The adaptable transcriptional response to changes in food availability not only ensures animal survival but also lets embryonic development progress. Interestingly, the CNS is preferentially protected from periods of malnutrition, a phenomenon known as “brain sparing.” However, the mechanisms that mediate this response remain poorly understood. To get a better understanding of this, we used Drosophila melanogaster as a model, analyzing the transcriptional response of neural stem cells (neuroblasts) and glia of the blood–brain barrier (BBB) from larvae of both sexes during nutrient restriction using targeted DamID. We found differentially expressed genes in both neuroblasts and glia of the BBB, although the effect of nutrient deficiency was primarily observed in the BBB. We characterized the function of a nutritional sensitive gene expressed in the BBB, the serine protease homolog, scarface (scaf). Scaf is expressed in subperineurial glia in the BBB in response to nutrition. Tissue-specific knockdown of scaf increases subperineurial glia endoreplication and proliferation of perineurial glia in the blood–brain barrier. Furthermore, neuroblast proliferation is diminished on scaf knockdown in subperineurial glia. Interestingly, reexpression of Scaf in subperineurial glia is able to enhance neuroblast proliferation and brain growth of animals in starvation. Finally, we show that loss of scaf in the blood–brain barrier increases sensitivity to drugs in adulthood, suggesting a physiological impairment. We propose that Scaf integrates the nutrient status to modulate the balance between neurogenesis and growth of the BBB, preserving the proper equilibrium between the size of the barrier and the brain.

**Key words:** blood–brain barrier; Drosophila melanogaster; glial cells; nutrient restriction; Scarface; serine protease homolog

**Significance Statement**

The Drosophila BBB separates the CNS from the open circulatory system. The BBB glia are not only acting as a physical segregation of tissues but participate in the regulation of the metabolism and neurogenesis during development. Here we analyze the transcriptional response of the BBB glia to nutrient deprivation during larval development, a condition in which protective mechanisms are switched on in the brain. Our findings show that the gene *scarface* reduces growth in the BBB while promoting the proliferation of neural stem, assuring the balanced growth of the larval brain. Thus, Scarface would link animal nutrition with brain development, coordinating neurogenesis with the growth of the BBB.
**Introduction**

The formation of the nervous system is a tightly regulated process that is controlled by complex mechanisms that mitigate external perturbations, such as temperature changes and food availability. For example, the developing mammalian brain is protected against intrauterine growth restriction by a phenomenon known as brain sparing in which the animal adapts to a decrease in nutrient availability. For example, the developing mammalian brain is very effective in reducing the levels of carbohydrates and glutathione after complete starvation, a phenomenon that resembles mammal sensitivity to undernourishment and continues proliferating even during early larval development block neurogenesis by arresting NBs in response to nutrition (Kanai et al., 2018). BBB glia mediate the reactivation of neural expression is sensitive to nutrional during larval development. scaf is expressed by the SPG and is required for controlling SPG growth and PG proliferation. Moreover, Scaf is necessary for the proper rate of neurogenesis, and its overexpression enhances NB proliferation and brain growth in animals subjected to NR. Finally, we show that knocking down scaf in SPG affects the resistance of flies to drugs, suggesting that the function of the BBB is impaired when scaf is lost.

**Material and Methods**

Fly stocks and husbandry. Drosophila melanogaster stocks were cultured in fly food medium at 25°C. Our fly food contains the following ingredients per liter of medium: 100 g yeast, 80 g glucose, 50 g wheat flour, 11 g agar, 6 ml propionic acid, and 12 ml 20% Nipagin (methylparaben). All RNAi experiments were performed at 29°C.

For Targeted DNA adenine methyltransferase identification (DamID; TaDa) analysis, we used tub-GAL80ts, UAS-LT3-NDam-RiP215 (NDam-Pollu), and tub-GAL80ts, UAS-LT3-NDam (Southall et al., 2013) crossed to the drivers wor-GAL4 (Albertson et al., 2004), mdr65/R54C07-GAL4 (Jenett et al., 2012; Spéder and Brand, 2014), or sema5C/R71C08-GAL4 (Jenett et al., 2012). For functional experiments we used w¹¹¹є as experimental control, UAS-shScafRNAi (catalog #330286, Vienna Drosophila Resource Center), UAS-IhScafRNAi (catalog #13249, Vienna Drosophila Resource Center), UAS-DlgG::EGFP (Koh et al., 1999); UAS-GFP::nls (catalog #107-870, Kyoto Stock Center), UAS-mCD8-GFP, UAS-mdCD8-RFP, UAS-lam-GFP (catalog #7378, Bloomington Drosophila Stock Center), and UAS-Scaf:GFP (Roussel et al., 2010). For fluorescent reporters and fusion proteins we used the following stocks: scafHG (Scaf::GFP; Bonin and Mann, 2004), 10xSTAT92E-GFP (Bach et al., 2007), TRE-RFP (Chatterjee and Bohmann, 2012), Lanα::GFP (catalog #318155, Vienna Drosophila Resource Center), Lanα::GFP and mdr5-mtdTomato (Benmimoun et al., 2020). We used the glial drivers R54C07-GAL4 (mdr65:GAL4; catalog #50472, Bloomington Drosophila Stock Center) and moody-GAL4 (Schwabe et al., 2005) for knockdown and overexpression experiments. We used the mutant alleles scafF1 (Roussel et al., 2010) and scafF1 (catalog #53101, Bloomington Drosophila Stock Center).

**Nutrient restriction protocol.** Nutrient restriction (NR) experiments were performed as previously described (Contreras et al., 2018). Briefly, 68–72 h after larval hatching (ALH), larvae were transferred to a tube with fly food (Fed) or 1% Agarose in PBS (NR). Tubes were left at 25°C for 24 h or until pupariation. For adult dissection, pupae were transferred to food tubes, and adult fly brains were dissected 1–3 d after eclosion.

**Sample collection for targeted DamID.** Briefly, we crossed tub-GAL80ts; UAS-LT3-NDam and tub-GAL80ts; UAS-LT3-NDam-RiP215 to the respective drivers. We collected embryos for 3 h and let them develop for 8 d at 18°C. Third instar larvae were transferred to 29°C for 1 d to induce the expression of the TaDa construct in either Fed or NR (1% agarose, 1 × PBS) conditions. Fifty to 70 brains of each condition were dissected to extract genomic DNA. Two to five replicates were processed.

**DamID sequencing, processing and data analysis.** The isolated genomic DNA from each cross was digested and amplified to generate libraries for next-generation sequencing as described (Marshall et al., 2016). Libraries were sequenced in an Illumina HiSeq 1500 with at least 8 million reads per library.

Here, we performed cell-type-specific transcriptional analysis of the BBB glia under fed and nutrient restriction (NR) conditions. Among the differentially expressed genes in the SPG, we found scarface (scaf), a member of the serine protease homolog (SPH) family. We show that scaf expression is sensitive to nutrition during larval development. scaf is expressed by the SPG and is required for controlling SPG growth and PG proliferation. Moreover, Scaf is necessary for the proper rate of neurogenesis, and its overexpression enhances NB proliferation and brain growth in animals subjected to NR. Finally, we show that knocking down scaf in SPG affects the resistance of flies to drugs, suggesting that the function of the BBB is impaired when scaf is lost.
TaDa reads were aligned to *Drosophila melanogaster* genome release dm6 and normalized using the damidseq pipeline (Marshall and Brand, 2015). Genome browser views of bedgraph files were displayed as Log2(Dam-PolII/Dam) using Integrative Genomics Viewer software (Robinson et al., 2011). RNA polymerase II (PolII)-bound genes were called using the poli.gene. call R script (Marshall and Brand, 2015; Marshall et al., 2016) using *Drosophila* genome Annotation Release dm6:06. Genes with a false discovery Rate (FDR) lower than 0.01 were considered as bound by PolII. All genes from two datasets were compared using poli.correlation.plot R script (Marshall and Brand, 2015). Differentially bound genes have a difference between Log2(Dam-PolII/Dam) values of 0.3, an FDR < 0.01 in at least one dataset, and a ratio of PolII binding bigger than 2. Student’s test was used to analyze *p* values between Fed and NR replicates.

