Pharmacological And Genetic Reversal Of Age-Dependent Cognitive Deficits Attributable To Decreased Presenilin Function

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Alzheimer’s disease (AD) is the leading cause of cognitive loss and neurodegeneration in the developed world. Although its genetic and environmental causes are not generally known, familial forms of the disease (FAD) are attributable to mutations in a single copy of the Presenilin (PS) and amyloid precursor protein genes. The dominant inheritance pattern of FAD indicates that it may be attributable to gain or change of function mutations. Studies of FAD-linked forms of presenilin (psn) in model organisms, however, indicate that they are loss of function, leading to the possibility that a reduction in PS activity might contribute to FAD and that proper psn levels are important for maintaining normal cognition throughout life. To explore this issue further, we have tested the effect of reducing psn activity during aging in Drosophila melanogaster males. We have found that flies in which the dosage of psn function is reduced by 50% display age-onset impairments in learning and memory. Treatment with metabotropic glutamate receptor (mGluR) antagonists or lithium during the aging process prevented the onset of these deficits, and treatment of aged flies reversed the age-dependent deficits.

Introduction

Alzheimer’s disease (AD) is a neurodegenerative disease characterized by progressive impairments in memory and cognitive abilities with a typical late age onset, although in cases of early onset familial Alzheimer’s disease (FAD), the onset can be as early as the third decade of life. The histopathological hallmarks of AD are amyloid plaques and neurofibrillary tangles in the brains of afflicted patients (Alzheimer, 1907; Selkoe, 2002; Mattson, 2004; Hardy, 2006; Small and Gandy, 2006). The majority of FAD cases are linked to mutations in the Presenilin 1 and Presenilin 2 (PS1/PS2) genes, with additional cases linked to mutations in the amyloid precursor protein (APP) gene. FAD cases exhibit dominant inheritance pattern in which the disease is caused by a mutation in a single copy of one of these three genes (Mattson, 2004; Shen and Kelleher, 2007).

In humans as well as in animal and cell culture models of AD, FAD-linked PS and APP mutations generally, although not in all cases, result in an enhanced ratio of β-amyloid 1-42 (Aβ1–42) to Aβ1–40 (Moehlmann et al., 2002; Schroeter et al., 2003; Qitakahara et al., 2005; Walker et al., 2005; Kumar-Singh et al., 2006; Sambamurti et al., 2006; De Strooper, 2007; Isoo et al., 2007; Shen and Kelleher, 2007; Shiio et al., 2007). This finding and the dominant inheritance pattern of FAD has led to a model suggesting that FAD-linked mutations in PS1, PS2, and APP lead...
to gain-of-function or change-of-function forms of PS or APP protein (Hardy, 2006; Small and Gandy, 2006).

Studies in mice, *Drosophila*, and *Caenorhabditis elegans* indicate that FAD-linked presenilin mutations have reduced function with respect to cleavage substrates (De Strooper, 2007; Shen and Kelleher, 2007). Both loss-of-function mutations in presenilin (psn) and FAD-linked mutations lead to an increase in GSK-3β activity, which is normally negatively regulated by presenilin and may be a γ-secretase-independent function (Baki et al., 2004; Serban et al., 2005; Shioi et al., 2007). Finally, the high number of different mutations in presenilin that give rise to FAD is more consistent with a loss-of-function mechanism as gain-of-function or change-of-function mutations for a protein are normally considered to be rare, particularly because the mutations in *presenilin 1* are spread throughout different portions of the protein (Saura et al., 2004; De Strooper, 2007; Shen and Kelleher, 2007). Together, these results suggest the possibility that a reduction in the functional activity of PS1 and PS2 contributes to the pathogenesis of FAD, including cognitive loss.

We investigated the effect of reducing psn function on cognition during aging as an approach to determine whether lowered psn function (but not completely absent function) may contribute to cognitive impairment. We find that reduction in psn function leads to age-onset cognitive deficits. Through pharmacological and genetic studies, we identify misregulation of metabotropic glutamate receptor (mGluR) signaling and inositol trisphosphate receptor (InsP₃R)-mediated calcium release, as causal for the observed age-onset cognitive dysfunction. The implications of these studies toward understanding and possibly treating certain aspects of AD are discussed.

**Materials and Methods**

*Drosophila* strains and drug testing. *Drosophila* strains used in this study are described in supplemental Table 1 (available at www.jneurosci.org as supplemental material). The *Drosophila* strains were cultured as in the study by McBride et al. (2005). Drugs were obtained from Tocris Cookson and solubilized according to the instructions of the manufacturer. They were added to the fly food after cooling to the appropriate concentration. Concentrations used were based on dose–response testing performed in a previous study on the *Drosophila* fragile X mental retardation gene dfmr1 mutants (McBride et al., 2005). Vehicle for each drug was added to the appropriate control food for each experiment.

