procollagen C-endopeptidase enhancer            | -1.88163707 | Mostly OPCs, also endothelial cells |
| Igfbp1   | Insulin-like growth factor binding protein-like 1 | -2.077167083 | Neurons                     |
| Col1a2   | Collagen, type I, alpha 2                       | -2.330267894 | OPCs                        |
| Col3a1   | Collagen, type III, alpha 1                     | -2.41594823 | OPCs                        |
| Slc22a2  | Solute carrier family 22, member 2              | -2.423198076 | N/S                         |
| Col1a1   | Collagen, type I, alpha 1                       | -2.619212838 | OPCs                        |
| Igfbp2   | Insulin-like growth factor binding protein 2    | -2.760020834 | Mostly astrocytes, also OPCs and neurons |
| Igf2     | Insulin-like growth factor 2, transcript variant 2 | -2.910105434 | Mostly OPCs, also endothelial cells and neurons |

To identify commonly deregulated genes among our study and those identified in other age-related studies in animals, we performed an enrichment analysis from the 210 deregulated genes with an aging perturbation based database available at Enrichr online resource. This approach enabled us to identify a
set of genes associated with the aging process, significantly deregulated across male mice and male rat models (Fig. 3D, Supporting Information File 2).

Given the aforementioned association with immune-related processes, we developed a gene expression signature–based algorithm to predict the immune component among hippocampal samples. Interestingly, ESTIMATE predicted higher immune scores on old hippocampal samples compared with young brain samples \( (P < 0.01, \text{Fig. 3E}) \).

In order to validate our RNA-Seq data we performed quantitative RT-PCR analysis of eight representative transcripts (Fig. 3F). Genes such as Tyrobp, C3, Itgb2, or gpr183, displayed an

**FIGURE 4.** Hippocampal 11 gene-signature of deregulated genes conserved among species during aging (A) Heat map of the 30 DE genes with aging in human and rat hippocampus. (B) Correlation matrix of the 30 deregulated genes, note the cluster of 11 genes showing the highest degree of correlation (11-gene signature). Dark blue intensity denotes high expression whereas circle diameter is proportional to the correlation value between the pair of genes in turn. (C) Expression correlation between tyrobp and itgb2 in human hippocampi. (D) Expression correlation of the 11 gene-signature related to aging in human hippocampi. (E) Expression of the 11 gene-signature in human hippocampi from old human individuals. [Color figure can be viewed at wileyonlinelibrary.com]
induction of 3- to 8-fold increase in the old animals, whereas 
Kcnj13, Cdh3, Mfrp, Scl22a2 behaved in the opposite way, 
with a 4- to 40-fold repression in old animals (Fig. 3F).

To further explore the differentially expressed genes found in 
rat hippocampus across aging, an in silico analysis in human 
hippocampal samples was performed. The comparison between 
the young and old groups yielded 30 deregulated genes among 
the 210 genes analyzed ($P < 0.05$), of which 24 were upregu-
lated and 6 down-regulated in the group named old (Fig. 4A, 
Supporting Information File 2). The most prominent change 
was evidenced towards the 7th decade, with a strong activation 
of genes such as CLEC7, TYROBP, CD74, ITGB2, C3, 
ITGAM, GPR183 and GAPT. Additionally, we computed a 
correlation matrix with the 30 deregulated genes and found 
that a cluster of 11 genes showed the highest degree of correla-
tion. We named this cluster the 11-gene signature (Figs. 4B,C; 
Table 2). Next, we evaluated the 11-gene signature with respect 
to sex and found that 7 out of the 11 genes are significantly 
upregulated in the group “old” of both sexes, implying that the 
gene signature is independent of sex. Functional analysis of the 
11-gene signature indicated an enrichment in processes related 
with the activation of immune response and MHC protein 
binding. These were the ontology terms more strongly enriched 
in the 210 deregulated genes identified in our animals, con-
fiming the robustness and reproducibility of these 11 genes 
across species. Moreover, in the category of aging enrichment, 
6 out of the 11-gene signature are significantly associated with 
the signature of the CA3 region of the old male rat hippocam-
pus ($P < 0.001$, GSE14724). The performance of the 11-gene 
signature was also evaluated with respect to sex and 
neurodegenerative conditions, such as Huntington’s and Alzheimer’s 
diseases. In both situations, a significant positive association 
was obtained (Figs. 4D,E; $P < 0.001$).

