

1 **Title: High Sucrose Diets Contribute to Brain Angiopathy with Impaired Glucose**

2 **Uptake, and Psychosis-related Higher Brain Dysfunctions in Mice**

3 **Short title: Angiopathy in mice and humans with psychosis**

4 **Authors: Shinobu Hirai^{1,†}, Hideki Miwa^{1,2}, Tomoko Tanaka¹, Kazuya Toriumi³,**

5 **Yasuto Kunii⁴, Hiroko Shimbo¹, Takuya Sakamoto⁵, Mizuki Hino⁴, Ryuta Izumi⁴,**

6 **Atsuko Nagaoka⁴, Hirooki Yabe⁴, Tomoya Nakamachi⁶, Seiji Shioda⁷, Takashi Dan⁸,**

7 **Toshio Miyata⁸, Yasumasa Nishito⁹, Kazuhiro Suzuki³, Mitsuhiro Miyashita³,**

8 **Toshifumi Tomoda¹⁰, Takatoshi Hikida¹¹, Junjiro Horiuchi¹², Masanari Itokawa³,**

9 **Makoto Arai³, Haruo Okado^{1,†}**

10

11 **Affiliations:** ¹Neural Development Project, Department of Brain Development and Neural

12 Regeneration, Tokyo Metropolitan Institute of Medical Science, Tokyo 156-8506, Japan.

13 ²Molecular Neuropsychopharmacology Section, Department of Neuropsychopharmacology,

14 National Institute of Mental Health, National Center of Neurology and Psychiatry, Tokyo

15 187-8553, Japan.

16 ³Schizophrenia Research Project, Department of Psychiatry and Behavioral Sciences, Tokyo

17 Metropolitan Institute of Medical Science, Tokyo 156-8506, Japan.

- 1 ⁴Department of Neuropsychiatry, School of Medicine, Fukushima Medical University,
- 2 Fukushima 960-1295, Japan.
- 3 ⁵Department of Applied Biological Science, Faculty of Science and Technology, Tokyo
- 4 University of Science, 2641 Yamazaki, Noda, Chiba 278-8510, Japan.
- 5 ⁶Laboratory of Regulatory Biology, Graduate School of Science and Engineering,
- 6 University of Toyama, Toyama 930-8555, Japan.
- 7 ⁷ Department of Clinical Pharmacy, Shonan University of Medical Sciences, Yokohama
- 8 244-0806, Japan
- 9 ⁸Division of Molecular Medicine and Therapy, Tohoku University Graduate School of
- 10 Medicine, Miyagi 980-8575, Japan.
- 11 ⁹Center for Basic Technology Research, Tokyo Metropolitan Institute of Medical Science,
- 12 Tokyo 156-8506, Japan.
- 13 ¹⁰Centre for Addiction and Mental Health, Campbell Family Mental Health Research
- 14 Institute, University of Toronto, Toronto, Ontario M5T 1R8, Canada.
- 15 ¹¹Department of Laboratory for Advanced Brain Functions, Institute for Protein
- 16 Research, *Osaka* University, Osaka 565-0871, Japan.
- 17 ¹²Tokyo Metropolitan Institute of Medical Science, Tokyo 156-8506, Japan.
- 18

1 †These authors contributed equally to this work.

2 Correspondence should be addressed to Shinobu Hirai and Haruo Okado, Neural

3 Development Project, Department of Brain Development and Neural Regeneration, Tokyo

4 Metropolitan Institute of Medical Science, Tokyo 156-8506, Japan.

5 E-mail: hirai-sn@igakuken.or.jp (S.H.) and okado-hr@igakuken.or.jp (H.O.)

6

7 **Abstract:** Metabolic dysfunction is thought to contribute to the severity of psychiatric
8 disorders; however, it has been unclear whether current high-simple-sugar diets contribute
9 to pathogenesis of these diseases. Here we demonstrate that a high-sucrose diet during
10 adolescence induces psychosis-related behavioral endophenotypes, including hyperactivity,
11 poor working memory, impaired sensory gating, and disrupted interneuron function in mice
12 deficient for glyoxalase-1 (Glo1), an enzyme involved in detoxification of sucrose
13 metabolites. Further, the high-sucrose diet induced microcapillary impairments and reduced
14 brain glucose uptake in brains of *Glo1* deficient mice. Aspirin protected against this
15 angiopathy, enhancing brain glucose uptake, and preventing abnormal behavioral
16 phenotypes. Similar vascular damage to our model mice was found in the brains of
17 randomly collected schizophrenia and bipolar disorder patients, suggesting that psychiatric

1 disorders are associated with angiopathy in the brain caused by various environmental
2 stresses, including metabolic stress.

3

4 **Introduction:**

5 Considering the global increase in consumption of simple sugars, the World Health
6 Organization published guidelines that addressed concerns regarding body weight gain and
7 dental caries development in 1995 (1). High-sugar intake alone increases the risks of
8 numerous chronic diseases, including diabetes, hypertension, and kidney disease. However,
9 there are few studies on the effects of high-sugar intake during adolescence on future mental
10 health. The daily caloric intake from simple sugar by teenagers is higher than that observed
11 for other age groups (~20% of total daily caloric intake) (2). Most chronic psychiatric
12 disorders, including schizophrenia (SZ) and bipolar disorder (BD) discussed in the present
13 study, develop before the age of 30 via complex interactions among multiple genetic and
14 environmental risk factors. Interestingly, patients with SZ and BD consume approximately
15 2-fold more sugar than age-matched healthy individuals, and patients with SZ who consume
16 more sucrose exhibit more severe symptoms (3-5). Moreover, the odds ratios for mental
17 distress, hyperactivity, and behavioral disorders were the highest among adolescents who
18 self-reported the highest consumption of soft drinks (6).

1 Advanced glycation end products (AGEs) and reactive carbonyl compounds are
2 produced from irreversible non-enzymatic crosslinking reactions between
3 carbonyl-containing molecules (e.g. sugars, reactive carbonyl compounds) and other
4 molecules (e.g. amino groups of proteins), and AGEs can lead to the induction of
5 inflammation and oxidative stress in various tissues, leading to a positive feedback loop that
6 further increases AGEs (7). There is substantial evidence that patients with psychiatric
7 disorders have elevated proinflammatory cytokines (e.g. interleukin (IL)-1 β , IL-6) and
8 increased oxidative stress (8-11). Glyoxalase I (GLO1) is a zinc metalloenzyme that
9 protects cells from AGE toxicity by catalyzing the reaction between the reactive carbonyl
10 compound methylglyoxal to glutathione to form S-lactoyl-glutathione (12). Patients with
11 depressive-state BD and major depressive disorder have reduced amounts of GLO1,
12 suggesting that this reduction, in combination with an increase in sugar consumption, may be
13 a cause of the increased inflammation and oxidative stress observed in these diseases (13).
14 Furthermore, a SZ patient exhibiting poor convalescence is found to harbor a frameshift
15 mutation in *GLO1* leading to reduced enzyme activity (14, 15).

16 Despite accumulating evidence, it is still unproven that excessive sugar intake
17 contributes to the pathogenesis of psychiatric disorders among susceptible individuals. We
18 addressed this question by generating mice with psychosis-related phenotypes associated

1 the gene \times environment interactions ($G \times E$) to determine whether excessive sucrose
2 consumption during adolescence is a novel environmental risk factor for SZ and BD. We
3 further identified angiopathy in mice with high-sucrose intake and genetic vulnerability, and
4 confirmed similar vascular damage in randomly collected brain samples from BD and SZ
5 patients. This suggests that a variety of genetic and environmental factors can adversely
6 affect brain capillaries. In addition, we found that chronic anti-inflammatory drug treatment
7 prevented core phenotypes of psychiatric disorders as well as vascular damage, suggesting
8 that angiopathy may have an important influence on the development and/or condition of
9 psychiatric disorders.

10

11 **Results**

12 **Psychosis-related behavioral phenotypes in mice on a high-sucrose diet**

13 We fed mice one of two diets containing the same total calories and caloric proportions
14 of carbohydrates, fat, and proteins, but with either starch or sucrose as the main
15 carbohydrate (Fig. 1A). We investigated four groups of mice fed these diets for 50 days
16 immediately after weaning (from postnatal day 21, corresponding to the juvenile/adolescent
17 stage): starch-fed wild-type (WT) mice (control, CTL), sucrose-fed WT mice
18 (environmental stressor, Env), starch-fed *Glo1* heterozygous knockout (*Glo1*/+) mice

1 (genetic factor, Gen), and sucrose-fed *Glo1*^{+/+} mice ($G \times E$) (Fig. 1A). Reduced *Glo1*
2 expression in *Glo1*^{+/+} mice was confirmed in the cerebral cortices, including the
3 hippocampus, using western blotting (fig. S1A, B). The body weight trajectories of these
4 mice were similar to those of control mice for at least 11 weeks of age, indicative of normal
5 structural development (fig. S1C) and removing obesity as a possible cause of the observed
6 group differences described below. We tested eight different behaviors in mice, including
7 open-field locomotor activity (Fig. 1B), acoustic startle response (Fig. 1C), pre-pulse
8 inhibition (PPI; a measure of sensory-motor gating) (Fig. 1D), object location performance
9 (used as a test of working memory) (Fig. 1E), self-grooming (Fig. 1F), nest building (Fig.
10 1G), elevated plus maze activity (fig. S2A), and social interaction (fig. S2B). These
11 behaviors could be classified into those that are not affected by either diet or genotype,
12 those that are affected by genotype but not diet, those affected by diet but not genotype, and
13 those affected by the combination of diet and genotype. Thus, neither diet nor genotype, nor
14 a combination of diet and genotype had any significant effect on social interactions (fig.
15 S2B), while acoustic startle responses were reduced in *Glo1*^{+/+} mice but not significantly
16 affected by feeding type (Fig. 1C). Self-grooming (Fig. 1F), nest building (Fig. 1G), and
17 elevated plus maze activity (fig. S2A) were affected by diet, but not genotype, while open
18 field locomotor activity, PPI, and object location performance required a combination of

1 sucrose feeding and a *Glo1*/+ genotype to be affected. In the open field test (Fig. 1B),
2 neither starch-fed *Glo1*/+ mice, nor sucrose-fed WT mice, showed significant differences in
3 distance traveled compared to starch-fed WT mice, while sucrose-fed *Glo1*/+ mice showed
4 significant increases. Similarly, starch-fed *Glo1*/+ mice and sucrose-fed WT mice had
5 similar PPI scores compared to starch-fed WT mice, while sucrose-fed *Glo1*/+ mice showed
6 significant decreases (Fig. 1D). Sucrose-fed WT mice had a non-significant reduction in
7 object location test scores, and this reduction became significant in sucrose-fed *Glo1*/+
8 mice (Fig. 1E). Altogether, our results suggest that a combination of sucrose feeding and
9 *Glo1*/+ mutation can generate novel phenotypes reminiscent of psychiatric disease
10 phenotypes that are not found by altering diet or genotype alone.

11 To further support our idea that synthetic G × E interactions may induce or enhance the
12 severity of psychosis-like symptoms, we examined the effects of the high sucrose diet on an
13 additional major psychosis-associated mutation in *Disrupted-in-schizophrenia 1 (DISC1)*
14 locus (*16*) (fig. S3). *DISC1* plays important roles in neurodevelopment and synaptic activity,
15 but a loss-of-function mutation in *DISC1* gene alone is not sufficient to cause psychiatric
16 disorders (*17*). Similar to the results for *Glo1*/+ mice, *Disc1*/+ mice fed a high sucrose diet (G
17 × E mice) showed severely impaired object location performance (working memory) and PPI
18 (fig. S3C, D), whereas WT mice fed a high sucrose diet or *Disc1*/+ mice fed control diet did

1 not, again suggesting that a combination of genetic and environmental influences can
2 induce psychosis-like behaviors.

3 Schizophrenia (SZ) is associated with increased amphetamine-induced dopamine release
4 in the striatum (18, 19) and unbalanced dopamine regulation between the medial prefrontal
5 cortex (mPFC) and striatum (20). To examine the effects of high sucrose diet on dopamine
6 release in *Glo1*^{+/+} mice, we next measured dopamine levels in the nucleus accumbens (NAc)
7 and mPFC of these mice using *in vivo* microdialysis. We found that the high sucrose diet
8 increased both basal and amphetamine-induced dopamine release in the NAc specifically in
9 *Glo1*^{+/+} mice (Fig. 1H), while reducing basal dopamine release in the mPFC in either genetic
10 background (fig. S2C). To assess whether enhanced dopamine release in the NAc could
11 induce the observed behavioral phenotypes, we examined the effects of aripiprazole, a D2
12 receptor partial agonist and clinical antipsychotic (21), administered during the last 7 days (0.5
13 mg/kg/day) of sucrose feeding (Fig. 1A, I, J and fig. S2D–I). We found that the
14 hyper-locomotion and increased striatal dopamine release in $G \times E$ mice were completely
15 reversed by aripiprazole treatment (Fig. 1I, J), while other behavioral defects were not
16 improved (fig. S2D–I). Thus, aripiprazole treatment selectively improves a subset of
17 behavioral abnormalities in $G \times E$ mice that are likely caused by dysregulated dopamine
18 signaling.

1

2 **Dysfunction of parvalbumin-positive inhibitory interneurons in G × E mice**

3 A reduction in density of parvalbumin (PV)-positive GABAergic interneurons is
4 reported for schizophrenia (22-25), and the precisely coordinated activity of these neurons
5 is crucial for the regulation of PPI and working memory (26). Thus, we next investigated
6 PV expression using immunohistochemistry and western blotting to determine whether
7 alterations in PV signaling may contribute to our observed psychiatric disease-associated
8 phenotypes. A high sucrose diet caused a reduction in PV-positive cells in both *Glo1*/+ and
9 WT mice (Fig. 2A, B) and significantly reduced PV expression in sucrose-fed *Glo1*/+ mice
10 (Fig. 2C, D), suggesting that the sucrose diet and the *Glo1*/+ mutation had additive or
11 synergistic effects on PV signaling. Since gamma oscillations are produced via the
12 synchronous activation of PV neurons, we measured gamma oscillations (30–45 Hz) using
13 surface electroencephalography (EEG) to determine whether downregulation of PV protein
14 was accompanied by functional abnormalities in neural activity. Interestingly, sucrose-fed
15 mice (WT and *Glo1*/+ mice) exhibited elevated baseline gamma oscillation power
16 compared with control mice (starch-fed WT) in the home cage (Fig. 2E). Further, only
17 sucrose-fed *Glo1*/+ mice failed to exhibit an increase in the gamma oscillation power when
18 approaching a novel object (Fig. 2F). These results are consistent with findings in patients

1 with SZ and BD as well as other mouse models of psychosis demonstrating increased
2 baseline gamma oscillations and decreased sensory stimulus-evoked synchronized gamma
3 power (27, 28). Thus, our results suggest that $G \times E$ mice mimic the pathophysiological
4 changes of PV neurons observed in psychiatric disorders.

5 To summarize, the administration of a high-sucrose diet to *Glo1*^{+/+} mice induces
6 behavioral, histological, and pathophysiological phenotypes reminiscent of phenotypes
7 observed in psychiatric disorders. This suggests that excessive sucrose intake during
8 adolescence is a potential environmental risk factor for these diseases.

9

10 **AGE accumulation and impaired astrocyte function in $G \times E$ mice**

11 How does a decrease in GLO1 expression and an increase in sucrose consumption
12 lead to defects in activity of PV and dopaminergic neurons? We first examined GLO1
13 expression and detected high expression in astrocytes (fig. S4A–G), especially those
14 surrounding capillaries (fig. S4B, C), moderate expression in neurons (fig. S4H, I), and low
15 expression (below detection limits) in vascular endothelial cells (fig. S4A–G) and
16 microglial cells (fig. S4H, J). Expression was not detected in the brains of *Glo1*
17 homozygous mice (fig. S3K) indicating that the antibody we used was specific for GLO1.

1 Since both GLO1 and a high sucrose diet are associated with glycation toxicity, we
2 next examined AGE immunoreactivity to detect levels of AGE products of glucose
3 metabolism and/or chemical reaction of glucose. We found stronger AGE immunoreactive
4 signals in the vascular endothelial cells of G × E mice compared with control mice (Fig.
5 3A–H). Moreover, we detected AGE products of methylglyoxal metabolism and/or
6 chemical reaction of methylglyoxal using AGE-4 antibody, in the microglia of sucrose-fed
7 mice (fig. S5A–E). Compared to controls, AGE accumulation in sucrose-fed *Glo1*^{+/+} mice
8 was accompanied by elevated IBA1 immunofluorescence intensity and an enlarged
9 CD68-positive area (fig. S5F, G, H), both of which are phenotypes identified in activated
10 microglia (29, 30). Thus, sucrose feeding in *Glo1*^{+/+} mice results in microglial and
11 endothelial AGE accumulation and microglial activation.

12 Since astrocytes express *Glo1* most strongly (fig. S4A–G), we next used mice
13 expressing green fluorescent protein (GFP) under control of the glial fibrillary acid protein
14 (GFAP) promoter (31, 32) to evaluate how astrocytes are affected in our G x E condition.
15 Typically, cellular damage from AGEs is caused by inflammatory responses induced by
16 Receptor of AGEs (RAGE) activation or by a loss of normal protein function following
17 AGE-forming reactions (33). Astrocytes exhibit a well-described reactive phenotype in
18 response to pathogenic conditions including neuroinflammation characterized by enhanced

1 GFAP expression (34, 35). Strongly enhanced *GFAP* promoter function was observed in G
2 × E mice, without changes in the number of GFAP-positive astrocytes (Fig. 3I–K),
3 indicating that the astrocytes in G × E mice are in the reactive pre-condition during the
4 high-sucrose feeding. Taken together, these results suggest that a high sucrose diet enhances
5 AGE production in endothelial cells and microglia (Fig. 3A–H and fig. S5A–E), and
6 converts astrocytes into a pre-inflammatory state in *Glo1*-deficient mice (Fig. 3I–K).