Behavioral training and testing. Virgin male flies were collected under ether anesthesia within 4 h of eclosion. Virgin XX, 1f females were collected on the day of eclosion and kept in food vials in groups of 10–15 flies. Flies were aged in a 12 h light/dark cycle before behavioral training and testing. All testing was performed during the relative light phase. Mated females were 5 d old and observed to mate with a male the night before training. The virgin females that were used as targets were 4 d old. All male subjects were transferred to fresh control food the day before testing. Male flies were assigned to random groups for behavior training and testing, which was performed blind (Siegel and Hall, 1979; Kane et al., 1997; McBride et al., 1999). The total amount of time a male was engaged in courtship activity while paired with an unanesthetized target female during a test period of 10 min or until successful copulation occurred was scored. A courtship index (CI) was calculated as the percentage of total observation time spent courting (Siegel and Hall, 1979). Binning of naive courtship behavior was performed as in the study by McBride et al. (2005), except that the percentage of flies advancing to a particular stage of courtship during the courtship interval was scored and compiled. Locomotor testing was done as in the study by McBride et al. (2005). Cell death and mushroom body morphological analysis. Terminal deoxynucleotidyl transferase-mediated biotinylated UTP nick end labeling (TUNEL) assays were performed according to Ye and Fortini (1999) using a S7110 kit from Oncor. Brains were treated following the protocol for imaginal discs. Acridine Orange stainings were performed according to Hérichè et al. (2003). Analysis of stained brains was performed by three-dimensional reconstruction of optic stacks taken at 0.5 µm using a Leica scanning confocal microscope. The number of cell death foci and their relative position were tabulated for each genotype. Analysis of mushroom body (MB) morphology was performed as described by McBride et al. (2005).

**Results**

Examination of the effect of decreased presenilin activity during aging

*Drosophila* has a single *psn* gene ortholog, the protein sequence of which is ~50% identical to human PS1 and PS2 (Boulianne et al., 1997; Hong and Koo, 1997). The encoded presenilin protein is expressed in a wide range of tissues, including the brain (Ye and Fortini, 1998). Isolation and characterization of several loss–of-function mutants have revealed that *psn* is an essential gene, required for proper neurogenesis and Notch processing (Struhl and Greenwald, 1999; Ye et al., 1999). Biochemical studies of the *Drosophila* presenilin protein have shown that it is a component of the γ-secretase complex like its mammalian counterparts (De Strooper, 2003; Hu and Fortini, 2003; Takasugi et al., 2003).

The underlying nature of the mutations in PS1 and PS2 that give rise to FAD are unclear (Saura et al., 2004; Qi-Takahara et al., 2005; Walker et al., 2005; Kumar-Singh et al., 2006; Sambamurti et al., 2006; De Strooper, 2007; Hardy, 2007; Isoo et al., 2007; Shen and Kelleher, 2007; Wolfe, 2007). However, studies in model organisms indicate that the FAD-linked mutations may result in a reduction of PS1 and PS2 activity (De Strooper, 2007; Shen and Kelleher, 2007). This possibility suggests that some phenotypes associated with Alzheimer’s disease, including age-onset cognitive loss, may be attributable to a reduction in overall PS activity levels. To investigate this possibility further, we tested the effect of reducing psn activity using known *psn* loss–of-function alleles. We evaluated the learning and memory capabilities of young adult flies heterozygous for a given *psn* mutant allele (hereafter referred to as “*psn*-het flies”), with only ~50% of the *psn* found in wild-type (WT) control flies [the wild-type control strain is Oregon-R (OR-R), the background strain from which the *psn* alleles we used in this study were derived]). We then reevaluated their capabilities at an older age to determine whether any age-related cognitive decline could be observed. We took advantage of the large number of *psn* alleles with diminished function that have been obtained from several genetic screens. In total, 10 different loss–of-function alleles of *psn* were examined, to ensure that any observed effects were not attributable to second site mutations or other background effects and crossed each to our WT OreR line (for a description of the alleles used, see Materials and Methods and Supplemental Table 1, available at www.jneurosci.org as supplemental material).

**Examining naive courtship, learning, and memory in aged control flies**

Before embarking on an examination of the effects of aging on *psn*-het flies, using the courtship based assays, we first needed to establish a timeline in which courtship–based learning and memory were not affected in our control OreR (WT) flies. The effect of aging on naive courtship and courtship conditioned learning and memory has not been extensively characterized. The lifespan of *Drosophila* is ~60 d (after eclosion, in adult form); therefore, a
The 5-d-old adult fly is relatively "young," a 30-d-old fly is "moderately aged," and a 45-d-old fly is relatively "old."

Courting *Drosophila* males perform a characteristic sequence of behaviors when paired with a female: orienting toward and following the female, tapping her with his forelegs, vibrating one wing, licking her genitalia, and attempting to copulate (Hall, 1994). The percentage of time the male spends performing any of these behaviors toward a target during a defined period of time is referred to as the CI (Siegel and Hall, 1979). We tested progressively older naive males and found that 45-d-old WT flies perform naive courtship as robustly as young 5-d-old flies (Fig. 1A). This indicates that there are no sensory impairments with age that diminish courtship. Furthermore, the quality of this courtship was not different between 5- and 45-d-old WT flies (supplemental Fig. S1, available at www.jneurosci.org as supplemental material), nor was the level of locomotor activity (supplemental Fig. S2, available at www.jneurosci.org as supplemental material).