We also evaluated the performance of the 11-gene signature 
on a dataset of Low Grade Brain Tumors obtained from the 
Cancer Genome Atlas (TCGA) ($n = 530$), which includes 
astrocytomas, oligoastrocytomas and oligodendrogliomas. The set 
of samples was clustered into two groups (named high and 
low) according to the expression of the 11-gene signature. A 
significant association between astrocytomas and the group 
high and oligodendrogliomas and the group low was observed 
($P < 0.01$; Supporting Information Fig. 1). A survival analysis 
was then performed which revealed that the group of samples 
belonging to the group high, mostly astrocytomas, had worse 
prognosis in overall survival and disease free survival ($P < 0.01$; 
Supporting Information Fig. 1).

Finally, a protein-protein interaction network-based analysis 
employing STRING online resource indicated strong interactions 
among 7 out of 11 members of the 11-gene signature in rats. More importantly, protein-protein interactions between C3, ITGB2, ITGAM, and TYROBP were also conserved between rat and 
human species (Fig. 4C), suggesting that they are involved 
in an evolutionarily conserved bioprocess (Fig. 5).

### DISCUSSION

#### Functional Changes in the Aging Brain

In rats, two important age-related changes take place in the 
brain during aging. At the functional level, there is an impair-
ment in spatial memory performance as revealed by various 
tests of spatial learning and memory such as the radial arm 
maze (de Toledo-Morrell et al., 1984), the Morris Water Maze 
(MWM; Morris, 1984), and the Barnes maze (Barnes, 1979; 
Morel et al., 2015). This functional decline occurs in parallel 
with a marked reduction in hippocampal neurogenesis in aged 
rats (Barnes, 1979; Morel et al., 2015). Our results confirm 
that like in males, aging in female rats is associated with a decline in spatial memory performance as assessed by the 
Barnes maze, a less physically demanding test than the MWM, 
which is important as animals age. Our data also confirmed 
that in the hippocampal DG, aging brings about a severe fall 
in the number of immature neurons, a change that is likely to 
be causally involved in the decline of certain memory functions.
in old rats (Yau et al., 2015). Some authors divide aging rats in age-impaired and age-unimpaired. We preferred not to use such a subdivision as we wanted to describe typical age-changes. Almost at every level, aging is known to lead to data scattering, be it serum hormone levels, circadian rhythms, etc. As expected, this also happened with some, but not all, cognitive parameters in young versus old rats and this led to a higher variance in the old than in the young animals. Had we subdivided the aging group, we would have been unable to tell which of the two groups represented typical age-related changes, marked impairment or lack/very mild impairment. In fact in our old rats we did not observe a spontaneous segregation into two groups. Rather, we observed a continuum from the less impaired old rats through the more impaired ones. Any subdivision would have to be based on an arbitrarily chosen limit.

Immune reactivity and inflammatory processes increase with aging in the brain (Lucin et al., 2009), with microglia playing a central role in this immune dysregulation (Luo et al., 2010; Norden et al., 2013). Innate immunity within the central nervous system is primarily provided by resident microglia, brain cells that are essential in immune surveillance and that also mediate the coordinated responses between the immune system and the brain. With normal aging, microglia develop a more reactive phenotype. For instance, in several models of aging there are increased levels of proinflammatory cytokines in the brain and increased expression of inflammation-associated receptors on microglia (Luo et al., 2010). There are also major differences in microglial biology between young and old age when the immune system is challenged and microglia activated. In this context, microglial activation is amplified and prolonged in the aged brain compared to adults. Prolonged microglial activation leads to the release of pro-inflammatory cytokines that exacerbate neuroinflammation, contributing to neuronal loss and impairment of cognitive function (Streit et al., 2004; Perry et al., 2010; Jiang et al., 2012). Our data are consistent with these alterations and demonstrate that in aging female rats there is a significant activation of reactive microglia in the hippocampus.

**FIGURE 5.** Biological interactions of the 11-gene signature members. String 10.0 bioinformatics software (http://stringdb.org) was used to search for protein-protein interactions with the 11-gene signature members (left: rat, right: human) in public databases and for the schematic representation of the network of functional interactions. Small nodes indicate protein of unknown 3D structure, while large nodes indicate that some 3D structure is known or predicted. In STRING, each protein-protein interaction is annotated with one or more scores. These scores are indicators of confidence. All scores rank from 0 to 1, being 1 the highest possible confidence. [Color figure can be viewed at wileyonlinelibrary.com]
Gene-Expression Changes in the Hippocampus during Aging