7

8 **Microcapillary angiopathy and impaired glucose intake in G × E mice**

9 Endothelial cells and astrocytes are functional and physical components of the
10 blood-brain barrier (BBB) which tightly controls the parenchymal environment by
11 modulating the selective passage of nutrients and various factors (36). Thus, endothelial
12 AGE accumulation and astrocyte reactivity may impair BBB function. To examine changes
13 in endothelial function, we first conducted a transcriptome analysis of the prefrontal cortex
14 (PFC), a region strongly implicated in psychiatric impairments, using microarrays (Fig. 4A,
15 B). The coagulation factor V, essential for the production of fibrin from fibrinogen, ranked
16 seventh on a list of transcripts exhibiting more than doubled expression in G × E mice
17 compared with the other three groups (Fig. 4A, Supplementary Table 1, 2). Fibrin controls
18 hemostasis via polymerization with platelets to form blood clots, and deposits of this protein

1 are indicative of endothelial abnormalities (37). In early stages of endothelial cell
2 impairment, fibrin accumulates in capillaries. Therefore, we investigated vascular fibrin
3 accumulation using immunohistochemistry and confirmed the presence of significant fibrin
4 accumulation on the vascular lumen side of endothelial cells in brain capillaries of $G \times E$
5 mice (Fig. 4C–G). Fibrin leakage and deposition in the brain parenchyma, as observed in
6 Alzheimer’s disease, was not detected in $G \times E$ mice, indicating that physical BBB
7 disruption did not occur (38).

8 We next speculated that the abnormal vascular endothelial cells and astrocytes
9 observed in $G \times E$ mice could alter glucose uptake from the plasma into the brain
10 parenchyma. Extracellular glucose concentrations in the brain parenchyma were measured
11 under three conditions: 1) fasting, 2) 1 h after feeding, and 3) 2 h after feeding. We first
12 measured a baseline glucose concentration in the mPFC in starch-fed WT mice after
13 starvation for 16 hrs (Fig. 4H). This concentration increased significantly within an hour
14 after starch feeding, and decreased close to baseline 2 hrs after feeding. Sucrose-fed WT
15 mice and starch-fed *Glo1*^{+/+} mice had similar baseline glucose concentrations, similar
16 increases 1 hr after starch or sucrose feeding, and similar decreases back to baseline 2 hrs
17 after feeding. In contrast, sucrose-fed *Glo1*^{+/+} mice had both significantly decreased basal
18 glucose concentrations in the mPFC, and no apparent increase in glucose concentrations

1 after sucrose feeding. We measured vascular diameter and expression of glucose transporter
2 1, the major glucose transporter expressed in vascular endothelial cells and astrocytes, and
3 observed no differences among the four groups (fig. S6). Further, no differences were
4 detected in plasma glucose after starvation and feeding, or in fasting plasma insulin levels
5 among the four groups (Fig. 4I, J), indicating that the lower parenchymal glucose
6 concentrations in $G \times E$ mice is due to reduced uptake across the BBB rather than
7 dysregulation of plasma glucose or insulin signaling.

8

9 **Protective effects of chronic low-dose aspirin against behavioral abnormalities and**
10 **angiopathy**

11 Previous reports have shown that adjunct non-steroidal anti-inflammatory drug (NSAID)
12 treatment can improve psychiatric disorder scores (39, 40). Aspirin, an NSAID, is routinely
13 used for the prevention and alleviation of vascular-related adverse events associated with
14 high blood pressure, ischemia, and cardiovascular diseases (41, 42). Here, we examined
15 whether aspirin treatment can protect against the development of psychiatric phenotypes in
16 $G \times E$ mice (Fig. 1A, Fig. 5A–D and fig. S7A–I). Low-dose aspirin (1 mg/kg/day) fully
17 prevented hyperlocomotor activity, deficits in PPI, working memory, grooming duration
18 (Fig. 5A–D), and partially prevented abnormal enhancement of DA release after

1 methamphetamine administration and reduced PV expression among $G \times E$ mice (fig.
2 S7D–F), but did not improve acoustic startle responses, nest building and elevated plus
3 maze scores and abnormal astrocyte activation (fig. S7A–C, G–I). These improvements
4 were accompanied by a decrease in endothelial fibrin accumulation (Fig. 5E, F) and a
5 partial restoration of glucose intake into the brain parenchyma (Fig. 5G). Collectively, these
6 results suggest that aspirin treatment in $G \times E$ mice significantly improves angiopathy and
7 brain glucose availability, contributing to prevention of several behavioral abnormalities.

8

9 **Angiopathy in postmortem brains of patients with psychiatric disorders**

10 Finally, we compared immunostains of brain slices from healthy controls and patients with
11 SZ or BD to examine whether these patients exhibit angiopathic fibrin accumulation in
12 vascular endothelial cells. Similar to our findings in $G \times E$ mice, we found significantly
13 elevated fibrin accumulation in the vascular endothelium of brain slices from randomly
14 collected SZ and BD patients (Fig. 6A–C). Therefore, fibrin-related angiopathy and
15 vascular damage may be a novel and common phenotype of psychiatric illness contributing
16 to disease progression.

17

18 **Discussion**

[ここに入力]

1 Using a mouse model, we demonstrated that high dietary sucrose consumption during
2 adolescence is a potential risk factor for the development of behavioral phenotypes
3 associated with psychiatric illnesses, such as SZ and BD. These behavioral phenotypes
4 include impaired sensory gating, dysfunctions in PV interneurons and working memory,
5 hyperactivity, and increased basal and stimulus-evoked striatal dopamine release. Second,
6 we identified endothelial fibrin accumulation (“angiopathy”) in both our mouse model and
7 in randomly collected postmortem brains of patients with SZ or BD. We also observed that
8 glucose intake from the plasma into the brain parenchyma was impaired in $G \times E$ mouse
9 model. Chronic low-dose aspirin treatment prevented fibrin deposition in the capillaries,
10 improved glucose transport, and reversed several behavioral phenotypes in $G \times E$ mice,
11 suggesting angiopathy as a seminal pathogenic event in mental illness.

12 In Fig. 6D we propose a possible model summarizing our results. A high sucrose diet
13 causes AGE accumulation in microglia and endothelial cells which express low amounts of
14 GLO1 (Fig. 3A–H and fig. S5A–E). Astrocytes are less sensitive to this diet in WT animals
15 because they express high amounts of GLO1 to relieve glycation stress (fig.
16 S4A–G). However, the reduced GLO1 in *Glo1*^{+/−} mice causes astrocytes to become
17 activated by a high sucrose diet (Fig. 3I–K). This activation may be caused directly by
18 glycation stress in astrocytes or may occur as a consequence of increased microglial and

1 endothelial AGEs. Astrocytic activation further increases microglial and endothelial AGEs
2 in a positive feedback loop (7). Increased microglial and endothelial AGEs and astrocytic
3 activation may induce fibrin accumulation and damage to the brain vasculature (33) (Fig.
4 4C–G). This results in decreased glucose import into the brain and dopaminergic and PV
5 neuron dysfunction which in turn cause psychosis-associated behavioral defects (Fig. 1B–H,
6 Fig. 2 and Fig. 4H).

7 Strikingly, administration of the NSAID aspirin, which inhibits inflammation and
8 oxidative stress, protected against the emergence of angiopathy as evidenced by reduced
9 fibrin accumulation, partially restored parenchymal glucose concentrations, and prevention
10 of several psychiatric-related phenotypes (Fig. 5 and fig. S7D–F). The increase in
11 coagulation factor V and the accumulation of fibrin in endothelial cells in G x E mice
12 indicate the presence of some vascular damage (43) (Fig. 4A–G). Aspirin may prevent this
13 damage through its antiplatelet coagulation and anti-inflammatory properties. Since
14 oxidative stress and chronic inflammation are also common features of psychiatric disorders,
15 aspirin may provide a protective effect against the etiology of these diseases as well (8, 11,
16 44, 45). We have tried to quantify ROSs in our experimental mice, but thus far have not
17 been able to see differences among the four groups (fig. S8), so a more sensitive detection
18 method such as FRET probe for ROS detection may be required.

1 Sucrose consists of glucose and fructose, when these sugars are ingested at the same time,
2 we find that they generate AGEs in different cell types. Glucose-metabolism-derived AGEs,
3 which can be detected using α -AGE polyclonal antibody, accumulated primarily
4 in vascular endothelial cells (Fig. 3A–H), while methylglyoxal-derived AGEs, recognized
5 by the α -AGE-4 monoclonal antibody accumulated specifically in microglia (fig. S5A–E).
6 Our results suggest that these two cell types may import and metabolize these two sugars
7 differently, but more detailed validation, such as one-cell metabolomic analysis, is required
8 to validate this idea.

9 Robust PV neuron function is required for PPI, working memory,
10 amphetamine-induced hyper-locomotion, dopamine (or 3,4-dihydroxyphenylacetic acid)
11 release regulation, and gamma oscillation generation, all of which are considered core
12 symptoms of psychiatric disorders (26). The PV neuron dysfunction in $G \times E$ mice may be
13 caused by the unique properties of these cells. PV-expressing interneurons are the
14 fast-spiking neuron subtype exhibiting a lower input resistance and higher-amplitude rapid
15 after-hyperpolarization than several projection neurons, and the combination of these
16 properties generates higher frequency action potentials compared with those by other
17 neuron types. This frequency of high action potentials in PV neurons can be exploited by
18 their fast spike phenotype to produce fast oscillations (30-100 Hz) in the gamma band of

1 neural oscillations (46). To maintain this rapid spiking property, PV neurons require high
2 energy expenditure as evidenced by mitochondrial and cytochrome c oxidase enrichment
3 (47). Therefore, reduced glucose within the brain parenchyma may preferentially disturb PV
4 neuron function, resulting in reduced PV expression. Indeed, reduced PV neuron number
5 has been repeatedly reported in postmortem brains of patients with psychiatric disorders
6 such as SZ and BP, and this reduction is proposed to be caused by a decrease in PV mRNA
7 or protein expression in these cells, rather than by a loss of PV neurons themselves (22-25).
8 Since PV is a Ca²⁺ binding protein, proper regulation of the expression of this protein is
9 considered to be important for maintaining the plasticity of PV neurons. For example, in PV
10 neurons of the hippocampus and mPFC, the expression of PV protein changes after
11 contextual fear conditioning, exposure to rich environments, and pharmacogenetic
12 activation of PV neurons, suggesting PV expression and plasticity are correlated (25, 48).
13 Further, it is known that the inhibitory activity of PV neurons is critical under
14 environmental stress to prevent a sequela of excessive excitatory activity such as oxidative
15 stress and inflammation (47). Pathogenic processes might more readily occur during the
16 critical adolescent prodromal period, wherein PV neurons may have increased vulnerability
17 to environmental stresses because of delayed maturation, which is another feature of PV
18 neurons (49).

1 In the present study, we identified capillary angiopathy in both $G \times E$ mice and
2 postmortem brains of patients with SZ and BD (Fig. 4C–G and Fig. 6A–C). In our $G \times E$
3 mice, angiopathy was caused by the high AGE production capacity of sucrose combined
4 with a GLO1 deficiency. Since high sugar consumption has been associated with SZ and
5 BD, and decreased GLO1 expression and function have been seen in SZ and BD, similar
6 interactions may be associated with some cases of psychiatric diseases. However, various
7 other environmental stresses and genetic conditions may also converge to induce
8 angiopathy. Several studies have reported that stressors, such as social defeat, isolation, and
9 viral infection, induce vascular defects (50-52). These stressors are also risk factors for SZ
10 and BD and induce PV neuron hypofunction (53), suggesting that angiopathy may be a
11 common trigger for psychiatric phenotypes.

12

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[ここに入力]

1 **Author contributions:** S.H. and H.O. designed the study. S.H. performed and analyzed all
2 experiments. H.M. assisted with the experimental design of EEG recording and analysis of
3 the results and edited the manuscript. T.T. coordinated the EEG recording. Y.K., M.H., R.I.,
4 A.N., and H.Y. helped with the design of experiments using human specimens and provided
5 fixed human brain sections. T.S. performed data and image analysis. T.N. and S.S. provided
6 GFAP-GFP mice. T.D. and T.M. generated and provided *Glo1* knockout (KO) mice. K.T.
7 and K.S. backcrossed *Glo1* KO mice to B6J mice for all experiments. Y.N. performed
8 cDNA microarray analysis of gene expression. H.S. assisted with behavioral experiments.
9 M.I., M.A., K.T., K.S., and M.M. provided important suggestions to this study. T. T. and T.
10 H provided critical input on *Disc1-L1* mouse experiments. J.H. edited the manuscript. S.H.
11 generated all figures, tables, and wrote the manuscript. H.O. edited the manuscript and
12 supervised this study.

13

14 **Competing interests:** The authors declare no competing interests.

15 **Data and materials availability:** All materials used in this paper are available upon request.

16 The accession number for transcriptome analysis data is provided in the Materials and

17 Methods.

18

1 **Figure Legends**

2 **Fig. 1 Generation of G × E model mice and psychosis-related phenotype analyses.**

3 (A) Experimental Timeline. See Materials and Methods for a complete description. (B–G)

4 Behavioral analyses in the four groups of mice (n = 18–23 mice per group). (B)

5 Spontaneous locomotor activity in the open-field test. (C) Acoustic startle responses. (D)

6 Pre-pulse inhibition (PPI) using a 70 dB pre-pulse. (E) Object location test to evaluate

7 working memory. (F) Duration of self-grooming in the home cage. (G) Quantification of

8 nest-building skills over 8 h. Post hoc Dunnett’s multiple comparisons test among groups at

9 each time point indicated $###p < 0.001$ for Suc *glol*/+ vs. Ctrl (Starch +/+), $$$p < 0.01$ for

10 Suc +/+ vs. Ctrl (Starch +/+). (H, J) Extracellular dopamine (DA) concentration in the

11 nucleus accumbens measured at 20 min intervals using an *in vivo* microdialysis system in

12 the presence or absence of aripiprazole (Aripip). Methamphetamine (1.25 mg/kg) was

13 administered via intraperitoneal (i.p.) injection at time 0 (arrow) (n = 6–11 mice per group).

14 See Materials and Methods for detailed statistical analyses. (I) Effects of Aripip treatment on

15 locomotor activity (n = 12–18 mice per group). (B–H) Starch +/+ group was used as a

16 control for post hoc Dunnett’s test. All data are presented as mean ± SEM. $***p < 0.001$,

17 $**p < 0.01$, $*p < 0.05$

18

1 **Fig. 2 Parvalbumin-positive interneuron dysfunction in G × E mice.**

2 (A) Parvalbumin (PV) immunohistochemistry in the medial prefrontal cortex (mPFC). (B)

3 Number of PV-positive cells in the mPFC (n = 4 mice per group). (C) Western blot analysis of

4 PV protein expression using tubulin as internal control. (D) Densitometric analysis of PV

5 protein expression: PV band intensities were divided by the corresponding tubulin band

6 intensities (n = 3 mice per group). (E) Average gamma band power in the home cage and

7 during novel object recognition in the open field (n = 7–8 mice per group). (F) Changes in

8 gamma band power from the home cage to the novel object phase in individual mice (n = 7–8

9 mice per group). Each *p* value indicates the result of repeated measures analysis of variance

10 (ANOVA). The effect of changes in gamma power at Starch +/+ (F1, 5 = 8.29, *p* = 0.035), at

11 Starch *Glo1*/+ (F1, 6 = 29.75, *p* = 0.0016), at Sucrose +/+ (F1, 6 = 12.04, *p* = 0.013), at

12 Sucrose *Glo1*/+ (F1, 7 = 4.038, *p* = 0.084). (B, D, H) Starch +/+ group was used as control for

13 post hoc Dunnett's test. All data are presented as mean ± SEM. ****p* < 0.001, ***p* < 0.01, **p*

14 < 0.05

15

16 **Fig. 3 AGE accumulation in the neurovascular endothelium and pre-inflammatory**

17 **status of astrocytes in G × E mice.**

1 (A) AGE immunohistochemistry in the medial prefrontal cortex. (B) Measurement of AGE
2 immunoreactive area. The mean intensity of the AGE-immunopositive area in the entire
3 image was measured for each section (n = 3 mice per group). (C–E) Colocalization of the
4 endothelial cell marker tomato lectin with the astrocytic marker ALDH1L1 or AGEs. (F)
5 Enlarged version of images in (C) presenting the region used for colocalization analysis.
6 (G) 2D intensity histogram of the two indicated channels for identifying the colocalization
7 of AGE with ALDH1L1 (Astrocyte) or tomato lectin (Endothelial cell). (H) Average
8 R-value of colocalization data including (G) three different locations of three mice. (I)
9 Immunohistological images of GFP-positive astrocytes and ALDH1L1 in the mPFC region
10 and merged GFP/ALDH1L1 image. (J) Percentage of GFP-positive cells per total
11 ALDH1L1-positive cells in each image in (I). (K) Mean fluorescent GFP intensities of 10
12 randomly selected cells per image in (I) (from four independent mice). (B, J, K) The Starch
13 +/- group was used as control for post hoc Dunnett's test. All data are presented as mean ±
14 SEM. *** $p < 0.001$, ** $p < 0.01$, * $p < 0.05$

15

16 **Fig. 4 Angiopathy and impaired glucose transport in G × E mice.**

17 (A, B) Venn graph showing overlap in prefrontal cortex genes exhibiting a >2-fold (A) or
18 <0.5-fold (B) expression change compared with the control (CTL) group (n = 3 mice per