Gene ontology analysis of differentially expressed genes was performed using the Database for Annotation, Visualization, and Integrated Discovery Bioinformatics Resources 6.8 (Huang et al., 2009a,b). Association networks were constructed using String (Szklarczyk et al., 2019). Plots and heatmaps using custom R scripts are available on request. PolII average Log2(Dam-PolII/Dam) from each replicate were correlated using Pearson’s method and displayed as a heatmap using R gplots library. Volcano plots were generated using the R ggplot2 library. Student’s test *p* values were used to compare every replicate from two conditions (Fed and NR) and the difference of Log2(Dam-PolII/Dam) for each gene in the conditions. Heatmaps were created using the R pheatmap library, and for clustering genes the hclust complete method was used.

**Immunostaining.** Third instar larval brains were fixed in 4% formaldehyde for 20 min and stained as previously described (Wu and Luo, 2006). The following primary antibodies were used: rat anti-N-cadherin (CadN; 1:20; catalog #DN-Ex #8, Developmental Studies Hybridoma Bank), guinea pig anti-Deadpan (Dpn; 1:5000; Andrea Brand Lab), rat anti-Elav (1:20; catalog #7E8A10, Developmental Studies Hybridoma Bank), mouse anti-Fas (1:20; catalog #1D4, Developmental Studies Hybridoma Bank), rabbit anti-phospho-Histone H3 (1:400; catalog #06–570 Merck), rat anti-phospho-Histone H3 (1:500; catalog #HTA28, Abcam), mouse anti-Repo (1:20; catalog #8D12, Developmental Studies Hybridoma Bank), and rabbit anti-Scaf N°2 (1:200; Rousset et al., 2010). DNA was stained using TO-PRO–3 (1:400; catalog #T3605, Invitrogen) or 0.2 μg/mL DAPI. FITC, Cy3, or Cy5 conjugated secondary antibodies were used at a final concentration of 1:200 (Jackson ImmunoResearch).

For BBB permeability assays, larvae were opened at the tail and inverted, exposing the brain. Larvae were incubated with 20 μg/mL Rhodamine-dextran (10,000 MW; #D1863, Invitrogen,) in Schneider’s insect medium for 30 min at room temperature. Brains were washed in 1 × PBS and fixed in 4% formaldehyde in 1 × PBS for 20 min. Samples were washed in PBT (0.3% Triton X-100, 1 × PBS). Brains were dissected and mounted in VECTASHIELD (Vector). For adult animals, males were anesthetized with CO₂, and ~60 nL of 12.5 μg/mL Rhodamine-dextran 10,000 MW was injected in the abdomen using a Nanoject II Auto-Nanoliter Injector (Drummond Scientific). Flies were left to recover 2 h at room temperature, and brains were dissected and fixed.

**Imaging.** Images were acquired using a Leica SP8 or an Olympus FluoView FV1000 scanning confocal microscope. Images of wings, pupae, and adult flies were taken using a Nikon stereomicroscope with a Canon Rebel T2i camera. Images, diograms, and figures were assembled using Fiji, Adobe Photoshop CC, and Adobe Illustrator CC.

**Behavioral assays.** Larval locomotion was performed by placing 10–12 larvae on a 1% agarose plate and recording a video for 1–2 min. Speed was calculating using the ImageJ plugin wrMTrck (Brooks et al., 2016). Adult climbing was performed using a six-tube countercurrent apparatus (Inagaki et al., 2010). Briefly, groups of 15–30 female flies were placed in the first tube of the apparatus and allowed to climb for 30 s. Flies that climbed were transferred to the next tube, and the process was repeated until the last tube was occupied. Then the number of flies in each tube was counted. The climbing index was calculated with the following equation: \( Ci=\frac{(N2+2N3+3N4+4N5+5N6)}{5(N1+N2+N3+N4+N5+N6)}, \) in which \( N1-N6 \) corresponds to the number of flies in each of the six tubes. Eight different groups of flies were tested for each genotype. For fly activity and sleep assay, we used Drosophila Activity Monitors (DAM2, TriKinetics) and loaded adult flies following a previously described protocol (Chiu et al., 2010). Activity monitors were placed in an incubator with a 12:12 h light:dark cycle at 25°C for 6 d; only the last 4 d were used for data collection. The data were analyzed using ShinyR-DAM (Cichewicz and Hirsh, 2018). Sleep was defined as inactivity in a period of 5 min.

For ethanol sedation sensitivity assay, we used groups of 8–10 male flies, placed in an empty vial with a cellulose acetate plug. One ml of ethanol was added to the plug, and the fraction of immobile flies was counted every minute until all flies were sedated (Sandhu et al., 2015). The time at which half the flies were sedated, or sedation time 50 (ST50), was estimated by fitting a sigmoidal curve. For malathion resistance assay, we placed groups of 20 female flies in tubes containing 0.5 g of instant dry food (Carolina Biological Supply) and 2 ml 0.01% malathion in dH₂O. Lethality was counted at 12, 24, and 36 h after malathion exposure.

**Quantifications and data analysis.** Pupal volume was estimated as previously described (Layalle et al., 2008), using the ellipsoid volume formula 4/3π(L/2)(d/2)² (L, length; d, diameter). Time of pupariation analysis was performed as we previously described (González-Itier et al., 2018). Larval and adult brain size was estimated using Imaris 7 (Bitplane) as described (Contreras et al., 2018). Neuropil size was estimated using the area of CadN staining in a maximal intensity Z-projection. The SPG nuclear area was measured by generating a Z-stack maximum intensity projection and calculating the area of each SPG nucleus using Fiji (Li et al., 2017). The number of SPG nuclei was counted in a three-dimensional (3D) reconstruction using the Surpass feature of Imaris 7 (Bitplane). The number of PG nuclei was measured by generating a Z-stack maximum intensity projection and calculating the area of each SPG nucleus using Fiji (Li et al., 2017). The number of SPG nuclei was counted in a three-dimensional (3D) reconstruction using the Surpass feature of Imaris 7 (Bitplane). The number of PG nuclei was measured by generating a Z-stack maximum intensity projection and calculating the area of each SPG nucleus using Fiji (Li et al., 2017). The number of SPG nuclei was counted in a three-dimensional (3D) reconstruction using the Surpass feature of Imaris 7 (Bitplane). The number of PG nuclei was measured by generating a Z-stack maximum intensity projection and calculating the area of each SPG nucleus using Fiji (Li et al., 2017). The number of SPG nuclei was counted in a three-dimensional (3D) reconstruction using the Surpass feature of Imaris 7 (Bitplane). The number of PG nuclei was measured by generating a Z-stack maximum intensity projection and calculating the area of each SPG nucleus using Fiji (Li et al., 2017). The number of SPG nuclei was counted in a three-dimensional (3D) reconstruction using the Surpass feature of Imaris 7 (Bitplane).
analysis were conducted using GraphPad Prism 8. Descriptions of each statistical test used are provided in the figure legends, and p values < 0.05, < 0.01, < 0.001, and < 0.0001 are shown in plots indicated by asterisks, whereas ns means nonsignificant (p > 0.05).