In male rodents, aging and, to a lesser extent, deficits in memory performance, have been associated with changes in hippocampal gene expression (Blalock et al., 2003; Verbitsky et al., 2004; Rowe et al., 2007; Burger et al., 2008). These differences consist mostly of gene upregulation in middle-aged male mice (15-month old) as compared with 2-month-old counterparts (Verbitsky et al., 2004). In the CA1 hippocampal region of old male rats, 233 genes were found to be differentially expressed with aging, 60% upregulated and 40% downregulated (Blalock et al., 2003). In the hippocampi from our female rats, 61% of the 210 transcripts differentially expressed in the old animals were downregulated suggesting that, in the hippocampus of very old animals, gene downregulation prevails. Therefore, a comparison between our females and old male rodents seems to point to a differential impact of age in males versus females concerning the proportion of hippocampal genes up and downregulated. Although this trend needs to be confirmed by further studies, a sex-related impact of age on the hippocampal transcriptome would, in general terms, be in line with the results of Sanguino et al. (2006) in rats and the observations reported in humans (Berchtold et al., 2008).

Our enrichment analysis from the 210 deregulated genes with aging revealed that there is a set of genes commonly deregulated during aging in male mice and rats. In aged male mice, the hippocampus was the brain region with the largest number of deregulated genes that were also present in the set of 210 deregulated genes we found in female rats.

As stated above, we and others have shown that one of most conspicuous changes associated with aging in the rat hippocampus is a sharp decline of neurogenesis in the hippocampal DG (Morel et al., 2015). It is not clear to us why such a substantial phenotypic change was not reflected by a significant deregulation of genes related with neurogenesis. It is likely that if the \( N \) of the hippocampal samples analyzed had been higher the changes could have become significant. Neurogenesis is restricted to the DG; therefore, had we performed RNAseq from solely the DG region, it is likely that significant differences in the expression of these genes had become detectable.

Nevertheless, we want to point out that, in a recent study, we demonstrated that intracerebroventricular (ICV) IGF-I gene therapy increases the neurogenic activity in the DG of old rats (Pardo et al., 2016). Furthermore, in a follow-up study, yet unpublished, we found that ICV IGF-I gene therapy in the whole hippocampus of old rats significantly upregulates the expression of Sypl2, Nnat, and Itga-8, three genes related with synaptic processes. Expression of the neurogenesis-related gene DCX was also significantly upregulated by IGF-I gene therapy. Therefore, we expected a marked impact of aging on the expression of these genes. Since aging involves a constellation of gene expression changes, it could be possible that somehow, this prevented the deregulation of the above genes whereas, IGF-I therapy, which affected the expression of a much smaller number of genes, was able to significantly overexpress the above genes.

From the ESTIMATE-based score it is clear that, at least at the hippocampal level, aging in our female rats is strongly associated with a deregulation of a high number of immune genes, particularly those related to antigen processing via MHC II, immune response, cell adhesion and integrin-mediated pathways. The most affected glial element of the aged hippocampus seems to be the microglia where the deregulation of the TYROBP and CD11b genes may be pivotal to microglia dysfunction in the aged rats (Zhang et al., 2013).

Hippocampal Aging- and Immune-Related Genes

Our RNA-Seq data, validated by quantitative RT-PCR analysis of eight representative transcripts, show that various genes involved in the immune response, including TYROBP, CD11b, C3, CD18, CD4, and CD74, are overexpressed in aged female rats. Among them, TYROBP (tyrosine kinase binding protein) protein also known as DAP-12 (DNAX activating protein of 12 kDa) or KARAP (killer cell activating receptor associated protein) is a key regulator of the immune system. TYROBP acts as an adapter protein signaling numerous cell surface receptors, which play an important role in signal transduction in dendritic cells, osteoclasts, macrophages, and microglia. Accordingly, TYROBP serves as a hub of the "microglial sensome" network, on which major molecular connections are concentrated (Hickman et al., 2013). Loss of function in TYROBP gene causes Nasu-Hakola disease, which is characterized by progressive dementia and multifocal bone cysts (Xing et al., 2015). TYROBP is significantly upregulated in the brains of patients with Alzheimer's disease (Zhang et al., 2013). In the central nervous system TYROBP expression predominates in microglia, enhancing their phagocytic activity, which is responsible for clearing the \( \beta \)-amyloid deposits (\( \beta \bar{A} \)) and apoptotic neurons (Xing et al., 2015). TYROBP is also involved in the suppression of proinflammatory cytokine production responses by the microglia. Thus, TYROBP overexpression during aging may result in imbalances of the neuroinflammatory process.