1 group). (C) Immunohistochemical images of fibrin and the endothelial cell marker tomato
2 lectin. (D) Measurement of the area of fibrin immunoexpression in (C). The mean intensity
3 of the fibrin-immunopositive area of the entire image was measured for each section (n = 4
4 mice per group). (E) Magnified immunohistochemical images of fibrin and the endothelial
5 cell marker tomato lectin. (F) 2D intensity histogram of the two indicated channels using
6 the left and middle panels in (E) for identifying fibrin colocalization with tomato lectin. (G)
7 Average R-value of colocalization data in (F) from three different locations of three mice.
8 (H) Extracellular glucose concentrations in the dialysis buffer from mPFC samples at each
9 time point (n = 5–6 mice per group). See the Materials and Methods for detailed statistical
10 analyses. (I) Plasma glucose concentrations (n = 6–7 mice per group). The first
11 measurement was performed after 16 h of fasting, and the second blood collection 30 min
12 after eating 0.05 g of carbohydrates. (J) Fasting plasma insulin levels (n = 5–6 mice per
13 group). (D, H–J) Starch +/+ group was used as a control for post hoc Dunnett's test. All
14 data are presented as mean ± SEM. **p* < 0.05

15

16 **Fig. 5 Protective effects of low-dose aspirin in G × E mice.**

17 (A–D) Results of behavioral tests performed to evaluate the effects of aspirin treatment (n =
18 12–21 mice per group). (A) Quantification of spontaneous locomotor activity in the open

1 field. **(B)** Pre-pulse inhibition at 70 dB. **(C)** Object location test of working memory. **(D)**
2 Quantification of nest-building skills over 8 h. **(E)** Immunohistochemical images of fibrin
3 and the endothelial cell marker tomato lectin. **(F)** Measurement of the area of fibrin
4 immunoexpression in **(E)**. The mean intensity of the fibrin-immunopositive area of the
5 entire image was measured for each section (n = 3 mice per group). **(G)** Extracellular
6 glucose concentrations in the dialysis buffer at each time point (n = 4–6 mice per group).
7 Post hoc Tukey's multiple comparisons test of groups at each time point, $^{####}p < 0.001$, $^{##}p <$
8 0.01 , $^{\#}p < 0.05$ for Starch $+/+$ vs. Suc $Glo1/+$, $^{$$$}p < 0.001$, $^{\$}p < 0.05$ for Starch $+/+$ Asp vs.
9 Suc $Glo1/+$, $^{\&}p < 0.05$ for Starch $+/+$ vs. Suc $Glo1/+$ Asp. All data are presented as mean \pm
10 SEM. $^{***}p < 0.001$, $^{**}p < 0.01$, $^{*}p < 0.05$

11
12 **Fig. 6 Angiopathy in postmortem brains from individuals with psychiatric disorders.**

13 **(A)** Representative immunohistochemical fibrin images in the BA9 region of postmortem
14 brains from controls and patients with schizophrenia (SZ) or bipolar disorder. **(B)**
15 Measurement of the area of fibrin immunoexpression in **(A)**. The mean intensity of the
16 fibrin-immunopositive area of the entire image was measured for each section. **(C)**
17 Representative immunohistochemical images of fibrin (magenta) and the endothelial cell
18 marker tomato lectin (green) in postmortem brains from a patient with SZ. Fibrin-positive

1 areas are merged with areas of vascular endothelial cell marker expression. **(D)** Diagrams
2 describing the hypothesis proposed to explain functional and behavioral abnormalities in
3 control (CTL) mice (left) and G × E mice (right) (see the Discussion for details). All data
4 are presented as mean ± SEM. ***p* < 0.01

5

6 **Materials and Methods**

7 **Experimental design (related to Fig.1A)**

8 After weaning (P21), wild-type (WT) and *Glo1* heterozygous mutant mice were fed either a
9 starch diet (control) or a sucrose diet (experimental). Diets were equal in total calories and
10 proportions of calories from carbohydrates, lipids, and proteins. The behavioral test battery
11 was administered starting at 2.5 months of age (upper panel). Middle panel: Macronutrient
12 composition of the two diets. We used *Glo1* heterozygous mice (or *Disc1* heterozygous
13 mice) to mimic patients with psychiatric disorders who exhibit decreased GLO1 activity or
14 expression (or decreased *Disc1* expression), whereas high-sucrose intake was used as an
15 environmental risk factor (bottom panel). We investigated four groups of mice: WT,
16 starch-fed mice (Starch +/+); *Glo1* (or *Disc1*) heterozygous, starch-fed mice (Starch *Glo1*/+
17 (or Starch *Disc1*/+)); WT, sucrose-fed mice (Suc +/+); and *Glo1* (or *Disc1*) heterozygous,
18 sucrose-fed mice (Suc *Glo1*/+ (or Suc *Disc1*/+)).

1

2 **Animals**

3 All experimental procedures were approved by the Animal Experimentation Ethics

4 Committee of the Tokyo Metropolitan Institute of Medical Science (49040). All mice were

5 maintained under a 12:12 h light:dark cycle (lights on at 8:00 AM) with free access to the

6 indicated diet. All efforts were made to minimize the number of animals used and their

7 suffering. *Glo1* knockout mice were generated as described previously (54, 55). Briefly,

8 *Glo1*-trapped ES cell lines from the International Gene Trap Consortium were used for the

9 generation of 3 founder mice, which were then backcrossed to C57BL/6 mice. In the

10 manuscript, trapped *Glo1* is referred to as *Glo1. Disc1-LI* (locus-impairment) heterozygous

11 mutant mice were generated as previously described (16). In the manuscript, *Disc1-LI* is

12 referred to as *Disc1*. Alternatively, mice were backcrossed to GFAP-GFP mice to monitor

13 astrocyte activation (31, 32). Male mice were exclusively used in the behavioral tests,

14 whereas mice of both sexes were used in histological, biochemical, and physiological

15 experiments.

16

17 **Diet preparation**

1 The two diets used in the present study were newly created in collaboration with Oriental
2 Yeast Co. Ltd. (Tokyo, Japan). We named the sucrose diet HSD-70 (# OYC 2405100) and
3 the starch diet HCD-70 (# OYC 2405000) (Supplementary Table 4). They contained the
4 same caloric proportions of carbohydrate, fat, and protein; however, all carbohydrate
5 calories are derived from either starch or sucrose.

6

7 **Drug preparation**

8 Aripiprazole was dissolved in acetic acid and diluted to 3.5 mg/L in water for
9 administration at 0.5 mg/kg/day. The final acetic acid concentration in the drinking water
10 was 0.7%. Aspirin was dissolved in ethanol and diluted to 70 mg/L in water for
11 administration at 1 mg/kg/day. The final ethanol concentration in the drinking water was
12 0.15%. The daily dose was based on the measurement of mean water consumption daily in
13 all sucrose-fed *Glo1* heterozygous mice and starch-fed wild-type (WT) mice (fig. S1D, E).
14 There were no between-group differences in terms of aripiprazole consumption (Student's
15 *t*-test, Starch +/+ and Suc *Glo1*/+: $p = 0.86$) or aspirin-containing water (Student's *t*-test,
16 Starch +/+ and Suc *Glo1*/+: $p = 0.61$).

17

18 **Behavioral tests**

[ここに入力]

1 Mice were habituated in the behavioral room for >30 min before each test. Behavioral tests
2 were performed in the following sequence of increasing stress: elevated plus maze,
3 grooming, nest building, open field, object location, social interaction, and pre-pulse
4 inhibition (PPI). All test apparatuses were cleaned using 70% ethanol and water between
5 trials, and the subsequent test session was started only after the ethanol vapor odors had
6 disappeared and the apparatuses had dried.

7 The elevated plus maze (EPM-04M, Muromachi, Japan) consisted of two opposing
8 open arms (297×54 mm) and two closed arms ($300 \times 60 \times 150$ mm) extending from a
9 central platform (60×60 mm). The entire apparatus was elevated 400 mm above the floor.
10 Each mouse was placed on the central platform facing a closed arm and allowed to freely
11 explore the maze for 10 min. Arm entry was defined as the entry of all four paws into the
12 arm. The time spent in the open arms over 10 min was recorded as an index of state anxiety.

13 For the self-grooming test, all mice housed in the same home cage were moved into a
14 new cage for 10 min. Each mouse was then individually placed in a standard mouse home
15 cage ($31 \times 16.5 \times 14$ cm) illuminated at ~200 lux. After a 10-min habituation period, each
16 mouse was scored for cumulative time spent grooming all body regions (56) over 10 min
17 using a stopwatch. Self-grooming behavior is conserved across species and is indicative of

1 certain pathological conditions or factors. In humans, for example, self-grooming increases
2 during stressful conditions and in certain psychiatric disorders (56).

3 For the nest building test, 200 g of corncob was spread across the bottom of each cage
4 for bedding, and a square-shaped piece of cotton was placed in the cage center as raw
5 material for the nest. Each mouse was individually placed in the cage for 8 h. Photos of the
6 constructed nest were acquired every 2 h, and the nest building process was evaluated by
7 measuring the proportion of loose cotton as follows: 1 point for 25% weight (Wt%)
8 loosened, 2 points for 50 Wt% loosened, 3 points for 75 Wt% loosened, and 4 points for
9 100 Wt% loosened. After 8 h, we checked the shape of the nest and added 1 point if the
10 mice had completed a nest with a bird's nest-like shape. The temperature of the room was
11 maintained at 25°C and illumination at 150–180 lux during nest building. Nest-building
12 behavior is an indicator of general well-being in mice (57), whereas poor nest building is an
13 indicator of psychological or physiological abnormalities (58, 59).

14 For the open field (OF) test, each mouse was placed in the center of the apparatus (40 ×
15 40 × 40 cm; 150–180 lux illumination) and allowed to move freely for 10 min. The
16 behavior of each mouse was monitored using a charge-coupled device camera mounted on
17 the ceiling above the OF. The total distance traveled (cm) was measured using CompACT
18 VAS software (Muromachi).

1 For the object location test (OLT) of working memory (60), mice first explored the
2 empty OF box, and then, two identical objects A and B (two 500-mL PET bottles filled with
3 blue-colored water) were placed in two corners 5 cm from the wall. After a 10-min
4 exploration/learning period, the mice were returned to their home cage for 5 min, and
5 Object A was moved to a new corner (Object A'). The animals were then placed back in the
6 OF box and allowed to explore for 5 min. The time spent exploring A' and B were
7 measured to calculate a discrimination index representing working memory according to the
8 following equation: Discrimination Index = (Novel Object A' exploration time - Familiar
9 Object B exploration time) / (Novel Object A' + Familiar Object B exploration times). The
10 OLT was performed under illumination at 10–15 lux.

11 The social interaction test was conducted as previously described (61) using a
12 specialized Sociability Apparatus (SC-03M, Muromachi). The time spent sniffing a novel
13 stimulus mouse or object was manually scored from videos recorded using an overhead
14 color USB camera (aa000080a02, Iroiro House). Stimulus mice (129Sv/SvJcl strain) age-
15 and sex-matched to test mice were habituated to the apparatus and to the enclosure cup for
16 30 min per day for 2 days prior to testing. The location (left or right) of the novel object and
17 novel mouse within an enclosure were alternated across test subjects. The test mouse was
18 allowed to acclimate to the apparatus for a total of 20 min before the sociability test—the

1 first 10 min in the central chamber with the doors closed and then 10 min in the empty arena
2 with the doors open. The test subject was briefly confined to the center chamber while a
3 novel stimulus mouse in an enclosure cup was placed on one of the side chamber and
4 another empty enclosure cup (novel object) was placed on the other side of the chamber.
5 The test subject was allowed to approach the novel object or mouse freely for 10 min. The
6 time spent interacting with the stimulus mouse versus the novel object was calculated as an
7 index of sociability.

8 The SR-LAB-Startle Response System (San Diego Instruments) was used to detect
9 acoustic startle reflexes and PPI. Startle responses were measured using 5 stimulus
10 intensities (80, 90, 100, 110, and 120 dB) delivered 10 times each for 40 ms over a white
11 noise background (65 dB). The stimuli were presented in quasi-random order at random
12 inter-trial intervals (10–20 s). In the PPI session, mice were exposed to 2 stimulus patterns:
13 1) a startle stimulus alone (120 dB, 40 ms) with no pre-pulse stimulus and 2) a startle
14 stimulus (120 dB, 40 ms) following a pre-pulse stimulus (70 dB for 20 ms; lead time, 100
15 ms). Each trial was repeated 10 times in quasi-random order at random inter-trial intervals
16 (10–20 s). PPI was defined as the percent decline in startle response because of pre-pulse
17 stimuli according to the following equation: $100 - [(120 \text{ dB startle amplitude after any}$

1 pre-pulse) / (120 dB startle amplitude only)] × 100. A new accelerometer was used for the
2 experiments with *Disc1* knockout mice.

3

4 **Immunohistochemistry**

5 Following transcardial perfusion with PBS and 4% paraformaldehyde, entire brains were
6 collected, post-fixed at 4°C overnight, and cryoprotected in 20% sucrose at 4°C overnight.

7 Serial coronal sections (50 μm) were then cut using a cryostat (CM3050 S; Leica

8 Microsystems). The antigens in the tissues were reactivated by heating in HistoVT One

9 solution (Nacalai Tesque) for 30 min at 70°C using a water bath. Sections were

10 permeabilized with 0.2% Triton X-100 and 1% Block Ace (DS Pharma Biomedical) in PBS

11 for 30 min at room temperature, following which they were incubated overnight with the

12 indicated primary antibodies at room temperature. For immunohistochemistry of

13 postmortem human brain tissues, paraffin blocks including BA9 (a region of frontal cortex)

14 were sliced into 10-μm sections, deparaffinized with xylene, and rehydrated with decreasing

15 concentrations of ethanol in water. Antigens were reactivated by heating in HistoVT One

16 solution for 30 min at 90°C using a water bath. Sections were treated with TrueBlack

17 Lipofuscin Autofluorescence Quencher (Biotium Inc.) for 30 s at room temperature and

18 blocked with 1% Block Ace (DS Pharma Biomedical) in PBS for 30 min at room

1 temperature. Thereafter, the mouse and human brain sections were subjected to the same
2 immunostaining procedures. The following primary antibodies were diluted in PBS
3 containing 0.4% Block Ace: goat anti-PV (Frontier Institute, PV-Go-Af860; 1:2000), mouse
4 anti-ALDH1L1 (Rockland, 600-101-HB6S; 1:200), FITC-conjugated tomato lectin
5 (VECTOR, FL-1171; 1:200), chick anti-GFP (Abcam, ab13970; 1:500), goat anti-IBA1
6 (Abcam, ab48004; 1:100), mouse anti-NeuN (Millipore, MAB377; 1:500), rabbit anti-AGE
7 (Abcam, ab23722; 1:2000), rabbit anti-IBA1 (Wako, WDJ3047; 1:300), rat anti-CD68
8 (Abcam, ab53444; 1:500), and rabbit anti-fibrin (Dako, A0080, 1:500). Thereafter, sections
9 were washed three times with PBS containing 0.05% Tween-20, incubated for 2 h with
10 fluorochrome-conjugated secondary antibodies in PBS containing 0.4% Block Ace, and
11 washed an additional three times in PBS containing 0.4% Block Ace. For enhanced
12 horseradish peroxidase (HRP) immunostaining, samples were treated with 3% H₂O₂ in PBS
13 for 20 min following the reactivation step to quench endogenous peroxidase activity and
14 washed in PBS. Sections were incubated with rabbit anti-GLO1 (Novusbio, NBP2-75514,
15 1:1500) and/or mouse anti-AGE4 (Trans Genic Inc, 14B5, 1:400), followed by incubation
16 with anti-IgG antibodies conjugated to biotin (Vector, 1:200). After washing as described
17 for other secondary antibodies, sections were incubated with streptavidin-conjugated HRP
18 (Jackson ImmunoResearch, 1:200) for 120 min and washed three times with PBS

1 containing 0.05% Tween-20. The TSA Plus Fluorescence System (PerkinElmer) was used
2 to detect HRP activity. All preparations were counterstained with DAPI (Nacalai Tesque) to
3 reveal cell nuclei, washed three additional times, mounted in Permaflow (Thermo
4 Scientific), and observed using a FluoView[®] FV3000 Confocal Laser Scanning Microscope
5 (Olympus).

6

7 **Image analysis**

8 Unless otherwise noted, all image analyses were performed using ImageJ version
9 2.0.0-rc-59/1.51n, and images were first binarized. PV-positive cells with a threshold
10 exceeding 30 were counted.

11 To measure the areas immunopositive for AGE, fibrin, and AGE-4, the threshold
12 settings were applied to ensure that only the correct immunopositive area was properly
13 selected, and the stained area of the entire screen was measured. For measuring the
14 fluorescence intensity of GFP and IBA1, after setting the ROI to select only target
15 immunoreactive cells, the intensity of >5 cells in each image were measured. To count
16 microglial cells containing a CD68-positive area exceeding $20 \mu\text{m}^2$, the threshold was set to
17 select only the CD68-positive area. Colocalization analysis of two different fluorescence
18 markers was performed using the Fiji plug-in Coloc 2 with the default settings. We adopted

1 Pearson's R value below the threshold to judge the strength of colocalization: $R \geq 0.7$,
2 strong; $0.7 > R \geq 0.4$, moderate; $0.4 > R \geq 0.2$, weak; and $R < 0.2$, none or very weak.