Data availability. DamID-Seq data were deposited in the Gene Expression Omnibus under the accession number GSE145055.

Results

A *Drosophila* brain-sparing model for understanding the response to nutrient restriction

To understand the transcriptional response of the *Drosophila* BBB to NR, we established the brain-sparing model (Cheng et al., 2011; Contreras et al., 2018) under the temperature conditions required for performing PolII targeted TaDa (Southall et al., 2013). Thus, we allowed development at 18°C until 6 d ALH and switched to Fed or NR conditions at 29°C for 1 d (Fig. 1A).

Following this protocol, the consequences of NR during larval development could be observed as a reduction in the size of the animal in both pupal (Fig. 1F–H) and adult (Fig. 1I–L) stages. The average size of the pupa after NR (pupal size reflects the maximum size that a larva reached before pupariation) corresponded to 34.51% of Fed animals (Fig. 1H). We found that under this condition, larval brains continued growing on NR but not at the same rate as their Fed counterparts (Fig. 1B–D). For the adult brain, we observed a significant reduction in the brain size of animals that were subjected to NR during larval development (Fig. 1M,N). The larval and adult brains of NR animals reached 67.08% and 60.62% of the size of the Fed brains, respectively (Fig. 1E,O). Therefore, we were able to replicate the brain-sparing effect under the conditions required to perform TaDa as the effect of NR over the growth of the whole animal was greater compared with the brain (body reduction to almost 35% compared with brain reduction to a 67% of the Fed animal), reflecting the preferential growth of the nervous system during starvation (Cheng et al., 2011).

Transcriptional profiling of neuroblasts and blood–brain barrier glial cells under nutrient restriction

To determine the transcriptional response of neuroblasts and the BBB to NR, we profiled *in vivo*, and in a cell-type-specific manner, the binding of RNA Polymerase II (Rpi215/PolII) using the TaDa technique. Briefly, TaDa uses the GAL4 system (Brand and Perrimon, 1993) to drive expression of a DNA-binding or chromatin factor fused to the *Escherichia coli* Dam methylase (Southall et al., 2013; Augey and Southall, 2016; van den Ameele et al., 2019; Extended Data Fig. 2-1A,B). We used two GAL4 drivers to target NDam-PolII expression in the SPG (R54C07-GAL4, also referred to as mdr65-GAL4) or the PG and SPG together (SG, R71C08-GAL4, sema5c-GAL4; Fig. 2B–B‴ for driver expression; Extended Data Fig. 2-1D–G for PolII binding in BBB genes), and wor-GAL4 for profiling NDam-PolII binding in all NBs (Extended Data Fig. 2-1, Fig. 2A). We performed TaDa under two nutritional conditions, Fed and NR, according to our established protocol (Fig. 1A).

Meta-analysis of our PolII occupancy data showed the characteristic profile of PolII binding across a gene (Southall et al., 2013; Extended Data Fig. 2-1C). Our TaDa replicates also correlated with each other as expected, with higher correlation among libraries of the same cell type and lower when NBs were compared with glial cells (Fig. 2C). Next, we analyzed the data and selected genes that were differentially bound by PolII between Fed and NR conditions in each dataset, considering genes with an FDR lower than 0.01, a PolII binding ratio higher than 2, and a difference higher than 0.3 (Fig. 2D–F, Extended Data Fig. 2-2).

In the NB comparison, we found 14 genes upregulated and 246 genes downregulated after NR (Fig. 2D,G,H). In the BBB, 12 genes were upregulated and 47 downregulated in SPG (Fig. 2E,I), whereas in the SG datasets, PolII TaDa showed 44 genes upregulated and 246 genes downregulated in the NR compared with Fed condition (Fig. 2F,J). Importantly, our data showed that PolII binding to genes associated to the drivers used (wor, mdr65 and sema5c) was not affected by NR in the corresponding experiment.

The brain-sparing model in *Drosophila* was previously shown to maintain NB proliferation on NR (Cheng et al., 2011). Our NB PolII TaDa gene ontology analysis showed that categories involved in ribosome, mitochondria, and oxidative phosphorylation metabolism were significantly enriched in downregulated genes (Fig. 2G, Extended Data Fig. 2-1H). This suggests that energy metabolism and protein synthesis were affected in neural stem cells during NR. However, we found no significant differences in the growth and proliferation of NBs between Fed and NR conditions (Fig. 3A,B), confirming that NB division and neurogenesis were not disrupted by starvation during third instar larval development (Cheng et al., 2011). In addition to this, we analyzed the neuropil of larval brains, finding that the relative size of the neuropil was increased in animals under NR (Fig. 3C–E), suggesting an effect of starvation over neuropil and cell body proportions.

In our BBB TaDa datasets, we found a group of proteins associated with the JNK and Janus kinase/signal transducer and activator of transcription (JAK/STAT) signaling pathways that were downregulated after NR (Extended Data Fig. 2-1I). These included *Matrix metalloproteinase 1* (*Mmp1*; Uhlirova and Bohmann, 2006), *scarface* (*scfa*; Rousset et al., 2010; Srivastava and Dong, 2015), *eiger* (*egr*, TNF homolog), *TGF-β activated kinase 1* (*Tak1*) and the JAK/STAT ligands *unpaired 1* (*upd1*), *unpaired 2* (*upd2*), and *unpaired 3* (*upd3*; Fig. 2I,J, Extended Data Fig. 2-1I). Therefore, we analyzed whether the JAK/STAT and JNK pathways were active in the larval BBB and if their activity was modulated by larval nutrition. We used the 10xSTAT92E-GFP (Bach et al., 2007) and *TRE-REP* (Chatterjee and Bohmann, 2012) reporters to evaluate JAK/STAT and JNK pathway activity, respectively. As expected, both reporters were expressed in the BBB during larval development; however, the expression of these reporters did not change after nutrient restriction (Fig. 3F–I), suggesting that these signaling pathways in the BBB are insensitive (not responding) to the nutritional status of the animal.

Nutrient restriction affects the development of the *Drosophila* blood–brain barrier

Because NB growth and proliferation were not affected by NR, we hypothesized that the larval BBB might adapt to starvation, being the first layer of the brain to sense and respond to a decrease in nutrients in the larval hemolymph.