We next determined an age range within which learning and memory would be intact in WT flies. Learning and memory can be examined in *Drosophila* by assaying conditioned courtship behavior. In conditioned courtship, a male fly learns to modify his courtship behavior after experience with an unreceptive female (Siegel and Hall, 1979; Hall, 1994). Virgin females generally respond to a courting male by mating. However, recently mated females are unreceptive and have an overlapping but altered pheromonal profile that naive males find less provocative than that of virgin female targets (Ejima et al., 2007). A naive male paired with a mated female will initially court her, but his courtship activity soon decreases. When this learning during training (LDT) is quantified, by comparing the CI during the first 10 min with a mated female to the CI of the last 10 min period of a 1 h pairing, wild-type flies typically show a \( \frac{1}{3} \) decrease in courtship activity (Joiner and Griffith, 1997; Kane et al., 1997). Hence, LDT is a form of behavioral plasticity but is distinct and separate from courtship suppression assayed after training, which is a form of associative memory (Tompkins et al., 1983; Ackerman and Siegel, 1986). When a male is paired with a virgin female after 1 h of experience with a mated female, his courtship remains depressed for 2–3 h (Siegel and Hall, 1979). This effect is not a general suppression of all courtship activity, because trained males do not modify their courtship of other pheromonally distinct targets (Ejima et al., 2005; Siwicki et al., 2005). After training with a mated female, memory is measured as a decrease in CI toward virgin females in trained males relative to naive controls.

In *Drosophila*, five phases of memory have been elucidated by a combination of genetic and pharmacological dissection. There is an immediate recall memory (immediate memory) at 0–2 min after training, short-term memory out to 1 h, medium-term...
memory out to 6 h, anesthesia-resistant memory out to 2 d, and long-term memory lasting up to 9 d after training that appears to be dependent on protein synthesis (Skoulakis and Grammenoudi, 2006). Intact short-term memory is dependent on intact immediate recall. However, immediate recall and short-term memory are distinct from LDT. Therefore, intact memory can occur without LDT, and learning during training can occur without posttraining memory (Joiner and Griffith, 1997; Kane et al., 1997; McBride et al., 2005).

To assess LDT, a male fly was placed in a training chamber with a previously mated female for 1 h, and the amount of time the male spent courting in the initial 10 min interval was compared with the time spent engaged in courtship in the final 10 min interval. Normal LDT in 5-d-old WT flies is illustrated by a significant ≥40% decrease in courtship during the training (Fig. 1B), and short-term memory is illustrated in 5-d-old WT flies by a significant decrease in courtship toward virgin females 1 h after training (Fig. 1D). We found that 45-d-old WT flies also demonstrated intact LDT and short-term memory (Fig. 1C,E). These studies indicate that the courtship-based learning and memory paradigm can be used to study aging-related issues of cognition and behavior in flies.

Naive courtship, LDT, and memory in young and aged adult psn-het flies

Courtship was robust in naive males of all 10 psn-het genotypes examined as young adults at 5 d of age (Fig. 1F). Naive males of these psn-het genotypes also displayed the normal steps of courtship behavior and normal levels of locomotor activity (supplemental Figs. S3, S4, available at www.jneurosci.org as supplemental material). LDT at 5 d of age in all 10 psn-het genotypes was normal, as evidenced by decreases of ≥40% from initial CI to final CI during training with mated females (Fig. 1G,H). To examine the immediate recall memory at 5 d of age, we took males that had just completed a 1 h training session with a previously mated female and immediately placed them in a new chamber with a virgin target female for a 10 min courtship assay. This CI was then compared with the courtship level of naive males that had been placed in a training chamber for 1 h with no female, before being introduced to a virgin target female. Immediate recall was intact in all 10 psn-het genotypes examined (Fig. 1I,J). To assay short-term memory, the trained male was placed in a holding chamber for 60 min and then subsequently placed in a testing chamber with a virgin female target. None of the psn-het genotypes were tested and found to have normal short-term memory at 5 d of age (Fig. 1K,L).

We next examined naive courtship and LDT in 30-d-old (late middle age) psn-het males. No significant effect on the level of naive courtship (Fig. 2A), its quality (supplemental Fig. S5, available at www.jneurosci.org as supplemental material), locomotor activity (supplemental Fig. S6, available at www.jneurosci.org as supplemental material), or phototaxis or chemotaxis (data not shown) was observed. However, males of all 10 different psn-het genotypes failed to demonstrate the typical decrease in courtship activity during the training session or a statistically significant reduction in courtship behavior. Therefore, psn-het flies have an age-dependent impairment in learning during training. D, E, Immediate recall remained intact in all psn-het males. F, G, psn-hets were also tested for short-term memory at 60 min after training. All lines failed to exhibit short-term memory, in contrast to their behavior at 5 d of age (Fig. 1K, L).
sults demonstrate an age-dependent impairment in LDT in psn-het male flies by 30 d of age.