3

4 **Immunoblotting**

5 Extracts from mouse hippocampi were homogenized in lysis buffer containing 40 mM Tris
6 base, 0.4% sodium dodecyl sulfate (SDS), 0.01 M EDTA (pH 8.0), 8 M urea, and 1 mM
7 phenylmethylsulfonyl fluoride. The total lysate protein content was quantified using a DC
8 Protein Assay Kit (Bio-Rad). Total protein (30 μ g per gel lane) was separated using
9 SDS-PAGE and transferred to PVDF membranes (Millipore). Membranes were blocked
10 with TBST buffer (137m M NaCl, 2.7 mM KCl, and 25mM Tris, pH 8.0) including 0.2%
11 Triton X-100 and 5% bovine serum albumin (BSA) for 30 min at room temperature with
12 slow shaking, followed by incubation overnight with primary antibodies in TBST including
13 2% BSA at 4°C. The primary antibodies used were rabbit anti-GLO1 (Santa Cruz,
14 sc-67351; 1:1000), mouse-anti-PV (Swant, PV-235; 1:1000), rabbit anti-glucose transporter
15 1 (Glut1, Frontier Institute, Af1020; 1:1000) and mouse anti-tubulin (Santa Cruz, sc-32293;
16 1:10000). After washing three times with TBST, membranes were incubated with the
17 secondary antibody (HRP-conjugated anti-mouse or anti-rabbit IgG antibody, GE
18 Healthcare; 1:2000) in TBST plus 2% BSA. After washing three times with TBST, blots

1 were processed for chemiluminescence using standard protocols (ECL Prime Western
2 Blotting Detection Reagent #RPN2236, GE Healthcare), and signals were detected using a
3 LAS 4000 Imager (Fuji Film).

4

5 **Microdialysis**

6 We used an *in vivo* microdialysis system for the measurement of extracellular dopamine
7 concentration (62) and collection of brain parenchymal dialysate (Fig.1H, J and fig. S7D).
8 After anesthesia by intraperitoneal injection of ketamine (80 mg/kg)/xylazine (16 mg/kg),
9 mice were fixed in a stereotaxic apparatus (Narishige) and a microdialysis guide cannula
10 (CXG-8, Eicom) was implanted in the medial prefrontal cortex (mPFC; antero-posterior
11 (AP), +1.8 mm; medio-lateral (ML), ± 0.15 mm; dorso-ventral (DV), -1.5 mm from
12 bregma), or nucleus accumbens (NAc) (AP, +1.5 mm; ML, ± 0.6 mm; DV, -3.5 mm from
13 bregma). After recovery for at least 10 days, a microdialysis probe (CX-I-8-01 for the
14 mPFC and CX-I-8-02 for NAc; Eicom) was inserted via the guide cannula. Following
15 insertion, the probe was connected to a syringe pump and perfusion was performed at 2 μ
16 L/min for NAc and 0.5 μ L/min for mPFC using Ringer's solution (147 mM NaCl, 4 mM
17 KCl, and 2.3 mM CaCl₂). Dialysate samples were collected every 10 min and automatically
18 loaded onto an HTEC-500EPS HPLC unit (Eicom). Constant 5-HT concentration in three

1 consecutive collection periods was first confirmed to rule out blood contamination before
2 initiating the measurements of dopamine concentration or collection of parenchymal
3 dialysates. Analytes were then separated on an affinity column (PP-ODS III, Eicom), and
4 compounds were subjected to redox reactions within an electrochemical detection unit
5 (amperometric DC mode; applied potential range, 450 mV). The resulting chromatograms
6 were analyzed using an EPC-500 data processor (Eicom), and actual sample concentrations
7 were computed based on the peak heights obtained using 0.01, 0.1, and 1 pg dopamine in
8 standard solution (Sigma). The locations of the microdialysis probes were histologically
9 confirmed.

10 For glucose measurements in Fig. 4H and Fig. 5G, we collected dialysates from the
11 mPFC for 1 h after 16 h of fasting, followed by 0–1 h after eating 0.05 g of carbohydrates
12 (starch or sucrose), and 1–2 h after eating 0.05 g of carbohydrates (starch or sucrose).

13

14 **EEG recordings**

15 For behavioral and video/EEG monitoring, mice were anesthetized by an intraperitoneal
16 injection of ketamine (80 mg/kg)/xylazine (16 mg/kg), fixed in a stereotaxic apparatus
17 (Narishige, Japan), and implanted with EEG and electromyography (EMG) electrodes. The
18 EEG electrodes were gold-coated stainless steel screws (SUS303) soldered with lead wires

1 (ANE-0190, Adler's Nest, Japan) implanted epidurally over the left frontal cortex (AP, 1
2 mm; ML, 1 mm) and the bilateral parietal cortex (AP, -2 mm; ML, ± 2 mm). All wires
3 were soldered to a multichannel electrical connector (R833-83-006-001, TOKIWA SHOKO,
4 Japan). The left parietal cortex electrode was used as a reference for monitoring the frontal
5 cortex EEG. The EMG electrodes were lead wires placed bilaterally into the trapezius
6 muscle. Following recovery for at least 10 days, EEG/EMG signals were amplified and
7 band-pass filtered (EEG: 1.5–1000 Hz; EGM: 15–3000 Hz) using a MEG-6116 system
8 (NIHON KOHDEN), digitized at a sampling rate of 200 Hz, recorded using a data
9 acquisition system (PowerLab 8/30, ADInstruments), and analyzed using LabChart
10 Software (ADInstruments). Behavioral activities were recorded using a USB camera
11 (aa000080a02, Iroiro House, Japan). Behavioral and electrophysiological responses to a
12 novel object (an empty 100 mL DURAN bin) were recorded in an OF chamber (20 × 20 ×
13 26 cm). The novel object was placed in one corner of the OF chamber to induce exploration.
14 The 30 s preceding the first contact with the novel object was analyzed for object
15 recognition ("object activity"). For EEG monitoring in the home cage, mice were first
16 habituated for 8 h. Home cage EEG data were then acquired for 2 min after awaking as
17 confirmed by clear EMG signals and movement images from an offline video camera
18 analysis ("home cage activity"). All recordings were converted into power spectra using a

1 fast Fourier transform (FFT) algorithm with a 5-s Hann cosine-bell window and 50%
2 overlap between successive window measurements. All FFTs were maintained at 1024
3 points to obtain 0.512 Hz resolution. The total signal amplitude or power (V^2) in each 5-s
4 period was measured as the power magnitude at each frequency. The mean of the grouped
5 power spectra was calculated over the following frequency ranges: 1–4 Hz (delta), 5–10 Hz
6 (theta), 30–45 Hz (low gamma), and 55–80 Hz (high gamma). The power values detected at
7 each frequency range for 30 s were further averaged over 30 s of total EEG power using the
8 mean values to remove potential noise. These analyses were performed using custom
9 software written in MATLAB (R2019b; MathWorks).

10

11 **Transcriptome analysis**

12 Three independent total RNA samples from each group were mixed and purified using a
13 RNeasy Mini Kit (Qiagen). RNA quality was assessed using a 2100 bioanalyzer (Agilent
14 Technologies). Cy3-labeled cRNA was prepared using a Low Input Quick Amp Labeling
15 Kit (Agilent Technologies), in accordance with the manufacturer's protocol. Samples were
16 hybridized to the SurePrint G3 Mouse Gene Expression v2 Microarray (G4852B; Agilent
17 Technologies). Thereafter, the array was washed and scanned using the SureScan
18 Microarray Scanner (Agilent Technologies). Microarray images were analyzed using the

1 Feature Extraction software with default settings for all parameters (Agilent Technologies).
2 Data from each microarray analysis were normalized by shift to the 75th percentile without
3 baseline transformation. Microarray results were deposited in the Gene Expression
4 Omnibus database under the accession number GSE141829.

5

6 **Insulin and glucose measurements**

7 Blood plasma was collected from the mouse cheek as described by Golde (63). Plasma
8 glucose concentration was measured using a Precision-Neo blood glucose meter (#71386-80,
9 Abbott Japan), plasma insulin concentration using an ELISA kit (#M1102, MORINAGA),
10 and glucose concentration in the dialysate samples using a different ELISA kit (#ab65333,
11 Abcam), all according to the manufacturers' instructions. Data were collected on a
12 microplate reader (Varioskan, Thermo Fisher Scientific).

13

14 **Human postmortem brain tissue collection**

15 Postmortem brain tissues from patients with SZ and BD were obtained from the Fukushima
16 Brain Bank at the Department of Neuropsychiatry, Fukushima Medical University.
17 Postmortem brain tissues from control individuals were obtained from the Section of
18 Pathology, Fukushima Medical University Hospital. The use of postmortem human brain

1 tissues in the present study was approved by the Ethics Committee of Fukushima Medical
2 University (No.1685) and Tokyo Metropolitan Institute of Medical Science (No. 18-20) and
3 complied with the Declaration of Helsinki and its later amendments. All procedures were
4 conducted with the informed written consent of the next of kin. Detailed demographic
5 information of the 10 patients with SZ, 9 patients with BD, and the 12 age- and sex-matched
6 control individuals is provided in Supplementary Table 3. No between-group differences
7 were observed in terms of sex (Fisher's exact test, Ctrl and SZ: $p = 1.00$, Ctrl and BD: $p =$
8 0.40), age (Student's t-test, Ctrl and SZ: $p = 0.69$, Welch's t-test, Ctrl and BD: $p = 0.66$),
9 postmortem interval (Student's t-test, Ctrl and SZ: $p = 0.89$, Ctrl and BD: $p = 0.98$), or a
10 history of diabetes mellitus (Fisher's exact test, Ctrl and SZ: $p = 0.59$, Ctrl and BD: $p =$
11 0.59). Each patient with SZ and BD fulfilled the diagnostic criteria established by the
12 American Psychiatric Association (Diagnostic and Statistical Manual of Mental Disorders,
13 DSM-IV) and did not have a past history of other neurological disorders or substance abuse.
14 Moreover, none of the control individuals had any record of mental disorders, neurological
15 disorders, or substance abuse.

16

17 **ROS detection**

1 DHE is oxidized by superoxide and other reactive species to produce a precipitate that emits
2 red fluorescence, whose intensity can be used to measure the degree of cellular oxidative
3 stress. Animals were quickly perfused with ice-cold phosphate-buffered salts (PBS), and the
4 decapitated brains were embedded in OCT compound (Sakura Finetek Japan), and 10 μm
5 thick coronal sections were prepared in a cryostat. These sections were incubated with
6 phosphate buffer containing 100 μM diethylene triamine pentaacetic acid (PB/DTPA) for 10
7 min, and then incubated with PB/DTPA containing 10 μM DHE (Invitrogen) for 20 min.
8 After washing with PB/DTPA twice, sections were fixed with 4% PFA for 15 min. Once
9 washed with PBS, PBS containing 1 μM TO-PRO 3 (Thermo Scientific) was applied to
10 each section followed by 3 washes with PBS. Then, sections were sealed with Permaflow
11 (Thermo Scientific) and observed using a FluoView® FV3000 Confocal Laser Scanning
12 Microscope (Olympus). For DHE detection, we used the following filter set: excitation 405
13 nm, emission 600–650 nm.

14

15 **Statistical analyses**

16 Statistical differences among ≥ 4 groups were determined using one-way analysis of variance
17 (ANOVA), two-way ANOVA, three-way ANOVA, or repeated measures ANOVA, followed
18 by the Bonferroni multiple comparison test or Tukey-Kramer test as a post hoc test, as

1 summarized in Supplementary Table 5. We used the same animals for all results coming from
2 Starch +/+ and Suc *Glo1*/+ groups according to the 3R rule. Data were collected at the same
3 time regardless of whether the drug was administered or not. The number of animals
4 employed for each experiment/analysis can be found in each figure legend. Detailed
5 descriptions of statistical analyses in Fig. 1H, J, and 4H are shown below. In Figure 1H, post
6 hoc Dunnett's multiple comparisons test for groups was performed at specific time points (0
7 min and 40 min), $^{###}p < 0.001$ for Suc *Glo1*/+ vs. Ctrl (Starch +/+), $^{\$}p < 0.05$ for Starch *Glo1*/+
8 vs. Ctrl (Starch +/+). In Fig. 1J, post hoc Bonferroni's multiple comparisons test for groups
9 was performed at specific time points (0 min and 40 min), $^{###}p < 0.001$ for Suc *Glo1*/+ vs.
10 Starch +/+, $^{$$$}p < 0.001$ for Suc *Glo1*/+ vs. Starch +/+ Aripiprazole, $^{\&\&\&}p < 0.001$ for Suc *Glo1*/+ vs.
11 Suc *Glo1*/+ Aripiprazole. In Fig. 4H, post hoc Dunnett's multiple comparisons test for groups was
12 performed at each time point, $^{###}p < 0.001$, $^{##}p < 0.01$ for Suc *Glo1*/+ vs. Ctrl (Starch +/+).

13

14 **Supplementary Figure Legends**

15 **Supplementary Figure 1. Characterization of regional GLO1 expression in WT and**
16 **heterozygous *Glo1* mutant mice fed starch or sucrose.**

17 (A) Western blot analysis of GLO1 protein expression using tubulin as internal control. The
18 cerebral cortex, including the hippocampus, was used as loading sample. (B) Densitometric

1 analysis of GLO1 protein expression (n = 3 or 4 mice per group). To quantify expression,
2 GLO1 band intensities in (A) were divided by the intensities of the corresponding tubulin
3 bands. (C) Body weight trajectories. No significant differences were observed among
4 groups (n = 6–10 mice per group). (D, E) Measurement of aripiprazole- or
5 aspirin-containing water consumption per day to estimate drug intake (n = 4 mice per
6 group). Student's *t*-test was used for statistical analysis. All data are presented as mean ±
7 SEM. ***p* < 0.01, **p* < 0.05

8

9 **Supplementary Figure 2. Analyses of psychiatric disease-related phenotypes in *Glo1***
10 **heterozygotic mice.**

11 (A, B) Behavioral analyses in the four groups of mice (n = 18–23 mice per group). (A)
12 Elevated plus maze test to evaluate anxiety. (B) Preference for novel mouse in the
13 three-chamber test, calculated as [(time spent exploring novel mouse)/(total time spent
14 exploring novel mouse and novel object)] × 100%. (C) Extracellular dopamine
15 concentration in the medial prefrontal cortex measured at 20-min intervals using an *in vivo*
16 microdialysis system (n = 7–10 mice per group). (D–I) Effect of aripiprazole treatment on
17 behavioral phenotypes (n = 12–18 mice per group). (D) Effects of aripiprazole treatment on
18 the acoustic startle response, (E) pre-pulse inhibition (PPI) using a 70-dB pre-pulse, (F) object

1 location test, (G) self-grooming, (H) elevated plus maze performance, and (I) nest-building
2 skill. In (I), post hoc Tukey's multiple comparisons test of groups at specific time points, ^{###}*p*
3 < 0.001, [#]*p* < 0.05 for Starch +/+ vs. Suc *Glo1*/+ Aripri, ^{\$\$}*p* < 0.01 for Starch +/+ Aripri vs. Suc
4 *Glo1*/+ Aripri, ^{\$\$\$}*p* < 0.001 for Starch +/+ vs. Suc *Glo1*/+, **p* < 0.05 for Starch +/+ Aripri vs.
5 Suc *Glo1*/+. All data are presented as mean ± SEM. ****p* < 0.001, ***p* < 0.01, **p* < 0.05

6

7 **Supplementary Figure 3. Analyses of psychiatric disease-related phenotypes in *Disc1***

8 **heterozygotic mice.**

9 (A-E) Behavioral analyses in the four groups of mice (n = 8–11 mice per group). (A)

10 Spontaneous locomotor activity in the open-field test. (B) Acoustic startle responses. (C)

11 Pre-pulse inhibition (PPI) using a 70 dB pre-pulse. (D) Object location test. (E) Preference for

12 novel mouse in the three-chamber test, calculated as fig. S2B. All data are presented as mean

13 ± SEM. ****p* < 0.001

14

15 **Supplementary Figure 4. GLO1 cellular localization in the prefrontal cortex of**

16 **sucrose-fed wild-type mice.**

17 (A-F) GLO1 localization in astrocytes. (A) GLO1 co-immunostaining with an astrocyte

18 marker (ALDH1L1) or an endothelial cell marker (tomato lectin). (B) Merged

1 GLO1/ALDH1L1 image from (A). Yellow arrows in (B) indicate cells with
2 GLO1/ALDH1L1 colocalization. (C) Merged GLO1/lectin image from (A). White arrows
3 denote representative GLO1-positive cells close to endothelial cells. (D, E) High
4 magnification images of GLO1 co-immunostaining with ALDH1L1 or with tomato lectin.
5 (E) Merged GLO1/ALDH1L1 image from (D). (F) 2D intensity histogram of the two
6 indicated channels to identify GLO1 colocalization with ALDH1L1 or tomato lectin,
7 interpreting the strength of a relationship based on R as follows: $R \geq 0.7$, strong; $0.7 > R \geq$
8 0.4 , moderate; $0.4 > R \geq 0.2$, weak; and $R < 0.2$, none or very weak. (G) Average R-value
9 of colocalization data including (F) from three different locations of three mice. (H) GLO1
10 co-immunostaining with the neuronal marker NeuN and the microglial marker IBA1. (I)
11 Merged GLO1/NeuN image from (H). Yellow arrows indicate neurons with mild GLO1
12 immunoreactivity. (J) Merged GLO1/IBA1 image from (H). (K) GLO1 immunostaining
13 and DAPI staining in *Glo1* homozygous mice.