The two types of glial cells that form the BBB, the SPG and PG have different mechanisms of growth. SPG grow by endoreplication resulting in polyploid cells (Unhavaithaya and Orr-Weaver, 2012), whereas PG proliferate, undergoing mitosis (Pereanu et al., 2005; Awasaki et al., 2008; Avet-Rochex et al., 2012). SPG endoreplication takes place by either endocycle or endomitosis (Unhavaithaya and Orr-Weaver, 2012; Øvrebø and
Edgar, 2018). Therefore, we checked the SPG for endomitosis by counting the number of SPG nuclei in third instar larval brains subjected to NR. We found a significant reduction in the number of nuclei in SPG (Fig. 4A) and a significant decrease in the percentage of multinucleated SPG (Fig. 4B). Similarly, the distribution of SPG with multiple nuclei changed significantly from a median of two in Fed animals to one after NR (Fig. 4C,D). Given that the size of the nucleus correlates highly with cellular size and...
Figure 2. Targeted DamID in neuroblasts and glial cells of the blood–brain barrier after nutrient restriction. A, Schematic representation of the different cell types in the larval brain. CG, Cortex glia; GMC, ganglion mother cells; N, neuron; NG, neuroplil glia (Extended Data Figure 2-1, schematic representation). B, B’, B’’, Confocal images of larval brain expressing mCD8-GFP in superficial glia (PG and SPG, sema5c-GAL4, UAS-mCD8-GFP, green and gray) and mtdTomato in SPG (mdr65-mtdTomato, red and gray). Arrowheads show the overlap of the green and red signals. Scale bar, 20 μm. PolII binding in BBB marker genes shown in Extended Data Figure 2-1. C, Heatmap diagram showing correlation of all Fed and NR TaDa libraries. D–F, Volcano plots showing PolII bound genes in NR and Fed conditions in (D) NBs, (E) SPG, and (F) surface glia (SPG+PG). Genes differentially bound by PolII between Fed and NR conditions are highlighted in red. G, Graph showing the gene ontology categories enriched in genes downregulated in NBs by NR (Extended Data Figure 2-1). H–J, Heatmaps showing the PolII binding of differentially expressed genes in the TaDa replicates of (H) NBs, (I) SPG, and (J) SG. Tables of PolII binding in all genes are shown in Extended Data Figure 2-2.
DNA content (Frawley and Orr-Weaver, 2015), nuclear size was measured as an indicator of the SPG endocycle. We found a significant reduction in average nuclear size after NR (Fig. 4E). We also counted the number of PG on NR to check PG proliferation and observed a reduction in PG proliferation (Fig. 4F). These results confirmed a major effect of nutrition on the BBB glial cells—reduction in SPG endoreplication and a decrease in PG proliferation after food deprivation.

A severe reduction in SPG endoreplication affects the maintenance of the subperineurial septate junctions, allowing paracellular diffusion inside the larval brain (Unhavaithaya and Orr-Weaver, 2012; Von Stetina et al., 2018). Therefore, we checked the integrity of SPG septate junctions after nutrient restriction using an endogenously tagged version of the septate junction structural protein \textit{lachesin} (lac::GFP; Morin et al., 2001). We found no major disruptions in the continuity of the septate junctions in larval brains under NR conditions (Fig. 4G–H). To test further the integrity of the BBB, we assessed BBB permeability by incubating larval brains with rhodamine-labeled dextran (10 kDa) and assayed for penetration inside the ventral nerve cord. Rhodamine-dextran did not enter the CNS and remained on top of the subperineurial layer, marked by \textit{mdr65-GAL4, UAS-mCD8-GFP} (Fig. 4I–J). These results suggest that the integrity of the larval BBB is not disrupted by larval NR, despite the reduction in BBB growth.

Figure 3. Nutrient restriction does not affect neuroblast proliferation and pathway activation in the BBB. A, NB diameter during mitosis (green dots) and interphase (magenta dots) of \textit{w^{1118}} larval brains in Fed and NR conditions. \( n = 9 \) and 7 brain lobes, respectively. Two-way ANOVA and Bonferroni’s multiple-comparisons test were performed. ns, Nonsignificant. B, Graph showing the NB mitotic index of \textit{w^{1118}} larval brains under Fed and NR conditions. \( n = 10 \) and 8 brain lobes, respectively. Unpaired Student’s \( t \) test was performed. ns, Nonsignificant. C, D, Larval CNSs immunostained for CadN (neuropil marker in green) and DNA (red) under (C) Fed or (D) NR conditions. E, Plot showing the quantification of the relative size of the neuropil (neuropil/whole brain) from C, D. \( n = 13 \) and 12 CNSs, respectively. Unpaired Student’s \( t \) test was performed, \(* * * p < 0.0001\). F–I, Ventral nerve cords of (F–G) JAK–STAT (10xSTAT92E-GFP) and (H–I) JNK (TRE-RFP) pathway reporters under (F, F’, H, H’) Fed and (G, G’, I, I’) NR conditions. F–G’, Immunofluorescences for GFP (green and gray), Repo (red) and Elav (blue). H–I, Immunofluorescences for Repo (green) and RFP (red and gray). Scale bars: 20 \( \mu \)m.
Scarface is expressed in subperineurial glia and is regulated by nutrient availability. Because the growth of the BBB is affected by NR, we focused on BBB genes that were differentially expressed as indicated by our TaDa analysis. Of those, we centered our interest on scarface (scaf), one of the most downregulated genes during NR in SPG (Figs. 2I, 5A). Scaf belongs to the SPH family (Ross et al., 2003; Bonin and Mann, 2004; Rousset et al., 2010; Sorrosal et al., 2010). Serine protease homologs resemble proteases but lack the amino acids required for enzymatic activity. Scaf controls epithelial polarity and morphogenesis during embryonic development and thorax formation, acting as a downstream target of the JNK...
Figure 5. Scarface is a nutrient-sensitive gene expressed in the Drosophila blood–brain barrier. **A**, Genome view of the scaf locus and the binding of PolII in SPG TaDa in Fed (blue) and NR (red) conditions. Note the reduction on PolII binding in NR compared with Fed. **B–E″**, Immunostaining of scaf::GFP (scafΔ), in green) larval brain lobes (optical section at the BBB level) under (**B–B″**) Fed and (**C–C″**) NR conditions stained for Repo (glial marker in red). **B″, C″**, are color-coded images of Scaf::GFP signal. **D–E″**, Immunostaining of larval brains under (**D–D″**) Fed and (**E–E″**) NR conditions stained for Scaf (green) and Repo (red). **D″, E″** are color-coded images of anti-Scaf antibody signal. **F, G**, Graph showing the quantification of the Scaf signal in the BBB using (**F**) Scaf::GFP reporter (n = 7 and 8 brains) or (**G**) anti-Scaf antibody (n = 11 brains). Unpaired Student’s t test was used. **H**, Confocal images of a larval brain stained with anti-Scaf (green) and Dpn (NB marker in red). **I, I″**, Larval brain expressing mCD8::GFP in SPG (mdr65-Gal4, UAS-mCD8-GFP, in red) and stained for Repo (glial marker in blue) and Scaf (green or gray). **J, J″**, Cross section of larval BBB of mdr65-Gal4, UAS-mCD8::GFP animals stained against Repo (blue), Scaf (green or gray) and mCD8::GFP (red or gray). **K–K″**, 3D reconstruction of the larval BBB of mdr65-Gal4, UAS-mCD8::GFP animals stained against Repo (blue), Scaf (green), and mCD8::GFP (red). **L–L″**, Representative confocal image of a cross section of a larval brain stained against Scaf (blue and gray), sema5-Gal4, UAS-mCD8::RFP (red) and LanA::GFP (green and gray) as a neural lamella marker. Scale bars, **B–E″**, **H–J″**, 20 μm; **L–L″**, 0 μm. **M**, Graphic representation of the regulation of scaf expression in the blood–brain barrier by nutrition.
pathway, and it has been reported to be a secreted protein (Rouset et al., 2010; Sorrosal et al., 2010; Srivastava and Dong, 2015; Kushnir et al., 2017).

To validate that scaf was differentially expressed on NR, we made use of a gene trap line (scafRBsu allele, Scaf::GFP; Bonin and Mann, 2004) and an antibody against Scaf (anti-Scaf; Rouset et al., 2010). Although, the scafRBsu is a semilethal allele, when used in heterozygosis it is a reliable tool for analyzing scaf expression. We observed Scaf expression in glial cells at the surface of the larval brain, corresponding to the BBB (Fig. 5B–B′, D–D′). As predicted by TaDa, the levels of Scaf were reduced after NR (Fig. 5C–C′, E–E′). Intensity quantification of the Scaf signal showed a significant reduction of an average of 47.56% and 59.37% of the Scaf intensity of Fed animals (Scaf::GFP and anti-Scaf, respectively, Fig. 5F,G).