We next examined immediate recall and short-term memory in the psn-het flies at 30 d of age. We found that males of all 10 different psn-het genotypes displayed intact immediate recall at 30 d of age (Fig. 2D,E). Intact immediate recall of courtship memory without intact LDT has been observed previously in young adult Drosophila (Joiner and Griffith, 1997; Kane et al., 1997). Although immediate recall was intact, examination of short-term memory in nine of the heterozygous mutant presenilin lines, at 30 d of age, revealed that this form of memory was no longer intact (Fig. 2F,G). This is in contrast to 5-d-old psn-het and 45-d-old WT flies, which exhibited intact short-term memory. This demonstrates that there is an age-dependent impairment of short-term memory in heterozygous psn loss-of-function flies.

Genetic interaction between psn and dfmr1
In an effort to identify novel pathways affected as a result of the reduction in psn levels, we looked for genetic interactions between psn and other genes known to affect cognition and behavior that might play a role in presenilin signaling based on the literature. Interestingly, we identified a strong genetic interaction between psn and the Drosophila fragile X mental retardation gene dfmr1. At 5 d of age, psn and dfmr1 heterozygous single-mutant flies display normal naive courtship levels relative to their given genetic background (supplemental Fig. S7, available at www.jneurosci.org as supplemental material). However, 5-d-old flies that are transheterozygous mutant for both psn and dfmr1 display a severe reduction in naive courtship levels, which is rescued by the introduction of one wild-type copy of a dfmr1 genomic rescue construct. Such heterozygous synergistic genetic interactions are rare and suggest that both genes act in one or more common pathways important for normal naive courtship activity.

Prevention of age-dependent LDT and short-term memory deficits with pharmacological treatment
To explore the possibility of pharmacologically rescuing some of the age-dependent psn-het phenotypes and in light of the genetic interaction between psn and dfmr1 described above, we focused on treatments that rescue naive courtship and cognitive deficits in dfmr1 mutants. One pathway that is known to be affected in both fly and mouse models of fragile X is mGluR signaling (Huber et al., 2002; McBride et al., 2005; Yan et al., 2005; Dönlen et al., 2007; Bolduc et al., 2008). In previous studies, we found that several of the phenotypes displayed by the dfmr1 mutants, including naive courtship and memory, could be rescued by treatment with antagonists of DmGluRA and LiCl (McBride et al., 2005). More recent studies have also observed this and have also found that similar rescue is obtained by genetic reduction of DmGluRA activity, the sole Drosophila mGluR (Bolduc et al., 2008; Pan et al., 2008).

Our working model is that treatment of flies with both lithium and antagonists of DmGluRA will increase CAMP signaling and may also decrease InsP3R-mediated calcium release (Fig. 3A). Lithium inhibits GSK-3β activity by competing with the magnesium-binding site and increasing phosphorylation at serine 9 (Gould and Manji, 2005; Huang and Klein, 2006; Jope and Roh, 2006). There is an antagonistic relationship between protein kinase A (PKA) and GSK-3β; therefore, this acts to upregulate the downstream effect of CAMP signaling because GSK-3β can inhibit CAMP response element-binding protein (CREB)-mediated gene transcription (Bullock and Habener, 1998; Grimes and Jope, 2001; Mai et al., 2002; Tanji et al., 2002; Hansen et al., 2004; Gould and Manji, 2005). Lithium also inhibits inositol polyphosphate 1-phosphatase (IPPPase) (Acharya et al., 1998) and inositol monophosphatase (IMPase) (Berridge, 1993), thereby reducing the pool of InsP3 available to stimulate InsP3R-mediated calcium signaling (Berridge, 1993; Takei et al., 1998; Williams et al., 2002). Previous heterologous cell-based studies have shown that DmGluRA can couple to heterotrimeric Gα leading to decreases in cAMP levels (Parmentier et al., 1996). More recent in vivo studies have shown that DmGluRA also has properties typically found associated with Gα-coupled group I mGluRs such as regulating synaptic morphology and AMPA receptor presentation (Bogdanik et al., 2004; Pan and Broaddie, 2007; Pan et al., 2008). Thus, antagonizing DmGluRA activity will increase cAMP levels (by relieving inhibition of cAMP after synaptic stimulation) and will possibly decrease InsP3R-mediated calcium signaling (Fig. 3A) (McBride et al., 2005). We therefore explored the possibility that the age-dependent cognitive deficits could be rescued by treatment with DmGluRA antagonists as well as lithium.