14

15 **Supplementary Figure 5. Fructose-derived AGE4 accumulation in microglia of**
16 **sucrose-fed mice.**

17 (A, B) Immunohistochemical images of the microglial marker IBA1 and AGE4 in the
18 medial prefrontal cortex region (mPFC). (C) Merged AGE4/IBA1 image. (D) High

1 magnification images of AGE4 co-immunostaining with IBA1. (E) Measurement of the
2 area of AGE-4 immunoexpression in (B). The mean AGE4-immunopositive area intensity
3 of the entire image was measured in each section (n = 4 mice per group). (F) Mean IBA1
4 fluorescence intensities of five randomly selected cells per image in four independent mice.
5 (G) Immunohistological images of IBA1- and CD68-positive microglia in the mPFC region.
6 White arrows in (G) indicate microglial cells containing CD68-positive areas $>20 \mu\text{m}^2$. (H)
7 Percentage of CD68-positive cells per total microglial cells in each image in (G). All data
8 are presented as mean \pm SEM. $***p < 0.001$, $**p < 0.01$

9

10 **Supplementary Figure 6. Blood vessels Characterization in the four groups.**

11 (A) Western blot analysis of glucose transporter 1 (Glut1) protein expression using tubulin
12 as internal control. The cerebral cortex, including the hippocampus, was used as loading
13 sample. (B) Densitometric analysis of Glut1 protein expression (n = 3 mice per group). To
14 quantify expression, Glut1 band intensities in (A) were divided by the intensities of the
15 corresponding tubulin bands. (C) Measurement of capillary diameter. Vessels $<10 \mu\text{m}$ in
16 diameter were defined as capillaries, and the mean diameter of 10 randomly selected
17 capillaries per image measured in Fig. 4C (n = 3 mice per group). All data are presented as
18 mean \pm SEM.

1

2 **Supplementary Figure 7. Protective effects of aspirin against the development of**

3 **psychiatric phenotypes in G × E mice.**

4 (A–C) Effects of aspirin treatment on the acoustic startle response (A), nest-building skill

5 (B), elevated plus maze test performance (C) and dopamine concentration (D). (E)

6 Parvalbumin (PV) Immunohistochemistry in the medial prefrontal cortex (mPFC). (F)

7 Number of PV-positive cells in the mPFC (n = 4 mice per group). (G) Immunohistological

8 images of GFP-positive astrocytes and ALDH1L1 in the mPFC region. (H) Percentage of

9 GFP-positive cells per total ALDH1L1-positive cells in each image in (G). (I) Mean

10 fluorescence GFP intensities of 10 randomly selected cells per image in (G) (from four

11 independent mice). In (B), post hoc Tukey's multiple comparisons test of groups at specific

12 time points, $^{###}p < 0.001$, $^{##}p < 0.01$ for Starch +/+ vs. Suc *Glo1*/+ Aspi, $^{$$}p < 0.01$, $^{$}p <$

13 0.05 for Starch +/+ Aspi vs. Suc *Glo1*/+ Aspi. In (D), post hoc Bonferroni's multiple

14 comparisons test of groups at specific time points (0 min and 40 min), $^{###}p < 0.001$ for Suc

15 *Glo1*/+ vs. Starch +/+, $^{$$$}p < 0.001$ for Suc *Glo1*/+ vs. Starch +/+ Aspi, $^{&&&}p < 0.001$ for

16 Suc *Glo1*/+ Aspi vs. Starch +/+, $^{††}p < 0.01$ for Suc *Glo1*/+ Aspi vs. Suc *Glo1*/+, $^{‡‡}p < 0.01$

17 for Suc *Glo1*/+ Aspi vs. Starch +/+ Aspi. All data are presented as mean ± SEM. $^{***}p <$

18 0.001 , $^{**}p < 0.01$, $^{*}p < 0.05$

1

2 **Supplementary Figure 8. ROS detection in the four groups.**

3 (A) Reactive oxygen species (ROS) detection with *dihydroethidium* (DHE). (B) Merged
4 DHE/Nucleus (TO-PRO 3) image. All cells are DHE positive, according to nucleus images.
5 (C) ROS detection images without DHE. (D) Fluorescent intensity of DHE positive cells in
6 (A). The mean DHE fluorescent intensity of each cell was measured in each section (n = 4
7 mice per group). All data are presented as mean \pm SEM.

8

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4

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Figure 1

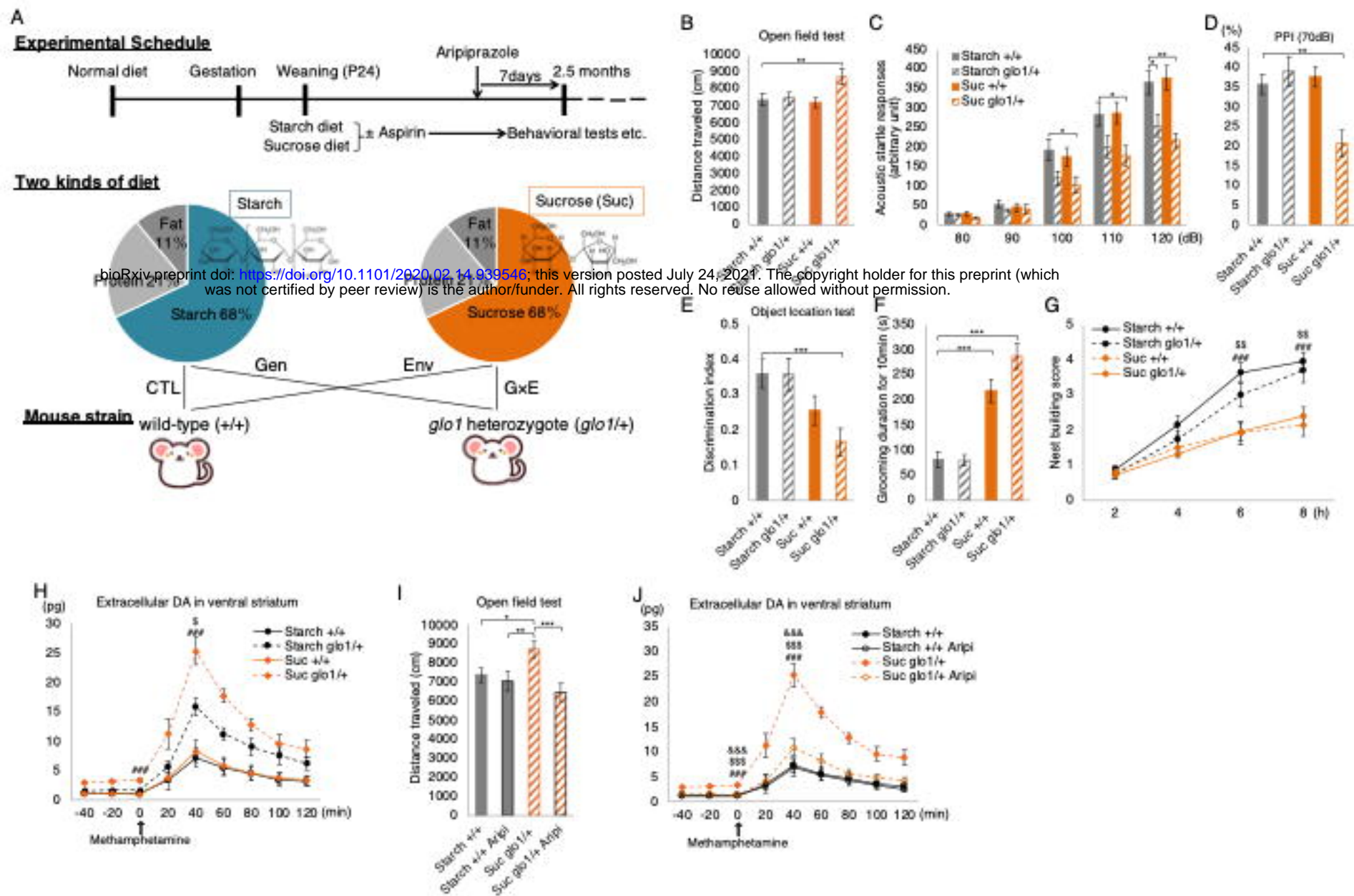


Fig. 1 Generation of G × E model mice and psychosis-related phenotype analyses.

(A) Experimental Timeline. See Materials and Methods for a complete description. (B–G) Behavioral analyses in the four groups of mice ($n = 18–23$ mice per group). (B) Spontaneous locomotor activity in the open-field test. (C) Acoustic startle responses. (D) Pre-pulse inhibition (PPI) using a 70 dB pre-pulse. (E) Object location test to evaluate working memory. (F) Duration of self-grooming in the home cage. (G) Quantification of nest-building skills over 8 h. Post hoc Dunnett's multiple comparisons test among groups at each time point indicated $###p < 0.001$ for Suc *glo1*^{+/+} vs. Ctrl (Starch *+/+*), $##p < 0.01$ for Suc *+/+* vs. Ctrl (Starch *+/+*). (H, J) Extracellular dopamine (DA) concentration in the nucleus accumbens measured at 20 min intervals using an *in vivo* microdialysis system in the presence or absence of aripiprazole (Aripip). Methamphetamine (1.25 mg/kg) was administered via intraperitoneal (i.p.) injection at time 0 (arrow) ($n = 6–11$ mice per group). See Materials and Methods for detailed statistical analyses. (I) Effects of Aripip treatment on locomotor activity ($n = 12–18$ mice per group). (B–H) Starch *+/+* group was used as a control for post hoc Dunnett's test. All data are presented as mean \pm SEM. $***p < 0.001$, $**p < 0.01$, $*p < 0.05$

Figure 2

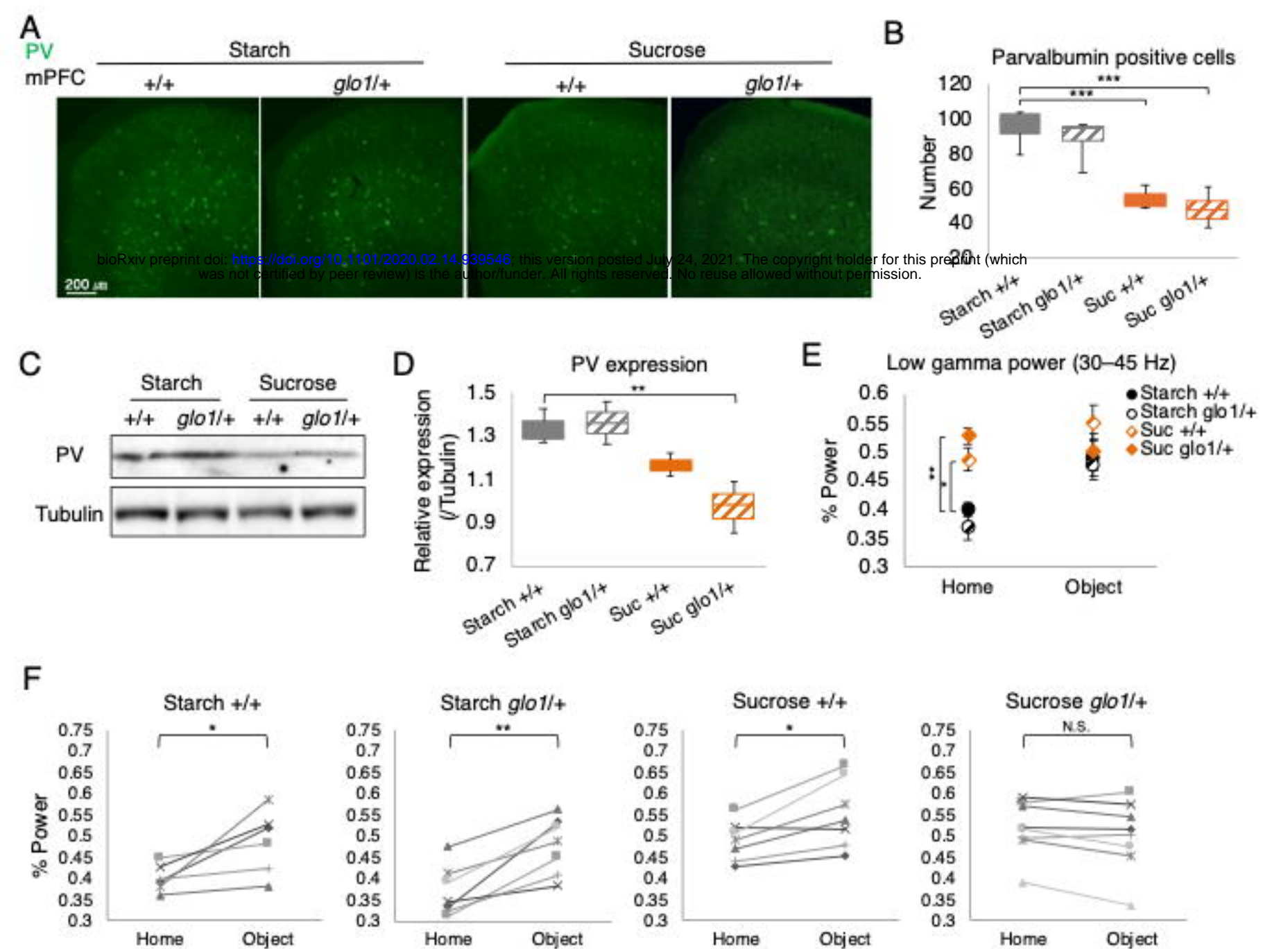


Fig. 2 Parvalbumin-positive interneuron dysfunction in G × E mice.

(A) Parvalbumin (PV) immunohistochemistry in the medial prefrontal cortex (mPFC). (B) Number of PV-positive cells in the mPFC ($n = 4$ mice per group). (C) Western blot analysis of PV protein expression using tubulin as internal control. (D) Densitometric analysis of PV protein expression: PV band intensities were divided by the corresponding tubulin band intensities ($n = 3$ mice per group). (E) Average gamma band power in the home cage and during novel object recognition in the open field ($n = 7-8$ mice per group). (F) Changes in gamma band power from the home cage to the novel object phase in individual mice ($n = 7-8$ mice per group). Each p value indicates the result of repeated measures analysis of variance (ANOVA). The effect of changes in gamma power at Starch $+/+$ ($F_{1, 5} = 8.29, p = 0.035$), at Starch $glo1/+$ ($F_{1, 6} = 29.75, p = 0.0016$), at Sucrose $+/+$ ($F_{1, 6} = 12.04, p = 0.013$), at Sucrose $glo1/+$ ($F_{1, 7} = 4.038, p = 0.084$). (B, D, H) Starch $+/+$ group was used as control for post hoc Dunnett's test. All data are presented as mean \pm SEM. *** $p < 0.001$, ** $p < 0.01$, * $p < 0.05$