As predicted by our SPG TaDa analysis, Scaf colocalised with an SPG membrane marker (mdr65-GAL4, UAS-mCD8-GFP, Fig. 5I–J′) at the surface of the brain, but not with NBs, labelled by Dpn (Fig. 5H). Interestingly, the Scaf signal appeared also on top of the SPG membrane marker in a 3D reconstruction (Fig. 5K–K′). To confirm this, we used an endogenously tagged Laminin A (LanA::GFP) line to label the brain extracellular matrix (neural lamella) and an SG membrane marker (sema3c-GAL4, UAS-mCD8-RFP). Because the thickness of the BBB is ~2 µm, we observed that the Scaf signal overlapped with LanA and most of SG membrane signals, thus the entire BBB (Fig. 5L–L′). These results showed that during third instar larval development scaf is expressed by SPG in a nutrition-sensitive fashion (Fig. 5M).

Scaf decreases the growth of the brain–brain barrier during larval development

As SPG expression of Scaf is sensitive to nutrition, we aimed to assess its function in the BBB during normal development. To accomplish this, we performed RNAi-mediated knockdown of scaf only in SPG using the mdr65-GAL4 driver. We tested two different RNAi lines, a short hairpin (shScafRNAi) and a long hairpin (lhScafRNAi), finding that both were able to significantly reduce the levels of Scaf in the BBB (Fig. 6A–D). Because shScafRNAi gave the most efficient knockdown (29.63% of control Scaf levels), we used this line in most of our experiments.

Knockdown of scaf in SPG produced normal larvae, which grew and pupariated as control animals and emerged into fertile adult flies (Fig. 6E–H). We checked the development of the BBB at late third instar larval stage, observing no morphologic defects (Fig. 6I,J). Thus, we analyzed endomitosis of SPG, finding a significant increase in the number of SPG nuclei after scaf knockdown (Fig. 6K) but without a significant change in the percentage of multinucleated SPG per brain lobe (Fig. 6L). This suggested that on scaf knockdown, the number of multinucleated SPG is similar, but they contain more nuclei than in control animals. This was confirmed by analyzing the distribution of SPG according to the number of their nuclei, showing an increase in the median from one nucleus in control animals to two nuclei in shScafRNAi brains (Fig. 6M,N).

Next, we checked the nuclear size in SPG, observing a significant increase on scaf knockdown (Fig. 6O), which suggests an increase in SPG endocycle. We also found a significant increase in the number of PG in shScafRNAi animals (Fig. 6P). We confirmed these results using the lhScafRNAi strain (Extended Data Fig. 6-1). Next, we assessed whether permeability of the BBB was altered in scaf knockdown larval brains using the Rhodamine-dextran assay described before. We did not detect the colorant inside the CNS (Fig. 6Q,R), ratifying that a reduction in the levels of Scaf increases the growth of the subperineurial layer as well as the number of PG without affecting the BBB permeability.

Given that scaf knockdown augmented the growth of the BBB, increasing the levels of Scaf should produce the opposite effect. To perform gain of function experiments in SPG during development, we used a line (UAS-Scaf) that rescues scaf mutant embryonic lethality (Rouset et al., 2010). Overexpression of Scaf generated normal and viable animals, without affecting larval growth under Fed or NR conditions. However, the timing of pupariation was significantly accelerated (see Fig. 7A–F). In the BBB, we found that the average number of SPG nuclei did not change on Scaf overexpression (Fig. 7G); however, the percentage of multinucleated SPG in a brain lobe was significantly decreased (Fig. 7H). We also observed an increase in the multinucleated fraction of SPG but without a change in the overall distribution (Fig. 7J). Interestingly, the size of the SPG nuclei was significantly reduced by Scaf overexpression (Fig. 7K). These results showed that increasing the levels of Scaf has a mild effect on SPG endoreplication. In the perineurial layer, the number of PG was not affected by Scaf overexpression (Fig. 7L), suggesting that there was no impact on PG proliferation. Our data infer a role for Scaf in diminishing the growth of the BBB through decreasing the levels of the SPG endocycle and endomitosis during development under normal nutritional conditions.

Blood–brain barrier–derived scraf increases neurogenesis in the larval brain

Nutrient restriction reduced the levels of Scaf in the BBB and also produced animals with smaller brains than the control Fed condition. We assessed whether Scaf function at the BBB has a nonautonomous effect on neurogenesis. Thus, we knocked down scaf in SPG and checked the mitotic index of larval central brain NBs, observing a significant decrease in the fraction of NBs in mitosis (Fig. 8A–C). To confirm this result, we used a scaf allelic combination of scaf+/−, a deletion resulting in a strong lethal allele (Rouset et al., 2010), with scaf+/+; sema5c−/−, a semilethal allele generated by a MiMIC insertion (Venken et al., 2011), producing a truncated Scaf protein (that retains the antibody epitope). This, scaf+/−; sema5c−/−, allelic combination is a hypomorphic condition that is viable and healthy during larval development but lethal at late pupal stage. Similar to scaf knockdown experiments, we found that scaf mutant brains had a significantly lower NB mitotic index than heterozygous animals (scaf+/−; Fig. 8D). In the same way, the hypomorphic allelic combination, scaf+/−; scaf−/−, also showed a decreased NB mitotic index compared with the heterozygous control (scaf+/+; Fig. 8E,G). Because scaf in the BBB was necessary for proper NB proliferation, we conducted a rescue experiment to determine whether the expression of Scaf only in SPG was sufficient to maintain the normal NB mitotic index in scaf mutant animals. As seen in Figure 8E, the overexpression of scaf was not able to rescue NB mitotic index in a scaf mutant background. Thus, Scaf from SPG is not sufficient to maintain NB proliferation in the mutant (Fig. 8E). Additionally, the knockdown of scaf only in SPG did not have a major impact on the overall size of the brain (Fig. 8F). These results suggest that Scaf is necessary in the BBB to maintain the rate of larval neurogenesis, but Scaf in other cell types is also required for NB proliferation.