For this experiment, we used the competitive group II antagonist LY341495 [(2S)-2-amino-2-[(15,2S)-2-carboxycycloprop-1-yl]-3-(xanth-9-yl) propanoic acid] and noncompetitive mGluR5 antagonist 2-methyl-6-(phenylethynyl)-pyridine (MPEP), as well as lithium at concentrations found previously to be optimal for treating dfmr1 mutants (McBride et al., 2005). LY341495 is a very specific group II antagonist that has been shown to antagonize DmGluRA in the nanomolar range (Bogdanik et al., 2004). Although MPEP was developed as a noncompetitive antagonist of the vertebrate group I mGluRs, we have included it in our studies because the MPEP binding pocket of mGluR5 is more highly conserved with DmGluRA than any other vertebrate mGluRs and in previous experiments has similar activity to the known DmGluRA antagonists LY341495, MPPG [(R)-α-methylserine-O-phosphate], and MTGP [(R)-α-methyl-4-tetrazolylphenylglycine] and shows similar effects to that obtained by genetically reducing DmGluRA activity (Parmentier et al., 1996; Pagano et al., 2000; Malherbe et al., 2003; Bogdanik et al., 2004; McBride et al., 2005; Bolduc et al., 2008; Pan et al., 2008). However, because this has not been formally shown to be an antagonist of DmGluRA, it is therefore a putative DmGluRA antagonist.

Two different psn-het genotypes (psn B3 and psn I2) were treated with different drugs or non-drug-containing control vehicle solutions from days 6 to 29 of adulthood (after eclosion). Flies were then transferred to control food containing no drug for 1 d before being tested. Flies were treated with 5 mM NaCl (ionic control for LiCl), 0.004% DMSO (the vehicle for LY341495), 5 mM LiCl (lithium), 8.6 μM MPEP, or 400 nM LY341495. Remarkably, treatment with LiCl, LY341495, or MPEP improved LDT performance in psn-het flies (Fig. 3H–J). In contrast, no improvement was observed in flies treated with NaCl or DMSO (Fig. 3K,L). These experiments demonstrate that the impairment in LDT exhibited by the untreated psn-het flies can be prevented by treatment with lithium, LY341495, and MPEP.

This result with LDT raised the possibility that treatment with lithium, LY341495, and MPEP might also prevent the age-dependent impairment in short-term memory that is exhibited by the psn-het males. Using the same set of pharmacologic treatments, we found that psn-het flies treated from 6 to 29 d after eclosion with 5 mM lithium, 8.6 μM MPEP, or 400 nM LY341495 demonstrated intact short-term memory (Fig. 3M–O), whereas those treated with 5 mM NaCl or 0.004% DMSO did not (Fig. 3P,Q). This result demonstrates that the impairment in short-
term memory exhibited by 30-d-old psn-het flies can be prevented by treatments aimed at reducing DmGluRA activity. In contrast, prolonged treatment of OreR (WT) flies with lithium, LY341495, and MPEP impaired LTD and short-term memory, whereas vehicle treatment did not (Fig. 3B–G).

The rescue that we observe when psn-hets are treated with LY341495, MPEP, or lithium is analogous to the results, we obtained when we treated dfmr1 mutants with these same drugs (McBride et al., 2005). The similarity of these results suggests that both psn-hets and dfmr1 mutants display cognitive deficits that are attributable to enhanced DmGluRA signaling. However, another possibility is that these drug treatments represent some type of panacea that can rescue any cognitive deficit observed in flies. With this second possibility in mind, we tested the effect of treating another disease model that displays age-onset loss of short-term memory with LY341495, MPEP, and lithium.

We have found that expression of human wild-type α-synuclein in the mushroom bodies of the fly leads to age-dependent loss of short-term memory. The mushroom bodies are required for short-term memory formation in the conditioned courtship paradigm (Joiner and Griffith, 1999; McBride et al., 1999). In controls for these flies, short-term memory is intact at 30 d of age (supplemental Fig. S8A, available at www.jneurosci.org as supplemental material). In flies expressing human wild-type α-synuclein in the mushroom bodies, short-term memory is intact at 5 d of age but is impaired at 30 d of age (supplemental Fig. S8B, available at www.jneurosci.org as supplemental material). Treatment of these flies with lithium, LY341495, or MPEP failed to rescue this deficit (supplemental Fig. S8C,D, available at www.jneurosci.org as supplemental material), indicating that rescue of cognitive deficits in the psn-hets by the drug treatments is disease specific and may not necessarily be extrapolated to other disease models displaying age-dependent cognitive impairments.