Integrin CD11b (also known as Integrin alpha M, ITGAM) is one protein subunit that forms the heterodimeric integrin alpha-M beta-2 (\( \alpha M \beta 2 \)) molecule, also known as complement receptor 3 (CR3) (Ling et al., 2014) which positively regulates TLR4-induced signaling pathways in dendritic cells (Ling et al., 2014). In the brain, CD11b integrin expression is restricted to microglia and is a hallmark of their activation (Block et al., 2007). CD11b has recently been implicated in the death of dopaminergic neurons on inflammation (Pei et al., 2007). It has been demonstrated that CD11b and TYROBP (DAP12) are involved in the induction of developmental neuronal death in the hippocampus and that microglia use innate immunity mechanisms to induce the apoptosis of neurons committed to death during development (Wakselman et al., 2008).
The relevance of the other immune genes that change with aging in the hippocampus is not obvious and will, therefore, not be further discussed here.

A Conserved 11-Gene Signature Overexpressed during Aging in Rats and Humans

An in silico analysis in human hippocampal samples performed to identify deregulated genes among the 210 genes analyzed in our rats, showed that from a group of 30 human hippocampal genes common to the deregulated female rat genes there is a cluster of 11 human genes showing the highest degree of correlation in a correlation matrix. When evaluated respective to age and neurodegenerative diseases, such as Huntington’s and Alzheimer’s, this 11-gene signature showed a significant positive association. Interestingly, a functional analysis of the 11-gene signature indicated that the deregulation of these genes is associated with an activation of immune response and MHC protein binding, which are the ontology terms more enriched by the 210 deregulated genes in the hippocampus of our aged female rats. The implication of this finding is that these 11 genes are deregulated in the aging hippocampus in rodents and humans and, since they are associated with an immune activation, it seems plausible to hypothesize that the activation of this 11-gene signature underlies a common dys-functional mechanism of the immune system in the hippocampus during aging.

Some of the 11 signature genes have been discussed above. Others like Pld4, Gpr183, Glec7a, Gpr34, and Gapt will be briefly discussed here. Phospholipase D4 (Pld4) belongs to the Phospholipase D (PLD) family (Terao et al., 2013). Phospholipase D plays an important pathophysiological role in the progression of neurodegenerative diseases, primarily through its capacity as a signal transducer in indispensable cellular processes such as cytoskeletal reorganization and vesicle trafficking (Lindsley et al., 2012). Dysregulation of PLD by the protein α-synuclein has been shown to lead to the specific loss of dopaminergic neurons in mammals. Disinhibition of PLD by α-synuclein may contribute to Parkinson’s deleterious phenotype (Peng et al., 2012). Abnormal PLD activity has also been suspected in Alzheimer’s disease, where it has been observed to interact with presenilin 1 (PS-1), the principal component of the γ-secretase complex responsible for the enzymatic cleavage of amyloid precursor protein (APP).

G-protein coupled receptor 183 (GPR183) also known as EBI2 (Epstein Barr-induced receptor 2) is a protein encoded by the GPR183 gene in humans (Schöneberg et al 1999). Growth factor receptor-bound protein 2 (also known as Grb2 or Gapt) is an adaptor protein involved in signal transduction and cell communication. In humans, the GRB2 protein is encoded by the GRB2 gene (Matuoka et al., 1992; Lowenstein et al., 1992).

It will be interesting to determine whether the 11-gene signature is also deregulated during aging in other species as well as in other pathologies.

Concluding Remarks

The comprehensive characterization of age-related gene expression changes in the hippocampus allowed us to identify a group of 11 related genes commonly deregulated (overexpressed) across species during aging. The 11-gene signature may represent a cluster of immune and regulatory genes that are deregulated in the hippocampus and possibly other brain regions during aging as well as in some neurodegenerative diseases and low-grade brain tumors. This finding may offer the opportunity to seek suitable therapeutic targets among them, for the treatment of some brain pathologies.

AUTHORS’ CONTRIBUTIONS

JP and GRM did the cognitive studies and the analysis of the behavioral data. JP and LF did the RT PCR analysis; MA, EL, TFO and RGG performed the analysis and interpretation of the RNA_seq data. JP, MA, EL and LF designed the different plots and graphs. JP, GRM, MA, EL, TFO, and RGG wrote different sections of the manuscript. GRM, TFO, and RGG assembled the final version of the paper.

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The authors declare no conflict of interest.

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