Figure 3

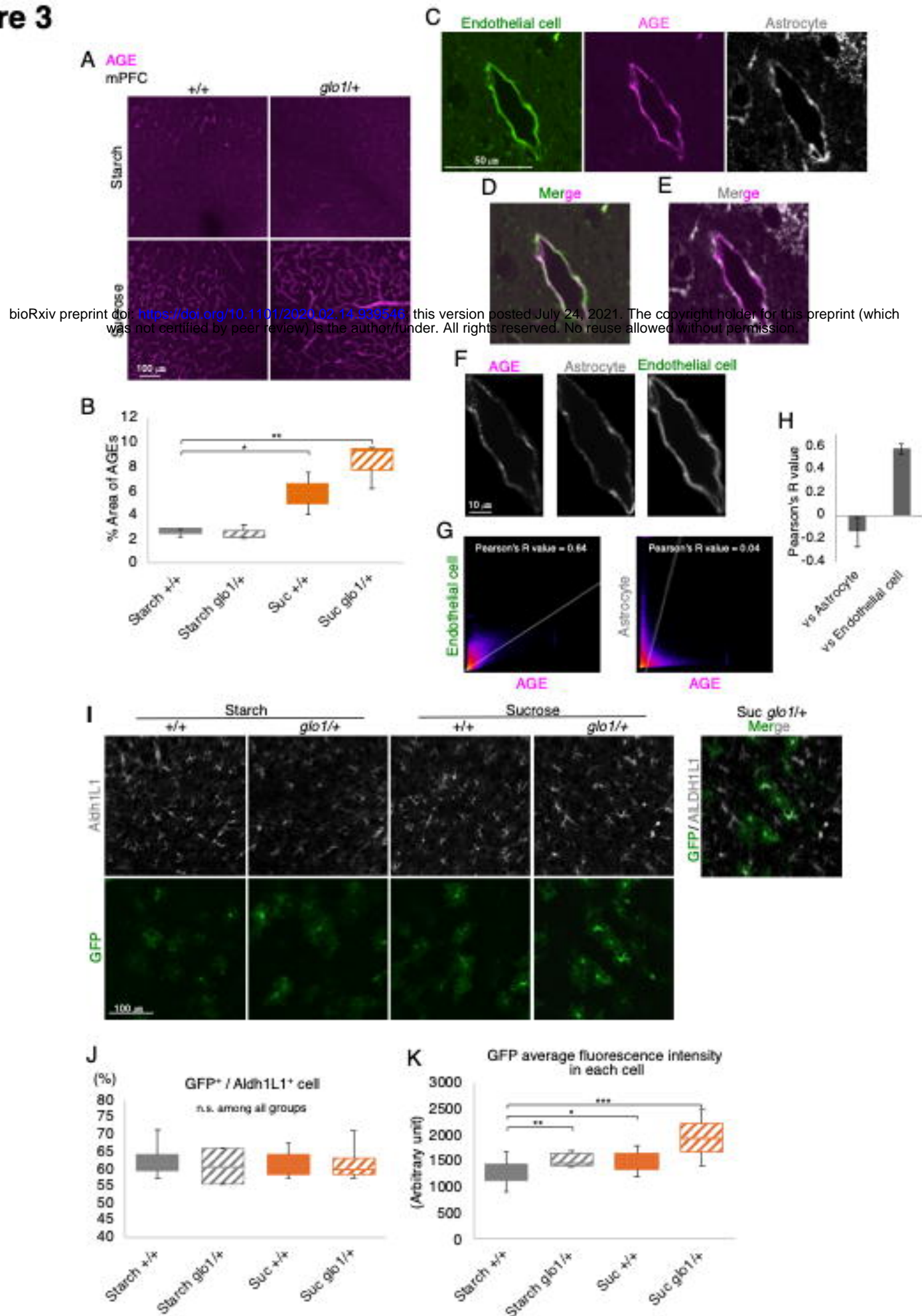
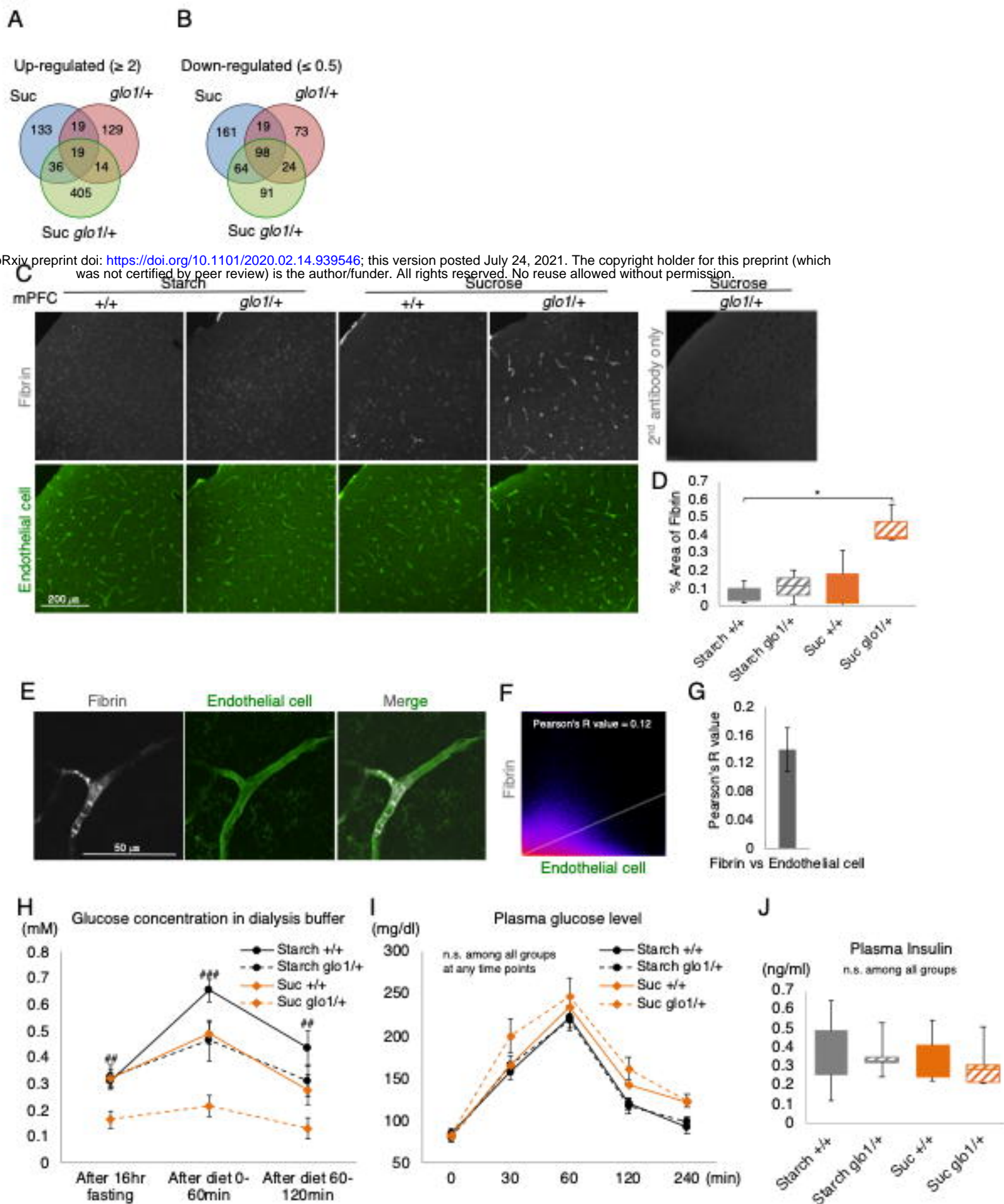


Fig. 3 AGE accumulation in the neurovascular endothelium and pre-inflammatory status of astrocytes in $G \times E$ mice.

(A) AGE immunohistochemistry in the medial prefrontal cortex. (B) Measurement of AGE immunoreactive area. The mean intensity of the AGE-immunopositive area in the entire image was measured for each section ($n = 3$ mice per group). (C–E) Colocalization of the endothelial cell marker tomato lectin with the astrocytic marker ALDH1L1 or AGEs. (F) Enlarged version of images in (C) presenting the region used for colocalization analysis. (G) 2D intensity histogram of the two indicated channels for identifying the colocalization of AGE with ALDH1L1 (Astrocyte) or tomato lectin (Endothelial cell). (H) Average R-value of colocalization data including (G) three different locations of three mice. (I) Immunohistological images of GFP-positive astrocytes and ALDH1L1 in the mPFC region and merged GFP/ALDH1L1 image. (J) Percentage of GFP-positive cells per total ALDH1L1-positive cells in each image in (I). (K) Mean fluorescent GFP intensities of 10 randomly selected cells per image in (I) (from four independent mice). (B, J, K) The Starch +/+ group was used as control for post hoc Dunnett's test. All data are presented as mean \pm SEM. *** $p < 0.001$, ** $p < 0.01$, * $p < 0.05$

Figure 4



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Fig. 4 Angiopathy and impaired glucose transport in G x E mice.

(A, B) Venn graph showing overlap in prefrontal cortex genes exhibiting a >2 -fold (A) or <0.5 -fold (B) expression change compared with the control (CTL) group ($n = 3$ mice per group). (C) Immunohistochemical images of fibrin and the endothelial cell marker tomato lectin. (D) Measurement of the area of fibrin immunoexpression in (C). The mean intensity of the fibrin-immunopositive area of the entire image was measured for each section ($n = 4$ mice per group). (E) Magnified immunohistochemical images of fibrin and the endothelial cell marker tomato lectin. (F) 2D intensity histogram of the two indicated channels using the left and middle panels in (E) for identifying fibrin colocalization with tomato lectin. (G) Average R-value of colocalization data in (F) from three different locations of three mice. (H) Extracellular glucose concentrations in the dialysis buffer from mPFC samples at each time point ($n = 5-6$ mice per group). See the Materials and Methods for detailed statistical analyses. (I) Plasma glucose concentrations ($n = 6-7$ mice per group). The first measurement was performed after 16 h of fasting, and the second blood collection 30 min after eating 0.05 g of carbohydrates. (J) Fasting plasma insulin levels ($n = 5-6$ mice per group). (D, H-J) Starch $+/+$ group was used as a control for post hoc Dunnett's test. All data are presented as mean \pm SEM. * $p < 0.05$

Remark: Please clarify the difference with B) in this same figure.

Figure 5

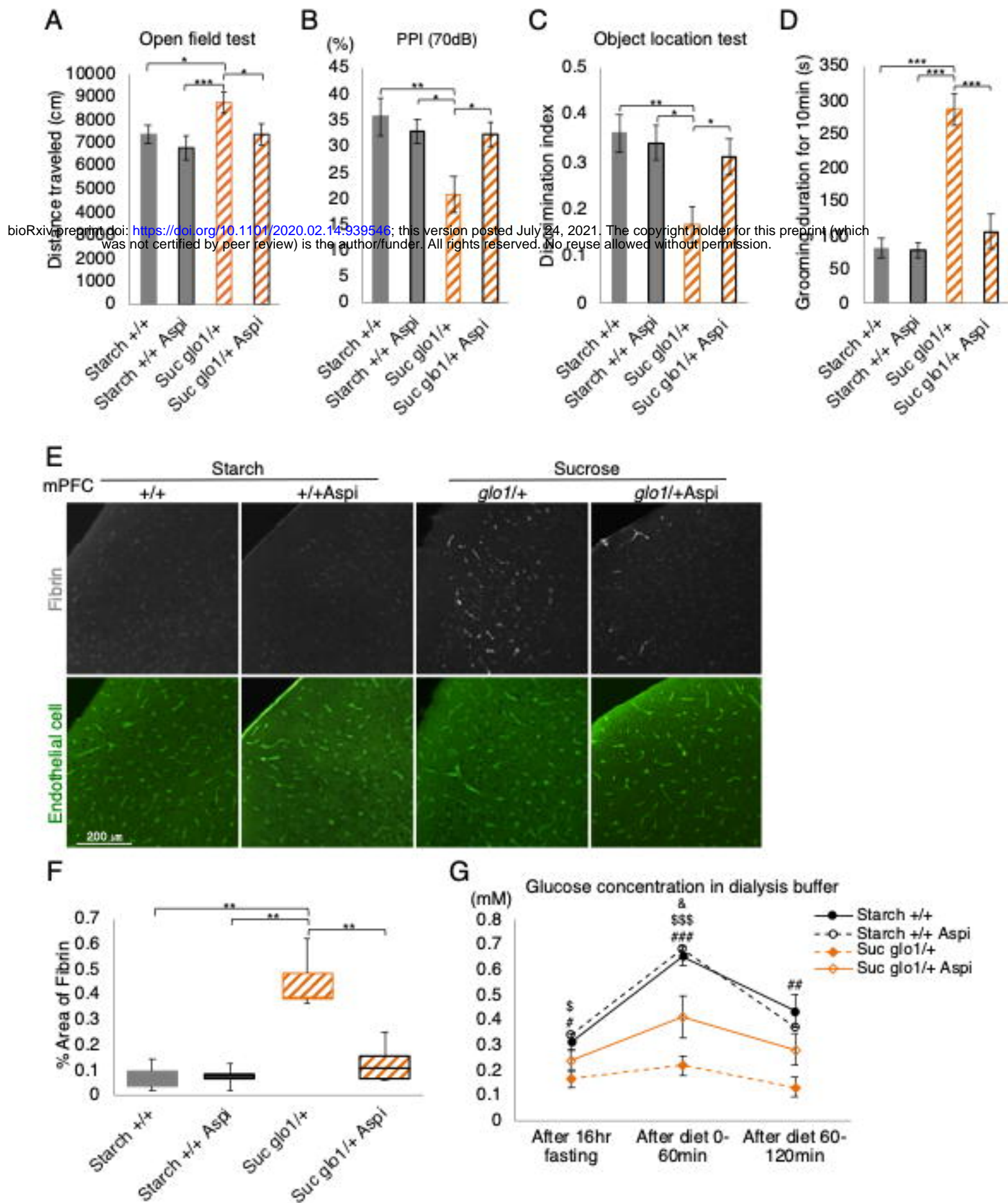


Fig. 5 Protective effects of low-dose aspirin in G × E mice.

(A–D) Results of behavioral tests performed to evaluate the effects of aspirin treatment (n = 12–21 mice per group). (A) Quantification of spontaneous locomotor activity in the open field. (B) Pre-pulse inhibition at 70 dB. (C) Object location test of working memory. (D) Quantification of nest-building skills over 8 h. (E) Immunohistochemical images of fibrin and the endothelial cell marker tomato lectin. (F) Measurement of the area of fibrin immunoexpression in (E). The mean intensity of the fibrin-immunopositive area of the entire image was measured for each section (n = 3 mice per group). (G) Extracellular glucose concentrations in the dialysis buffer at each time point (n = 4–6 mice per group). Post hoc Tukey's multiple comparisons test of groups at each time point, $^{***}p < 0.001$, $^{**}p < 0.01$, $^{*}p < 0.05$ for Starch +/+ vs. Suc glo1/+, $^{sss}p < 0.001$, $^{s}p < 0.05$ for Starch +/+ Aspi vs. Suc glo1/+, $^{&}p < 0.05$ for Starch +/+ vs. Suc glo1/+ Aspi. All data are presented as mean ± SEM. $^{***}p < 0.001$, $^{**}p < 0.01$, $^{*}p < 0.05$

Figure 6

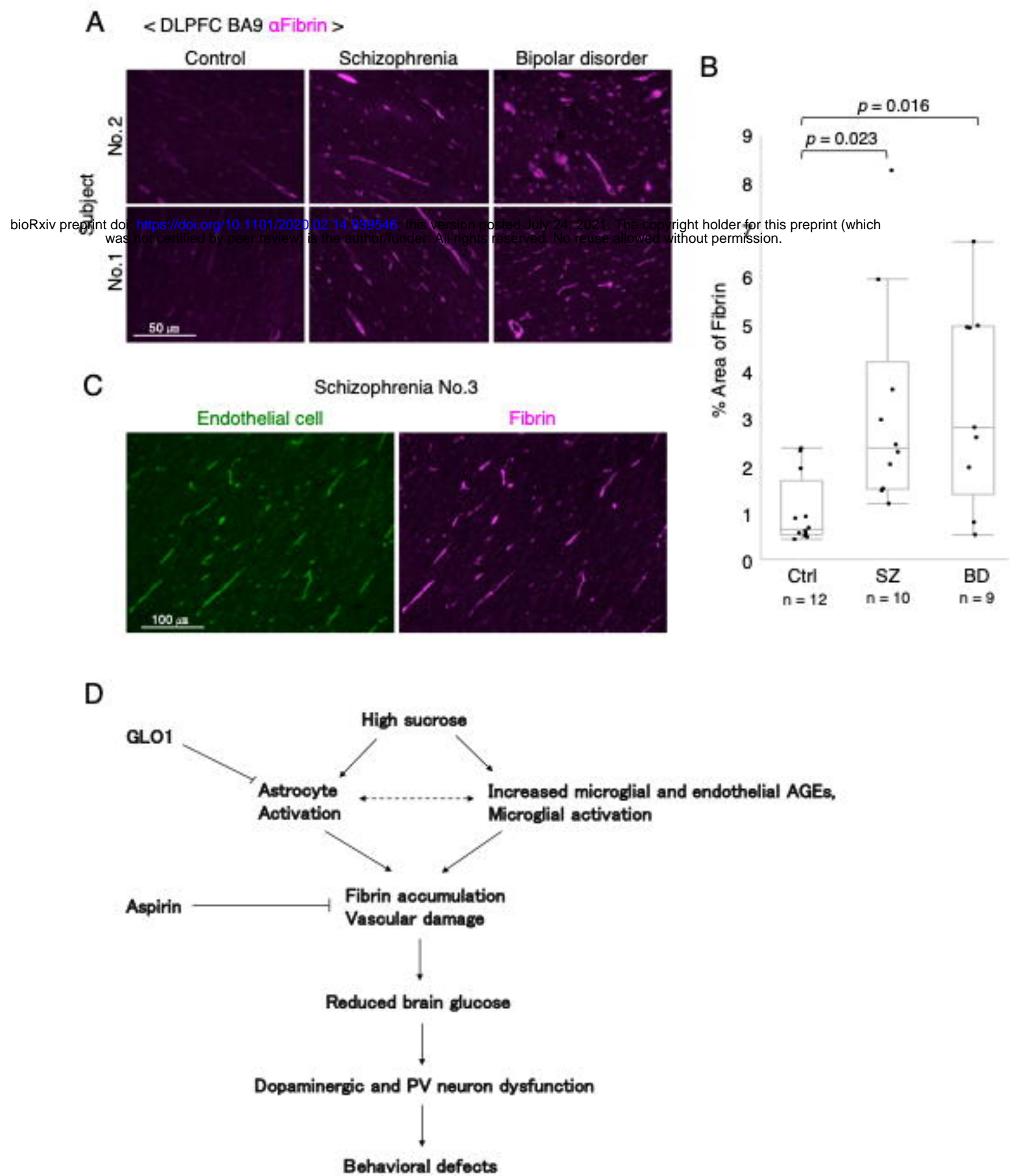
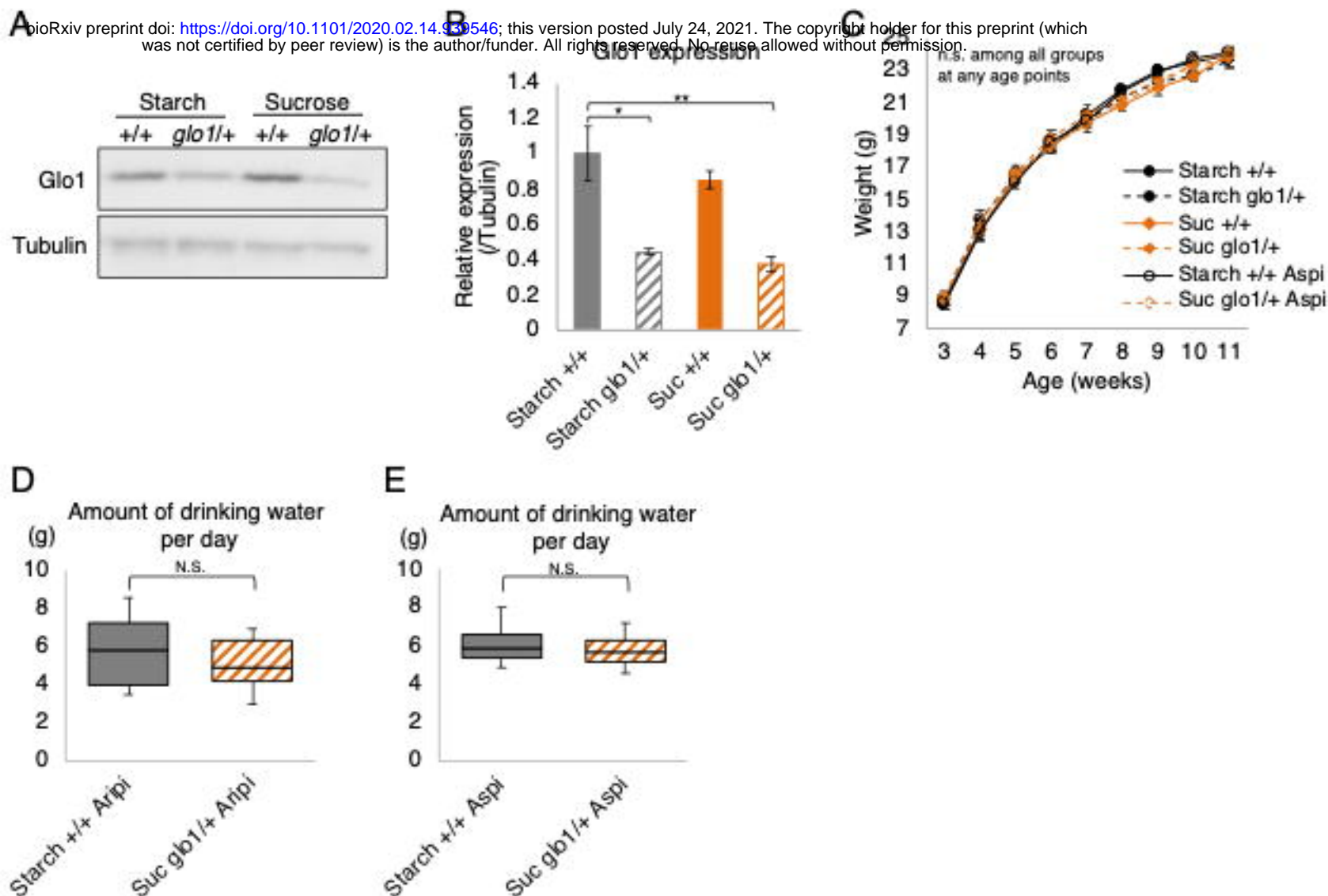