Morphological analysis of heterozygous psn brain tissues

The age-dependent impairments in LTD and short-term memory observed in psn-het flies could possibly be attributable to the loss of critical neurons as a result of cell death. There is a body of literature describing loss of synapses and neurons in the brains of Alzheimer’s patients and in related mouse models (Selkoe, 2002; Mattson, 2004; Walsh and Selkoe, 2004; Hardy, 2006; Small and Gandy, 2006). To determine whether neuronal loss could account for the cognitive loss, we performed TUNEL staining on...
Figure 4. Cell death and morphology in 50-d-old psn-het flies. Pharmacologic treatments can reverse impairments in learning during training and short-term memory in psn-het flies. A–C, Analysis of cell death in aged psn-het, OreR, and Adar mutant brains. TUNEL staining was performed on 50-d-old (negative control) OreR (A), 50-d-old psn-het (B), and 5-d-old (positive control) Adar mutant brains (C). The brains were examined by confocal microscopy to identify cells undergoing cell death. Foci containing at least one cell undergoing cell death are denoted by white arrows. Maximum projections, as shown in A–C, made from confocal stacks of entire brains were examined for cell death foci. The number of foci in the central brain region for each brain was scored. The distribution of the number of cell death foci versus the percentage of brains with such number of foci is shown in D. The majority of psn-het brains lacked any detectable cell death (n = 41). Those that did had less than five foci, and there was no consistent location in the brain in which the foci were detected. Similar results were obtained with the OreR brains (n = 33). Cell death foci were detected in all Adar mutant brains (n = 5), which have been described previously to undergo massive cell loss via cell death (Palladino et al., 2000). Fifty-day-old OreR (E) and psn-het (F) brains were stained with anti-FasII and examined by confocal microscopy to examine the integrity of the mushroom bodies. This antibody strongly labels the α-lobes (arrowhead shown in E) and weakly stains the γ-lobes (white arrow with #, shown in E). There was no detectable difference in the morphology of the mushroom bodies between OreR (n = 21) and psn-het (n = 18) brains. G–P, Treatment with LY341495, MPEP, or lithium reverses the age-dependent impairments in learning during training and short-term memory observed in psn-het flies. psn-hets for the psn B3 and psn I2 alleles were raised on control food and then, at day 30 after eclosion, were placed on food containing 8.6 μM MPEP (G, I), 400 mM LY341495 (H, M), 5.0 mM lithium (J, N), 5.0 mM NaCl (L, O), or 0.004% DMSO (K, P) until day 44 and were tested on day 45. Mean ± SEM CIs are plotted, and n values are indicated above each bar for all groups. The levels of significance are indicated (∗∗p < 0.01, ***p < 0.001). A triangle indicates a 40% or greater reduction in courtship activity during the training session with a previously mated female (G–P). psn-het flies that were treated with LY341495, MPEP, or lithium demonstrated intact learning during training (G–I) and intact short-term memory (J–K) at 45 d of age. In contrast, psn-het flies that were on food containing NaCl or DMSO did not display learning during training (J, K) or short-term memory (O, P) at 45 d of age.

Reversal of the learning during training and short-term memory impairments Because the brain tissue of the aged psn-het flies appears to be morphologically normal, we considered the possibility that the age-related behavioral phenotypes could be reversed after they have already become established. Because the age-dependent impairments in LDT and short-term memory could be prevented by treatment with LY341495, MPEP, or lithium, we next examined whether these phenotypes could be reversed by these treatments. Reversal of the deficits would further indicate that the age-dependent cognitive impairments were likely the result of physiological neuronal defects (disrupted intracellular signaling possibly followed by synapse loss) and not cell death. Using the same two genotypes (psn-hets with decreased presenilin activity) that were subjected to treatments as young adults (Fig. 3), we now began treatments at day 30 and treated until day 44, testing at 45 d of age. As above, flies were treated with 5 mM NaCl, 0.004% DMSO, 5 mM lithium, 8.6 μM MPEP, or 400 mM LY341495. Treatment with LY341495 and the putative DmGluRA antagonist MPEP, as well as lithium, reversed the LDT (Fig. 4G–I) and short-term memory (Fig. 4L–N) phenotypes in both lines tested. In contrast, treatment with 5 mM NaCl and the DMSO vehicle from days 30 to 44 failed to restore normal LDT (Fig. 4F, K) or short-term memory (Fig. 4O, P) in either of the two lines. This demonstrates that the impairments in LDT and short-term memory exhibited by untreated psn-het flies can be reversed by treatments aimed at reducing DmGluRA activity. This finding is significant considering that our results may be relevant to the pathogenesis of Alzheimer’s disease in humans, which is only
recognized after the onset of cognitive impairments, emphasizing the need to develop therapeutic strategies that can be implemented after the appearance of overt disease symptoms.

**Genetic reduction of DmGluRA prevents the onset of LDT and short-term memory defects in aged psn-hets**

To validate the pharmacological studies described above, we tested the effect of genetically reducing the level of DmGluRA, the only metabotropic glutamate receptor in the *Drosophila* genome (Bogdanik et al., 2004), in the psn-het background. Given the reported binding specificities of MPEP, the ability of MPEP treatment to phenocopy genetic loss of function in DmGluRA mutants, the reported actions of lithium, and the fact that LY341495 has been shown to be a very effective antagonist of DmGluRA, our drug treatments should reduce DmGluRA signaling activity (Bogdanik et al., 2004; McBride et al., 2005). If our observed rescue is occurring through reduction of DmGluRA signaling activity, our expectation is that genetic reduction of DmGluRA should similarly provide some level of rescue of the age-onset cognitive deficits. To perform this test, we used the previously described null allele of DmGluRA (*DmGluRA*122) and precise excision wild-type allele (*DmGluRA*2b) (Bogdanik et al., 2004).