As Scaf downregulation affects the rate of NB proliferation under normal nutritional conditions, we analyzed whether Scaf overexpression in SPG could enhance neurogenesis in animals subjected to NR. We found that in NR conditions, expression of Scaf in SPG resulted in a 13.78% increase in the size of the adult.
Scarface is necessary for controlling the growth of the blood–brain barrier. A–C, Immunostaining of larval brains of mdr65-GAL4, UAS-DlgA:EGFP animals crossed to (A, A’) w^{1118} (control), (B, B’) UAS-shScafRNAi, and (C, C’) UAS-lhScafRNAi, and stained for Repo (blue), Scaf (red and gray) and GFP (green). Scale bars: 50 μm. (D) Plot showing the quantification of scaf knockdown. n = 6, 7, 6 brains, respectively. One-way ANOVA and Dunnett’s multiple comparisons test were performed. E, F, Images of pupae of mdr65-GAL4 crossed to (E) w^{1118} (control) and (F) UAS-shScafRNAi. G, Plot showing the pupal volume for each group. n = 45 and 52 pupae, respectively. Mann–Whitney test was performed. H, Graph showing the timing of pupariation of mdr65-GAL4 crossed to w^{1118} (Ctrl), dark or UAS-shScafRNAi (red). n = 6 groups of ~35–40 larvae for each condition. I, J, Larval CNS of mdr65-GAL4, UAS-DlgA:EGFP animals crossed to (I) w^{1118} (control) and (J) UAS-shScafRNAi. UAS-DlgA:EGFP labels SPG nuclei and septate junction. Scale bars: 100 μm. K–N, Analysis of larval SPG endomitosis. mdr65-GAL4, UAS-DlgA:EGFP animals were crossed to w^{1118} (control) and scaf knockdown (UAS-shScafRNAi). K, Graph showing the average number of SPG nuclei per brain lobe. n = 43 and 46 brain lobes. Unpaired Student’s t test was performed. L, Plot showing the percentage of multinucleated SPG (2 or more nuclei) per brain lobe. n = 43 and 56 brain lobes. Mann–Whitney test was done. M, Plot showing the number of nuclei in each SPG analyzed, median (black line) and interquartile range are shown. Mann–Whitney test was done. n = 230 (Ctrl) 320 (shScafRNAi). SPG. N, Histogram showing the relative distribution (percentage) of SPG according to the number of nuclei. O, Graph showing the average size of the SPG nucleus of larval brains. mdr65-GAL4, UAS-DlgA:EGFP animals were crossed to w^{1118} (control) and UAS-shScafRNAi. ***, p < 0.001; **, p < 0.01; *, p < 0.05; ns, not significant; n = 20–60 lobi per condition.
brain in comparison to control NR animals (Fig. 9A–C). Furthermore, the larval central brain NB mitotic index was also increased on Scaf overexpression in animals under NR (Fig. 9D). For the BBB, Scaf overexpression did not prevent the reduction in the growth of the SPG or PG layers during NR (Fig. 9E–J). Importantly, we did not observe a difference in brain growth during normal Fed conditions on Scaf overexpression (Fig. 9K–M), showing that Scaf overexpression enhances brain growth only under starvation. In the case of the growth of other organs, reexpression of Scaf in the BBB during NR, induced a small increase in the size of adult wings (4.76% of increase compared with control animals; Fig. 9N–R).

Together, these results demonstrate that Scaf, expressed by the SPG layer, is required for maintaining the rate of neurogenesis. This highlights a nonautonomous role for Scaf in coordinating neurogenesis and the growth of the BBB. Additionally, Scaf reexpression is able to promote brain growth during nutrient deprivation.

Drug sensitivity is increased on scarface loss in the blood–brain barrier

We found that Scaf modulates the growth of the BBB and has a role in maintaining the rate of NB proliferation in the central brain. However, the loss of scarface seemed to have only a mild effect in the development of the CNS. Therefore, we wonder whether the phenotypes observed after scarface knockdown during larval development has an impact on the function of the BBB and the CNS of the adult animal.

With this goal, we analyzed locomotor activity in larval and adult animals on scarface knockdown in SPG. We found no significant differences in the larval crawling speed (average and maximum) of control and scarface knockdown animals (Fig. 10A,B). Similar results were obtained when we assessed climbing activity in adult flies (Fig. 10C). As BBB function has been associated with circadian activity rhythm and sleep (Artiushin et al., 2018; Zhang et al., 2018; Cuddapah et al., 2019), we analyzed locomotor activity and sleep in adult flies using DAM2. We found no differences in total activity or activity profile between control and scarface knockdown groups (Fig. 10D,F). Total sleep was not affected in scarface knockdown animals either (note that driver-alone sleep did not differ from scarface knockdown group in Fig. 10E), however, the sleep profile of these animals showed less sleep than both control groups a few hours before the end of the light cycle (Fig. 10G).

One of the major roles of the BBB is to control the influx and efflux of molecules across the brain, and it is known that the sensitivity to different drugs is altered in mutant animals with defects in BBB permeability (Bainton et al., 2005; Mayer et al., 2009). The BBB permeability is controlled by preventing paracellular diffusion (by septate junctions) and by the active extrusion of lipophilic molecules into the hemolymph. Hence, we assessed the sensitivity to ethanol and malathion as a manner to test for BBB function defects. Ethanol, although attractive for the flies, produces miscoordination and loss of equilibrium in flies (pass out), and its tolerance is regulated in the BBB by the action of a kinase anchoring protein in PG (Parkhurst et al., 2018). Malathion is an organophosphate insecticide that blocks acetylcholinesterase, inducing hyperexcitability as acetylcholine is the main excitatory neurotransmitter in the fly brain. In Drosophila, malathion sensitivity is increased by the loss of the ATP-binding cassette transporter mdr65 (Sun et al., 2017), which is necessary for BBB xenobiotic eﬄux (Mayer et al., 2009; Hindle et al., 2017). Similarly, in vertebrates malathion can cross and affect the structure of the BBB (Balbuena et al., 2010, 2011). We found that sensitivity to a single exposure of ethanol was significantly enhanced in animals with scarface knockdown in SPG (Fig. 11A,B), and animals exposed to malathion, showed a significant increase in lethality compared with control animals (Fig. 11C). Interestingly, adult flies that were subjected to larval NR, in which scarface is downregulated in the BBB, also showed an increased sensitivity to ethanol (Fig. 11D,E). Importantly, the sensitivity to drugs after scarface knockdown could not be attributed to an effect over the excretion of drugs because scarface RNAi was not expressed in the adult gut or renal (Malpighian) tubules (Fig. 11F,G). mdr65-GAL4 expression. Additionally, paracellular permeability of the adult BBB was not affected by scarface knockdown (Fig. 11H,I), suggesting that extrusion of xenobiotic agents may be affected by the loss of scarface in the BBB.

These results show that reducing the expression of scarface in the BBB, which has an effect over BBB growth, increases the sensitivity of the CNS to exogenous agents, supporting a role of Scarface in the regulation of the function of the BBB.

Discussion

The adaptive response of animals to periods of famine is essential for species survival. In the CNS, the brain is preferentially protected over other tissues during starvation. Here, we analyze the transcriptional response of Drosophila melanogaster NBs and BBB glial cells to NR during larval development using TaDa. Overall, we found differentially expressed genes in NBs, subperineurial and surface glial datasets. We also found that in our starvation model, the major effect of NR was on the growth of the BBB. Thus, we focused on the function of a gene that is highly regulated by NR in the BBB, the SPH scarface (scarface). We determined that scarface is expressed in the SPG, and its expression is modulated by the animal’s nutritional state. Tissue-specific knockdown of scarface showed that it is required to restrict the growth of the BBB and to maintain a proper rate of NB proliferation. In accordance with this, reintroducing Scaf expression in SPG enhances neurogenesis in animals subjected to nutritional stress. Finally, we showed that scarface knockdown in the BBB increases sensitivity to drugs in adult animals, suggesting defects in the control of the influx/efflux of molecules across the BBB.

A genetic response of the CNS to food scarcity

The maintenance of homeostasis during inanition permits animal survival and developmental progression. Among the different organs, the CNS, which is extremely sensitive to stress conditions, is protected from systemic nutrient reduction. The brain-sparing model is good example of this mechanism of protection during development. In Drosophila, neural stem cell proliferation depends on nutrition at early larval stages (Britton and
but it is independent of nutrition through the end of third instar larval stage (after acquiring the minimal viable weight; Cheng et al., 2011). Cheng et al. (2011) said that NBs maintain their proliferation rate by the action of Jelly belly/Anaplastic lymphoma kinase that activates the phosphatidylinositol 3-kinase pathway. However, these pathways do not respond to starvation, nor are they modulated by the nutritional state of the animal.
This raised the question of whether neural stem cells respond to, or are insensitive to, nutrient restriction. It is plausible that molecular adaptation to nutritional stress occurs in other cell types, such as glial cells, or in other organs.