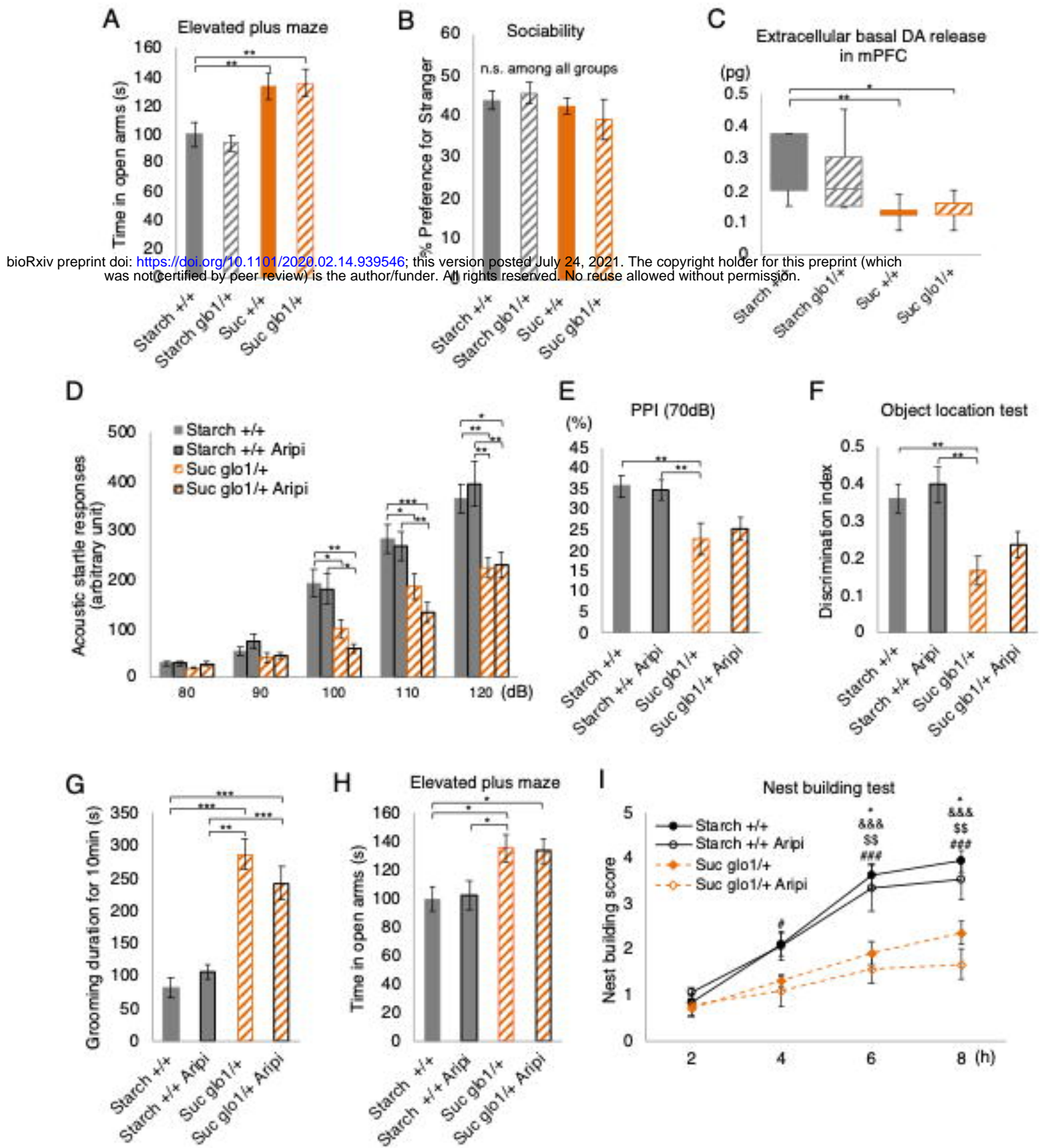
Fig. 6 Angiopathy in postmortem brains from individuals with psychiatric disorders.

(A) Representative immunohistochemical fibrin images in the BA9 region of postmortem brains from controls and patients with schizophrenia (SZ) or bipolar disorder. (B) Measurement of the area of fibrin immunopositive area in (A). The mean intensity of the fibrin-immunopositive area of the entire image was measured for each section. (C) Representative immunohistochemical images of fibrin (magenta) and the endothelial cell marker tomato lectin (green) in postmortem brains from a patient with SZ. Fibrin-positive areas are merged with areas of vascular endothelial cell marker expression. (D) Diagrams describing the hypothesis proposed to explain functional and behavioral abnormalities in control (CTL) mice (left) and G \times E mice (right) (see the Discussion for details). All data are presented as mean \pm SEM. ** $p < 0.01$



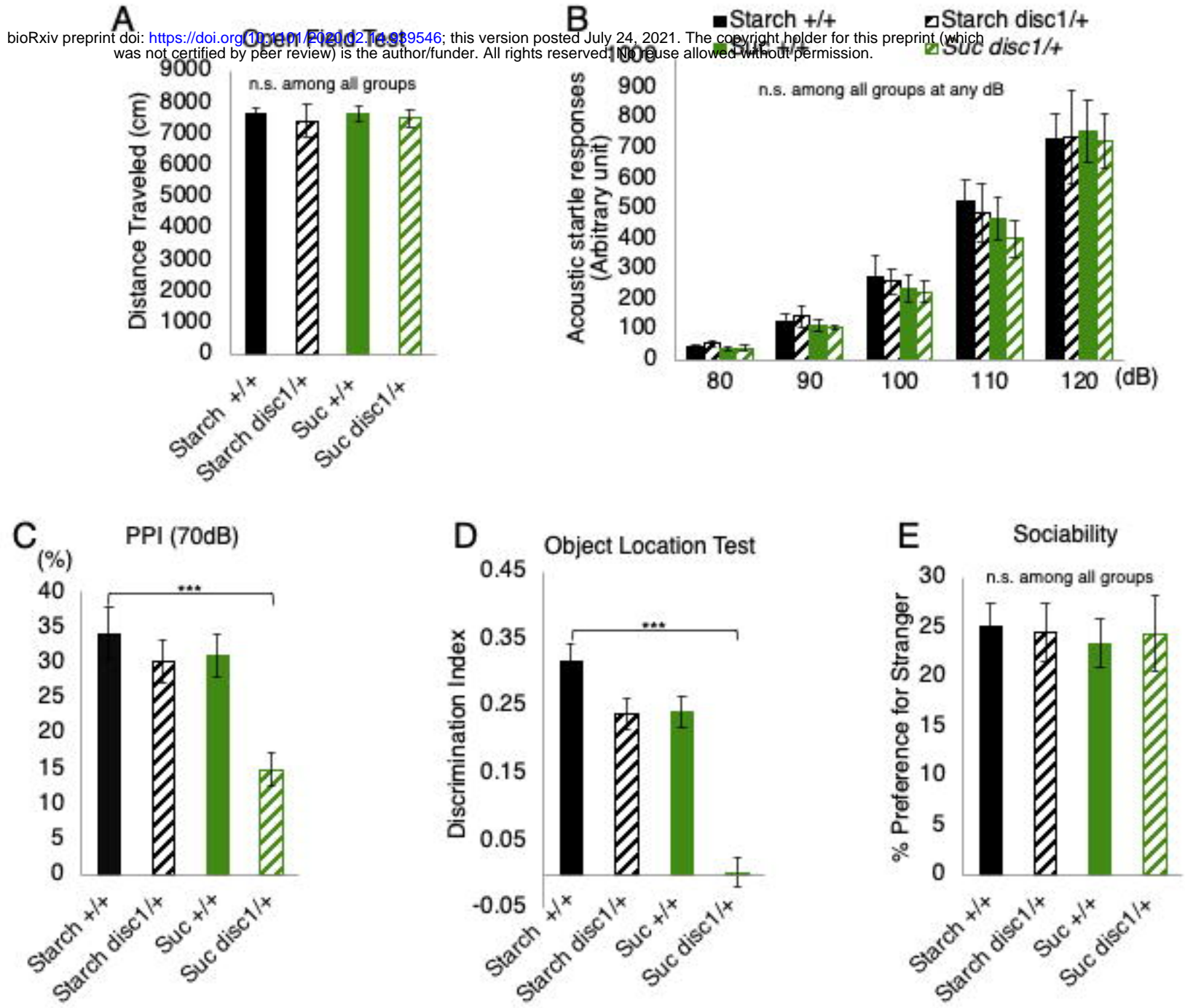
Supplementary Figure 1 Characterization of regional GLO1 expression in WT and heterozygous *Glo1* mutant mice fed starch or sucrose.

(A) Western blot analysis of GLO1 protein expression using tubulin as internal control. The cerebral cortex, including the hippocampus, was used as loading sample. (B) Densitometric analysis of GLO1 protein expression (n = 3 or 4 mice per group). To quantify expression, GLO1 band intensities in (A) were divided by the intensities of the corresponding tubulin bands. (C) Body weight trajectories. No significant differences were observed among groups (n = 6–10 mice per group). (D, E) Measurement of aripiprazole- or aspirin-containing water consumption per day to estimate drug intake (n = 4 mice per group). Student's *t*-test was used for statistical analysis. All data are presented as mean \pm SEM. ***p* < 0.01, **p* < 0.05



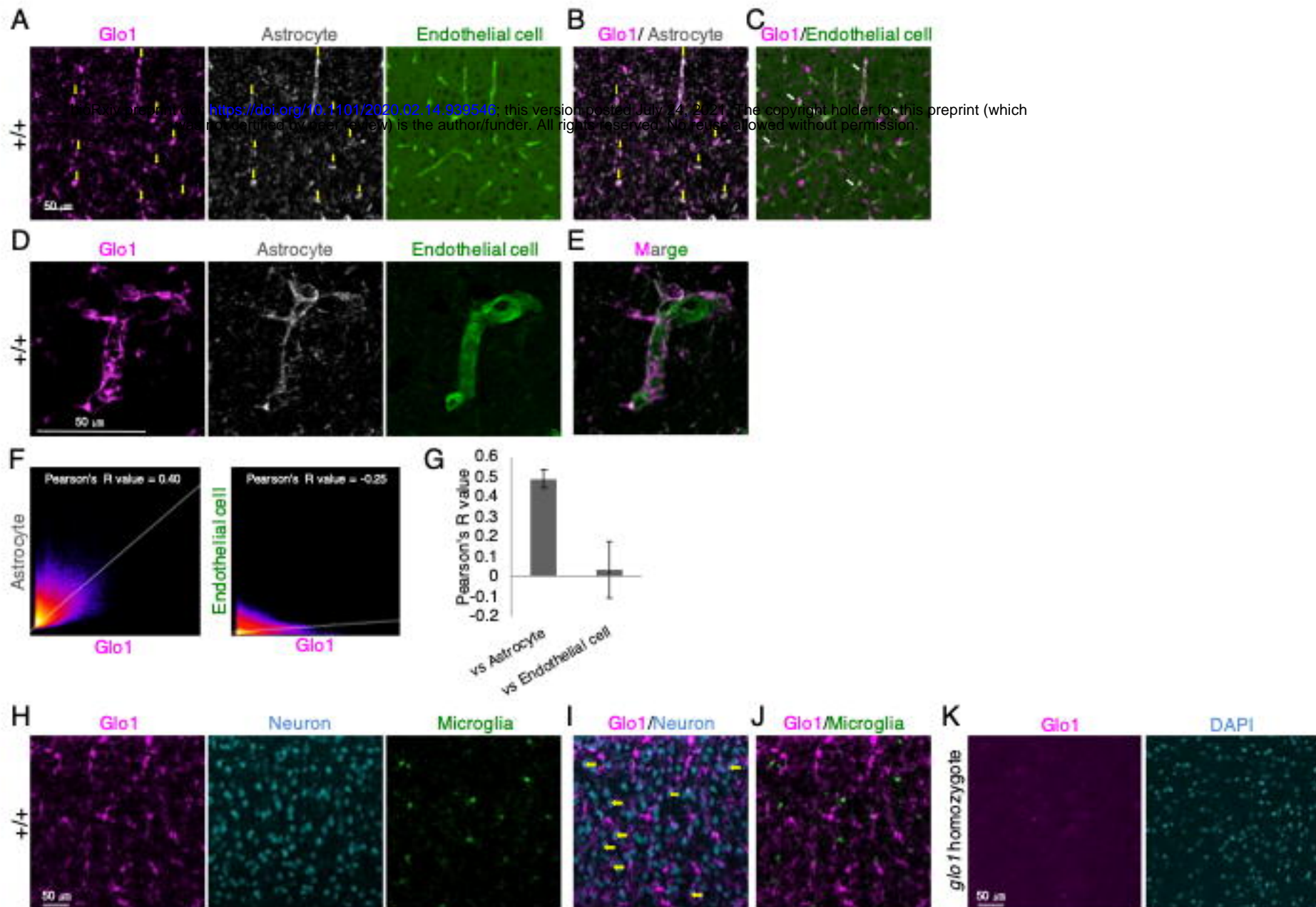
Supplementary Figure 2 Analyses of psychiatric disease-related phenotypes in *Glo1* heterozygotic mice.

(A, B) Behavioral analyses in the four groups of mice (n = 18–23 mice per group). (A) Elevated plus maze test to evaluate anxiety. (B) Preference for novel mouse in the three-chamber test, calculated as [(time spent exploring novel mouse)/(total time spent exploring novel mouse and novel object)] × 100%. (C) Extracellular dopamine concentration in the medial prefrontal cortex measured at 20-min intervals using an *in vivo* microdialysis system (n = 7–10 mice per group). (D–I) Effect of aripiprazole treatment on behavioral phenotypes (n = 12–18 mice per group). (D) Effects of aripiprazole treatment on the acoustic startle response, (E) pre-pulse inhibition (PPI) using a 70-dB pre-pulse, (F) object location test, (G) self-grooming, (H) elevated plus maze performance, and (I) nest-building skill. In (I), post hoc Tukey's multiple comparisons test of groups at specific time points, $^{***}p < 0.001$, $^p < 0.05$ for Starch +/+ vs. Suc *glo1*/+ Arip, $^{SS}p < 0.01$ for Starch +/+ Arip vs. Suc *glo1*/+ Arip, $^{SSS}p < 0.001$ for Starch +/+ vs. Suc *glo1*/+, $^*p < 0.05$ for Starch +/+ Arip vs. Suc *glo1*/+. All data are presented as mean ± SEM. $^{***}p < 0.001$, $^{**}p < 0.01$, $^*p < 0.05$



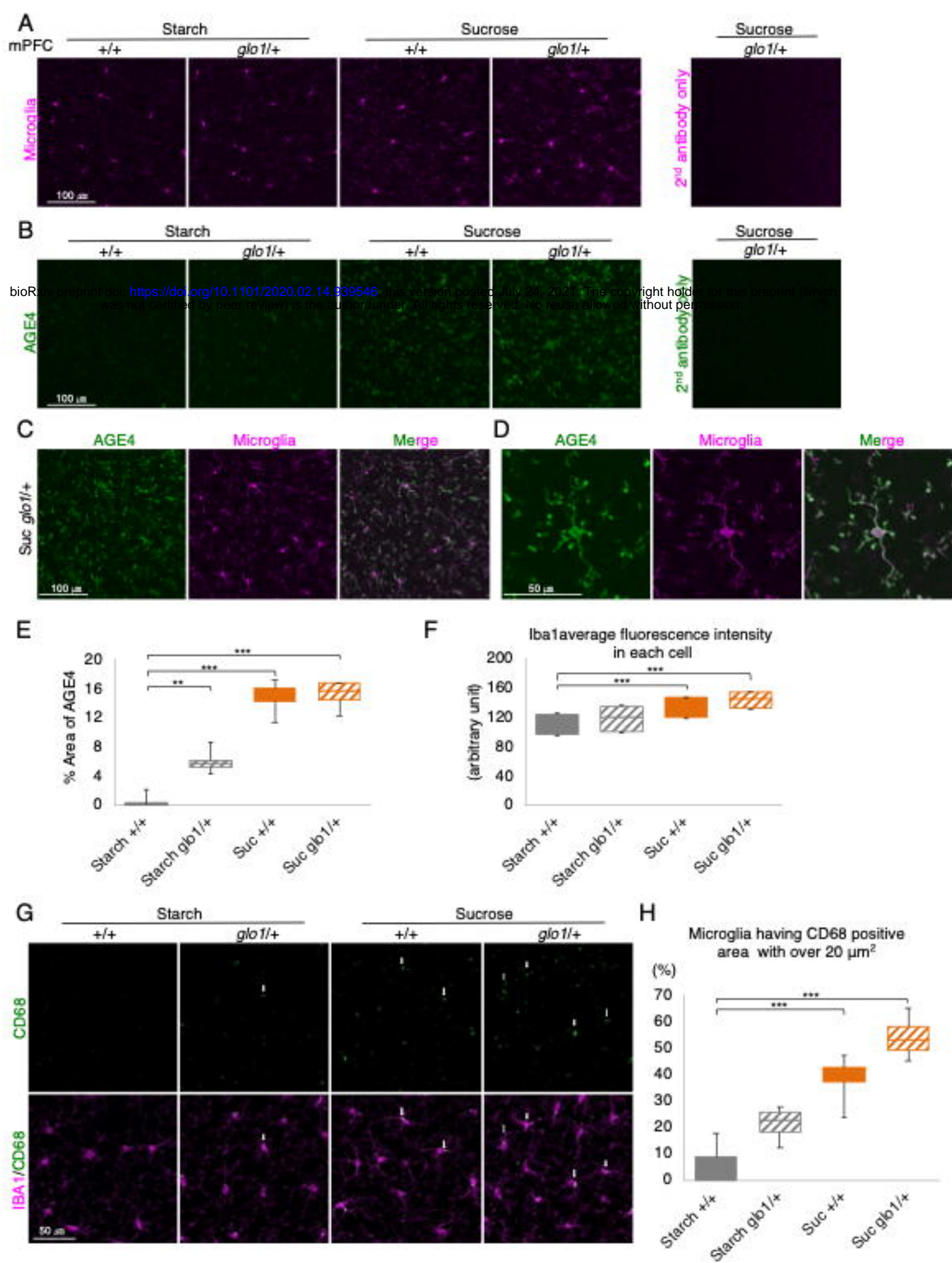
Supplementary Figure 3 Analyses of psychiatric disease-related phenotypes in *Disc1* heterozygotic mice.