In testing DmGluRA heterozygous flies alone, we found that, at 30 d of age, they display a defect in LDT as well as in short-term memory (Fig. 5A, D), which is similar to what we observed when OreR control flies were treated with LY341495, MPEP, and LiCl (Fig. 3B–G). In contrast, the wild-type controls for these lines displayed intact LDT and short-term memory at 30 d of age (Fig. 5A, D). Consistent with our hypothesis, we find that reducing the gene dosage of the DmGluRA receptor by 50% in the psn-het background prevents the age-onset loss of LDT and short-term memory observed previously at 30 d of age, whereas introduction of the precise excision allele has no effect (Fig. 5B, C, E, F). These results validate the pharmacological studies described above and again indicate that enhanced DmGluRA signaling contributes to the age-onset cognitive deficits. The results obtained from the DmGluRA heterozygous flies and OreR WT flies treated with MPEP, LY341495, and lithium, as described above, all indicate that LDT and short-term memory are sensitive to a reduction in the activity of this receptor as well.

**Genetic reduction of the inositol polyphosphate 1-phosphatase prevents age-onset cognitive defects in the psn-hets**

The results of our pharmacological and genetic analyses indicate that reducing DmGluRA signaling rescues age-onset cognitive deficits in psn-hets. Activation of DmGluRA through the G$_i$-coupled pathway lowers cAMP signaling and through potential G$_q$ coupling should increase InsP$_3$R-mediated calcium signaling (Fig. 3A) (McBride et al., 2005). Lithium as well as DmGluRA antagonists should act to increase cAMP-mediated signaling by increasing PKA activity and may also act to lower InsP$_3$R-mediated calcium signaling (Fig. 3A).

Previous studies indicate that presenilin proteins bind to the inositol triphosphate receptor, and expression of mutant forms of presenilin protein lead to elevated Ca$^{2+}$ signaling (Stutzmann et al., 2004; Cheung et al., 2008). The mechanism linking mGluR activity to increased InsP$_3$R activity is shown in Figure 3A. Decreased presenilin function by FAD mutations has been shown to increase the sensitivity of the InsP$_3$R to InsP$_3$. This means that, in response to glutamate release after synaptic stimulation, the mGluR will be activated, thereby activating G$_q$ and G$_i$. The G$_q$ activation will in turn generate InsP$_3$. Because there is less wild-type presenilin present to basally inhibit the sensitivity of the InsP$_3$R to InsP$_3$, there will be enhanced InsP$_3$R-mediated calcium release in response to physiologic synaptic stimulation. To test whether the rescue we observe with genetic and pharmacological manipulation aimed at reducing DmGluRA activity is possibly working by lowering enhanced InsP$_3$R activity, we have undertaken manipulations to directly lower InsP$_3$R activity. Toward this goal, we tested the effect of genetically reducing the levels of the InsP$_3$R locus. InsP$_3$ase as well as IMPases are directly inhibited by lithium and are required for InsP$_3$ recycling and synthesis (Berridge, 1993). Genetic reduction of InsP$_3$ase has been shown to reduce InsP$_3$R signaling in flies (Acharya et al., 1998). We tested two loss-of-function alleles (InsP$_3$ase1 and InsP$_3$ase3) (Acharya et al., 1998) as heterozygotes and found that, at 30 d of age, heterozygous males for one allele (InsP$_3$ase1) lacked significant LDT, but heterozygote males for both alleles displayed normal short-term memory (Fig. 6A, D). When crossed in the psn-het backgrounds, the InsP$_3$ase1 allele prevented loss of LDT in both psn alleles and the InsP$_3$ase4 allele prevented this loss of LDT in the psn122-het background (Fig. 6B, C). Both alleles of InsP$_3$ase prevented loss of short-term memory in both psn-het lines (Fig. 6E, F). These results indicate that reduction of InsP$_3$ase activity can prevent the age-onset cognitive deficits observed with the psn-het flies. Next we directly tested whether genetic reduction of the InsP$_3$R gene itself could modulate the age-dependent psn-het phenotypes. To test the effect of genetically reducing InsP$_3$R activity, we used three
different strength alleles: a hypomorph (wc361), a molecular null (90B0), and an antimorphic allele (wc703) (Venkatesh and Hasan, 1997; Deshpande et al., 2000). At 30 d of age, heterozygous males for these alleles display weak (wc703) or no detectable (wc361 and 90B0) LDT, and all display normal short-term memory (Fig. 7A,E). When introduced into the psn-het background, two of these alleles prevented loss of LDT in the B3 allele of psn (Fig. 7B–D), and all three alleles prevent the loss of short-term memory in both psn alleles at 30 d of age (Fig. 7E–H). The 90B0 allele continues to prevent this loss of short-term memory at 40 d of age (Fig. 7I). These results suggest that a reduction in InsP₃R-mediated Ca²⁺ release is also another potential route to ameliorate the age-onset cognitive deficits associated with reduced psn levels.