Our TaDa data showed that there are transcriptional changes in NBs during NR that could help to maintain proliferation and neurogenesis. We postulate that the decrease in the expression of genes associated with ribosome and mitochondrial metabolism could help to maintain the rate of NB proliferation on nutrient restriction. During pupal development, oxidative phosphorylation is required for NBs to reduce growth and exit the cell cycle (Homem et al., 2014; van den Ameele and Brand, 2019), but it is also necessary for larval NBs and NB-derived tumor proliferation (Genovese et al., 2019; van den Ameele and Brand, 2019). Hence, modulating mitochondrial metabolism could be an NB response for maintaining neurogenesis during nutrient restriction. However, the actual contribution of this metabolic regulation to neurogenesis is unknown.

On the other hand, it is well established that other organs contribute to the brain-sparing phenomenon. For instance, on starvation polyploid tissues such as salivary glands and body fat stop growing (Cheng et al., 2011), and body-fat-derived glycogen maintains trehalose levels in the hemolymph (Yamada et al., 2018). Therefore, nutrient stores are mostly available for the CNS to continue growing. The role of neurons and glial cells in this process is less understood. In the case of hypoxia, NB proliferation is protected by the formation of lipid droplets in glial cells (Bailey et al., 2015), supporting a protective role of glial cells.

In recent work, the Simon Sprecher group characterized the response of the first instar larval brain to starvation using single-cell sequencing. Although, under these conditions neurogenesis is completely blocked, the group found a transcriptional response in glial cells to promote lipid catabolism (Brunet Avalos et al., 2019). The glia forming the BBB are likely to be the first sensors of nutrient reduction in the hemolymph. The presence of different macromolecule transporters in the BBB (DeSalvo et al., 2014; Volkenhoff et al., 2015; Galagovsky et al., 2018) and their capacity to store neutral lipids (Bailey et al., 2015) supports the role of the BBB as the nutritional gatekeeper of the brain.

In our analysis, we observed that SPG showed an important transcriptional response to nutrient restriction. Among the differentially expressed genes, we found secreted proteins and components of the JNK and JAK/STAT signaling that are well-

Figure 8. Scarface modulates neurogenesis in the larval brain. A, B, Immunostaining of larval brains of mdr65-GAL4 crossed to (A) w1118 (control) and (B) UAS-shScafRNAi, stained for the NB marker Dpn (red) and phospho-Histone H3 (pH3 in green). Scale bar, 20 μm. C, Graph showing the NB mitotic index (percentage of pH3-positive NB) of control and shScaf knockdown brains. n = 34 and 35 brain lobes, respectively. Unpaired Student’s t test was performed. D, Graph showing the NB mitotic index of shScaf (heterozygous) and shScaf (homozygous). n = 12 and 11 brain lobes, respectively. Unpaired Student’s t test was performed. E, NB mitotic index of shScaf, mdr65-GAL4 (heterozygous); shScaf, mdr65-GAL4, and shScaf, mdr65-GAL4, UAS-Scaf:GFP animals. n = 26, 27, and 27 brain lobes, respectively. One-way ANOVA and Dunnett’s multiple-comparisons test were performed. F, Plot of the size of the adult brain of mdr65-GAL4 crossed to w1118 (control) and UAS-shScafRNAi. n = 39 and 41 adult brains respectively. Unpaired Student’s t test was performed. *p < 0.05, ****p < 0.0001. ns, Nonsignificant. G, Working model of Scaf function in the Drosophila blood–brain barrier. As a response to nutrient availability, the subperineurial glial cells in the BBB express Scaf. Scaf slows down the growth of SPG and diminishes the proliferation of PG but at the same time promotes NB proliferation.
Expression of Scarface in subperineurial glia increases neurogenesis during nutrient restriction. A, B, Adult brains stained for CadN (green) of mdr65-GAL4 crossed to (A) w1118 (control) and (B) UAS-Scaf::GFP (Scaf OE) under NR. Scale bar, 100 μm. C, Quantification of the adult brain area. mdr65-GAL4 animals were crossed to w1118 (control) and UAS-Scaf::GFP (Scaf OE) under NR. n = 38 and 37 adult brains, respectively. Unpaired Student’s t test was performed. D, Graph of the NB mitotic index of mdr65-GAL4 animals crossed to w1118 (control) and UAS-Scaf::GFP (Scaf OE) larval brains under NR. n = 11 and 13 brain lobes, respectively. Unpaired Student’s t test was performed. E–H, Analysis of larval SPG endomitosis of mdr65-GAL4, UAS-DlgA::GFP animals crossed to w1118 (control) and scaf overexpression (UAS-Scaf::GFP, Scaf OE) under NR. E, Graph showing the average number of SPG nuclei per brain lobe. n = 28 and 26 brain lobes. Unpaired Student’s t test was performed. F, Plot showing the percentage of multinucleated SPG (2 or more nuclei) per brain lobe. n = 28 and 26 brain lobes. Mann–Whitney test was used. G, Plot showing the number of nuclei in each SPG analyzed, median (black line) and interquartile range are shown. Mann–Whitney test was performed. n = 182 (Ctrl) and 183 (Scaf OE) SPG. H, Histogram depicting the relative distribution (percentage) of SPG according to the number of nuclei. J, Graph showing the average size of the SPG nucleus of larval brains of moody-GAL4, UAS-GFP.nls animals crossed to w1118 (control) and Scaf overexpression (UAS-Scaf::GFP, Scaf OE) under NR. n = 30 and 26 brain lobes. Unpaired Student’s t test was used. K, L, Immunostaining against CadN of adult female brains of mdr65-GAL4 crossed to (K) w1118 (Ctrl) or (L) UAS-Scaf::GFP (Scaf OE). Scale bar, 100 μm. M, Plot showing the size of the adult brain of mdr65-GAL4 crossed to w1118 (control) and UAS-Scaf::GFP (Scaf OE) under NR, n = 16 and 13 adult brains, respectively. Unpaired Student’s t test was performed. N–Q, Wings of male animals of mdr65-GAL4 crossed to (N, O) w1118 (Ctrl) or (P, Q) UAS-Scaf::GFP (Scaf OE) under NR conditions. Scale bar, 1 mm. R, Plot showing the quantification of the size of the wings for each conditions. n = 29, 30, 30, and 30 wings, respectively. Two-way ANOVA and Tukey’s multiple-comparison tests were done. *p < 0.05, ****p < 0.0001. ns, Nonsignificant.
characterized stress-response pathways. Recently, Prasad and Hens (2018) analyzed the transcriptional response of the adult brain to starvation and sugar absence, finding many secreted proteins, including serine protease homologs and Scaf secreted from a group of neurons (i.e., scarface neurons in the adult brain), which promote feeding behavior. Thus, it is plausible that changes in the expression of secreted proteins, including Scaf, are part of a signature of the brain response to nutrient restriction. Surprisingly, we found that the activity of the JNK and JAK/STAT pathways in the BBB was not affected by nutrient restriction. It would be interesting to evaluate whether during nutrient restriction a decrease in components of these pathways could affect these signaling pathways to reduce the growth of the BBB.