(A-E) Behavioral analyses in the four groups of mice (n = 8–11 mice per group). (A) Spontaneous locomotor activity in the open-field test. (B) Acoustic startle responses. (C) Pre-pulse inhibition (PPI) using a 70 dB pre-pulse. (D) Object location test. (E) Preference for novel mouse in the three-chamber test, calculated as Fig. S2B. All data are presented as mean ± SEM. ***p < 0.001



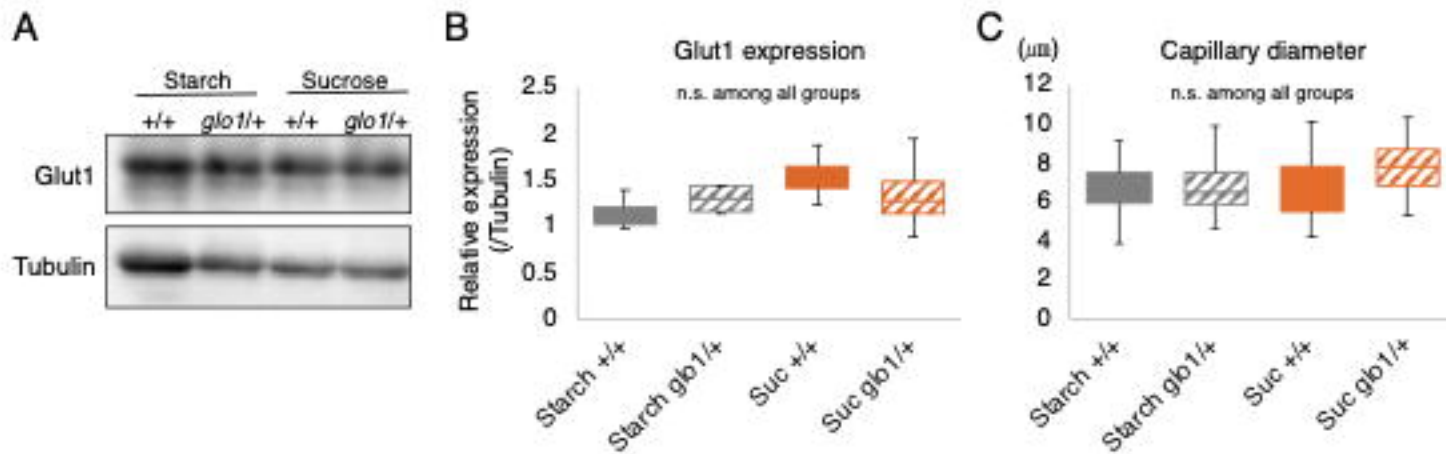
Supplementary Figure 4 GLO1 cellular localization in the prefrontal cortex of sucrose-fed wild-type mice.

(A–F) GLO1 localization in astrocytes. (A) GLO1 co-immunostaining with an astrocyte marker (ALDH1L1) or an endothelial cell marker (tomato lectin). (B) Merged GLO1/ALDH1L1 image from (A). Yellow arrows in (B) indicate cells with GLO1/ALDH1L1 colocalization. (C) Merged GLO1/lectin image from (A). White arrows denote representative GLO1-positive cells close to endothelial cells. (D, E) High magnification images of GLO1 co-immunostaining with ALDH1L1 or with tomato lectin. (E) Merged GLO1/ALDH1L1 image from (D). (F) 2D intensity histogram of the two indicated channels to identify GLO1 colocalization with ALDH1L1 or tomato lectin, interpreting the strength of a relationship based on R as follows: $R \geq 0.7$, strong; $0.7 > R \geq 0.4$, moderate; $0.4 > R \geq 0.2$, weak; and $R < 0.2$, none or very weak. (G) Average R-value of colocalization data including (F) from three different locations of three mice. (H) GLO1 co-immunostaining with the neuronal marker NeuN and the microglial marker IBA1. (I) Merged GLO1/NeuN image from (H). Yellow arrows indicate neurons with mild GLO1 immunoreactivity. (J) Merged GLO1/IBA1 image from (H). (K) GLO1 immunostaining and DAPI staining in *Glo1* homozygous mice.



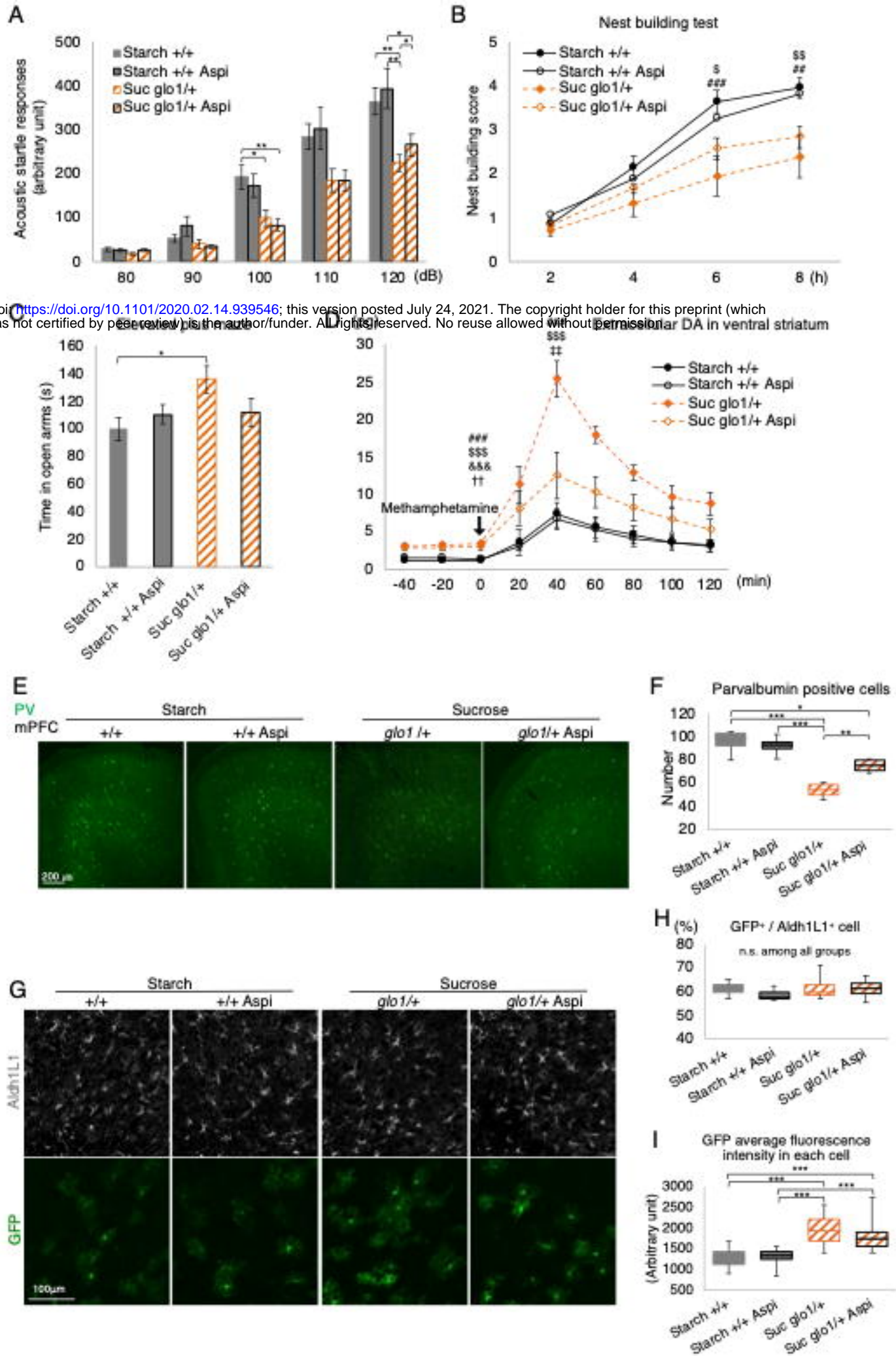
Supplementary Figure 5 Fructose-derived AGE4 accumulation in microglia of sucrose-fed mice.

(A, B) Immunohistochemical images of the microglial marker IBA1 and AGE4 in the medial prefrontal cortex region (mPFC). (C) Merged AGE4/IBA1 image. (D) High magnification images of AGE4 co-immunostaining with IBA1. (E) Measurement of the area of AGE4 immunopositivity in (B). The mean AGE4-immunopositive area intensity of the entire image was measured in each section ($n = 4$ mice per group). (F) Mean IBA1 fluorescence intensities of five randomly selected cells per image in four independent mice. (G) Immunohistological images of IBA1- and CD68-positive microglia in the mPFC region. White arrows in (G) indicate microglial cells containing CD68-positive areas $>20 \mu\text{m}^2$. (H) Percentage of CD68-positive cells per total microglial cells in each image in (G). All data are presented as mean \pm SEM. *** $p < 0.001$, ** $p < 0.01$



Supplementary Figure 6 Blood vessels Characterization in the four groups.

(A) Western blot analysis of glucose transporter 1 (Glut1) protein expression using tubulin as internal control. The cerebral cortex, including the hippocampus, was used as loading sample. (B) Densitometric analysis of Glut1 protein expression (n = 3 mice per group). To quantify expression, Glut1 band intensities in (A) were divided by the intensities of the corresponding tubulin bands. (C) Measurement of capillary diameter. Vessels <10 μ m in diameter were defined as capillaries, and the mean diameter of 10 randomly selected capillaries per image measured in Fig. 4C (n = 3 mice per group). All data are presented as mean \pm SEM.

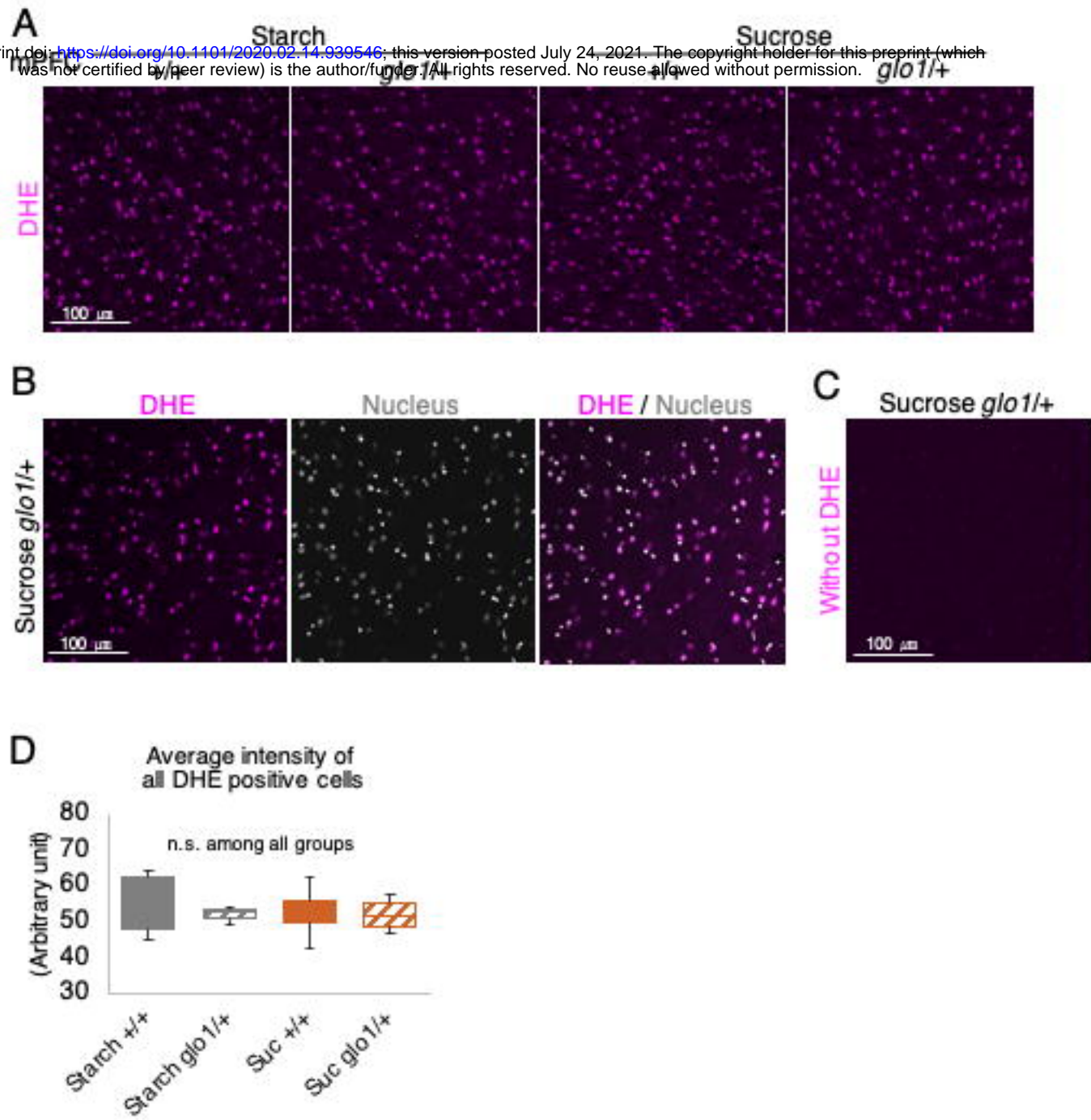


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Supplementary Figure 7 Protective effects of aspirin against the development of psychiatric phenotypes in G × E mice.

(A–C) Effects of aspirin treatment on the acoustic startle response (A), nest-building skill (B), elevated plus maze test performance (C) and dopamine concentration (D). (E) Parvalbumin (PV) Immunohistochemistry in the medial prefrontal cortex (mPFC). (F) Number of PV-positive cells in the mPFC ($n = 4$ mice per group). (G) Immunohistological images of GFP-positive astrocytes and ALDH1L1 in the mPFC region. (H) Percentage of GFP-positive cells per total ALDH1L1-positive cells in each image in (G). (I) Mean fluorescence GFP intensities of 10 randomly selected cells per image in (G) (from four independent mice). In (B), post hoc Tukey's multiple comparisons test of groups at specific time points, $^{***}p < 0.001$, $^{**}p < 0.01$ for Starch +/+ vs. Suc *glo1*+/+ Aspi, $^{SS}p < 0.01$, $^{S}p < 0.05$ for Starch +/+ Aspi vs. Suc *glo1*+/+ Aspi. In (D), post hoc Bonferroni's multiple comparisons test of groups at specific time points (0 min and 40 min), $^{***}p < 0.001$ for Suc *glo1*+/+ vs. Starch +/+, $^{SSS}p < 0.001$ for Suc *glo1*+/+ vs. Starch +/+ Aspi, $^{&&&p} < 0.001$ for Suc *glo1*+/+ Aspi vs. Starch +/+, $^{††}p < 0.01$ for Suc *glo1*+/+ Aspi vs. Suc *glo1*+/+, $^{††}p < 0.01$ for Suc *glo1*+/+ Aspi vs. Starch +/+ Aspi. All data are presented as mean \pm SEM. $^{***}p < 0.001$, $^{**}p < 0.01$, $^{*}p < 0.05$

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Supplementary Figure 8 ROS detection in the four groups.

(A) Reactive oxygen species (ROS) detection with *dihydroethidium* (DHE). (B) Merged DHE/Nucleus (TO-PRO 3) image. All cells are DHE positive, according to nucleus images. (C) ROS detection images without DHE. (D) Fluorescent intensity of DHE positive cells in (A). The mean DHE fluorescent intensity of each cell was measured in each section (n = 4 mice per group). All data are presented as mean \pm SEM.