Scarfase coordinates blood–brain barrier growth with neurogenesis in a cell nonautonomous manner

From our TaDa analysis, we selected Scaf as a good candidate to mediate BBB adaptation to nutrient restriction. Scaf belongs to the SPH family, which lacks amidase activity (Ross et al., 2003; Bonin and Mann, 2004). Scaf was previously described to be a transcriptional target of the JNK pathway, which controls epithelial polarity and morphogenesis during dorsal closure (Rousset et al., 2010; Sorrosal et al., 2010; Kushnir et al., 2017) and thoracic development (Srivastava and Dong, 2015). We showed that Scaf expression in SPG depends on nutrition, and one of its roles is to reduce SPG growth by endocycle/endomitosis as well as PG proliferation. Although, the effect of sacf loss is not strong enough to affect animal development to adulthood, sensitivity to drugs that cross the BBB is enhanced, revealing a physiological consequence to this defect in the growth of the BBB. We postulate that loss of sacf could affect transcellular diffusion of organic molecules across the BBB or the function of efflux transporters that eliminate xenobiotics from the brain (Mayer et al., 2009; Hindle et al., 2017; Sun et al., 2017).

How Scaf regulates the growth of the BBB is unknown. As Scaf has been shown to be a secreted protein (Rousset et al., 2010; Sorrosal et al., 2010), we propose that Scaf from SPG modulates the activity of growth factors for both SPG and PG. The balance between SPG endocycle and endomitosis is regulated by the Notch pathway (Von Stetina et al., 2018). This mechanism seems to be an autocrine signaling because δ ligand is also necessary in SPG for promoting endocycle over endomitosis (Von Stetina et al., 2018). On the other hand, PG proliferates in response to insulin and FGF signaling (Franzdóttir et al., 2009; Avet-Rochex et al., 2012). Therefore, it is possible that Scaf could affect these signaling pathways to reduce the growth of the BBB. Given that Scaf represses the JNK pathway during

Figure 10. Locomotion activity is not altered on sacf knockdown in the blood–brain barrier. A–C, Locomotion assays of mdr65-GAL4 animals crossed to w1118 (control) and shScafRNAi. Graphs showing the larval (A) average and (B) maximum speed. n = 67 and 65 larvae. C, Plot of the adult climbing index using a countercurrent apparatus. n = 8 groups of ~25 adult flies for each genotype. Unpaired Student’s t-test was performed. D, Activity measurements using the DAM2 system. Plot shows the daily number of counts per day of single adult flies of w1118 crossed to UAS-shScafRNAi (dark dots, control), mdr65-GAL4 crossed to w1118 (open dots, control) and mdr65-GAL4 crossed to UAS-shScafRNAi (red dots). n = 30, 32, and 30 adult flies, respectively. One-way ANOVA and Dunnett’s multiple-comparisons test were performed. E, Sleep assay using the DAM2 system. Graph showing the average sleep fraction per day of single adult flies of the genotypes w1118 crossed to UAS-shScafRNAi (dark dots, control), mdr65-GAL4 crossed to w1118 (open dots, control) and mdr65-GAL4 crossed to UAS-shScafRNAi (red dots). n = 30, 32, and 30 adult flies, respectively. One-way ANOVA and Dunnett’s multiple comparisons test were performed. ** p < 0.01. F, G, Activity (F) and sleep profiles (G) showing average values in each time point in a 12:12 light:dark cycle. Adult flies of the genotypes: w1118; mdr65-GAL4, shScafRNAi (dark dots, control), mdr65-GAL4 crossed to w1118 (open dots, control) and mdr65-GAL4 crossed to UAS-shScafRNAi (red dots). n = 30, 32, and 30 adult flies respectively. ZT, Zeitgeber time; ns, nonsignificant.
embryogenesis, it was postulated that Scaf could act on an unknown extracellular signal, including components of the extracellular matrix (Rousse et al., 2010). Spheroid, another SPH, regulates Toll pathway activation in a bacterial infection model (Pattnogic and Leclerc, 2017), whereas the SPH Masquerade regulates muscle attachment by stabilizing cell–matrix interactions (Murugasu-Oei et al., 1995). We showed that Scaf reaches the entire BBB; therefore, Scaf may play a similar role in modulating signaling pathways in the BBB as other SPHs. Alternatively, Scaf could also have a role intracellularly because it can be found in early, late, and recycling endosomes in neighboring cells (Sorrosal et al., 2010).

Interestingly, Scaf displays a cell nonautonomous effect over NB proliferation. We did not detect Scaf in NBs, therefore, this effect is indirect. One possibility is that reducing BBB growth could have an impact on nutrient availability inside the brain. This could explain the fact that Scaf overexpression during nutrient restriction enhances brain growth. Similarly, a reduction in BBB growth, mediated by Scaf, could increase the secretion of growth factors such as insulin-like peptides and activin that control NB proliferation (Zhu et al., 2008; Chell and Brand, 2010; Sousa-Nunes et al., 2011). Additionally, its overexpression in the BBB did not rescue the proliferation effect observed in the mutants, suggesting that other glial cells, such as cortex glia, could be expressing Scaf and contributing to the neurogenic niche.

We believe that the decrease in Scaf levels is part of a highly complex mechanism to maintain growth in the BBB during NR. Knocking down scaf in the BBB does not completely recapitulate the phenotype observed during starvation, suggesting that other pathways are affected. Moreover, reexpression of Scaf during NR does not rescue BBB growth. We propose that during normal

Figure 11. Drug sensitivity is affected by scaf knockdown in the blood–brain barrier. A, B, Ethanol sensitivity assay. A, Graph showing the inactive fraction of flies at different time points. Adult flies of mdr65-GAL4 animals crossed to w1118 (control, dark dots) and UAS-shScafRNAi (scaf knockdown, red dots). B, Sedation time point (ST50) was calculated from A. Bar chart showing ST50 for each genotype. n = 5 groups of 8–10 adult flies for each genotype. Unpaired Student’s t test was performed. C, Insecticide resistance assay. Adult flies of mdr65-GAL4 crossed to w1118 (control, dark dots), UAS-shScafRNAi (scaf knockdown, red dots) and UAS-IshScafRNAi (scaf knockdown, cyan dots). Plot shows the lethal fraction after 12, 24, and 36 h of exposure to 0.01% malathion. n = 4 groups of 20 adult flies for each genotype. Two-way ANOVA and Dunnett’s multiple-comparisons test were done. D, E, Ethanol sensitivity assay of male w1118 flies that grew under Fed or NR conditions. D, Graph showing the inactive fraction of flies at different time points. Fed (blue) and NR (red). E, Sedation time point (ST50) was calculated from D. Bar chart showing ST50 for Fed and NR conditions. n = 8 and 6 groups of 8 adult flies each. Unpaired Student’s t test was performed. **p < 0.01, ****p < 0.001. F, G, Expression of the mdr65-GAL4 driver in adult tissues. (F) Images of the adult brain and (G) hindgut (arrowhead) and malpighian tubules (arrow). DAPI was used to stain DNA (blue) and UAS-lam:GFP (green) for marking nuclei. H, I, Blood–brain barrier permeability assay in mdr65-GAL4 animals crossed to (H) w1118 (control) or (I) UAS-shScafRNAi. Images show adult brains; DNA was stained with DAPI in blue and 10 kDa dextran-Rhodamine in red. Scale bars: 100 μm.
development, Scaf responds to nutritional availability and acts as a negative feedback over BBB growth, contributing to the fine-tuning of the balance between BBB growth and neurogenesis in the CNS. During larval starvation, this balance is lost, and neurogenesis is prioritized at the expense of the growth of the BBB. Future experimental evidence will be needed to unveil the molecular mechanisms by which Scaf supports this balance of growth during the development of the nervous system.

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