Discussion

To date, all cases of early onset familial forms of Alzheimer’s disease are attributable to mutations in a single copy of PS1, PS2, or APP. The large number of different mutations in the PS1 and PS2 genes that cause AD as well as analysis of the protease function of FAD mutants is consistent with a loss of function of PS1 and PS2 as a contributing cause of FAD (De Strooper, 2007; Hardy, 2007; Shen and Kelleher, 2007). This interpretation of the data suggests that the levels or activity of PS1, PS2, and APP are crucial for maintaining normal cognition throughout life. Based on these studies, we have explored whether this dosage sensitivity of presenilin also exists in Drosophila by examining the cognitive capabilities of psn-het flies in young and progressively aged psn-het flies. Using a conditioned courtship paradigm, we have found that deficits in LDT and short-term memory develop in the psn-het flies with age. These results further strengthen the hypothesis that some aspects of FAD and possibly AD may be caused by reduced levels of psn activity.

The appearance of cognitive deficits before any detectable neuronal loss

Previous Drosophila AD models have revealed phenotypes that have parallels with those observed in mouse models and human patients. These fly models have been based on ectopic expression of either human tau or human AbB₃₄₀ or AbB₄₂ peptides and have been shown to exhibit either hyperphosphorylated tau or aggregations of Aβ that result in neurodegeneration (Wittmann et al., 2001; Iijima et al., 2004; Mershin et al., 2004; Crowther et al., 2005). Additionally, it has been demonstrated that expression of human tau, AbB₃₄₀, or AbB₄₂ leads to memory deficits (Iijima et al., 2004; Mershin et al., 2004). In these models, memory deficits are already present in young adult flies. In contrast, we have uncovered cognitive impairments that are age dependent in the psn-het flies and occur before any detectable loss of neurons. In studying the psn-het flies, we detect a clear deficit in LDT and short-term memory that appeared by 30 d of age. Our examination for cell death was done on psn-hets that were at least 50 d old, e.g., 20 d older than when the defects in LDT and short-term memory are detected. Consistent with this finding, we were able to reverse the cognitive deficits by initiating treatment at day 30, which would likely not occur if neuronal loss were the cause of the cognitive decline.

Although neuronal death is typical of AD, cognitive impairment precedes neuronal death in humans and animal models. Indeed, recent studies have indicated that synapse loss is better correlated with memory impairment than the histopathological hallmarks of plaques and tangles in AD patients (DeKosky and Scheff, 1990; Terry et al., 1991; Coleman and Yao, 2003). In fact, memory impairment in Alzheimer’s disease can occur in the absence of plaques and tangles in animal models (Oddo et al., 2003; Gong et al., 2004; Iijima et al., 2004; Mershin et al., 2004; Billings et al., 2005) or in FAD patients (Raux et al., 2000; Amtul et al., 2002; Dermaut et al., 2004; Halliday et al., 2005). Additionally, within 2–4 years of AD onset, 25–35% loss of synapses is observed in biopsies of frontal cortex in afflicted patients (Davies et al., 1987). Furthermore, synaptic function may be impaired even before synaptic loss occurs in animal models of AD as well as in patients (Oddo et al., 2003; Palop et al., 2003; Westphalen et al., 2003; Yoo et al., 2003). Thus, our results add to a growing body of literature that indicates that loss of synaptic plasticity can occur before or without plaque or tangle formation or neuronal loss. Our results also demonstrate that this loss of synaptic plasticity can occur solely as a result of a reduction in psn activity, in the absence of Aβ accumulation, which does not occur in Drosophila.

Pharmacological rescue of LDT and short-term memory

In a genetic search for pathways affected by a reduction in psn levels, we identified a strong genetic interaction between psn and dfmr1. Because this genetic interaction revealed a phenotype in courtship that was rescued previously in dfmr1 mutant flies by treatment with DmGluRA antagonists and lithium, we explored the possibility that these drugs might have efficacy in treating the
age-onset phenotypes of the psn-hets. Consistent with our findings, recent studies of mouse Alzheimer’s disease models and of afflicted humans have suggested that calcium signaling is enhanced and cAMP signaling may be decreased (Selkoe, 2002; Gong et al., 2004; Mattson, 2004; Walsh and Selkoe, 2004). We propose that these two signaling pathways are downstream of DmGluRA signaling (Fig. 3A).

In our studies, we demonstrated that treatment with lithium, the DmGluRA antagonist LY341495, and the putative DmGluRA antagonist MPEP prevented age-dependent impairments in psn-het flies at 30 d of age and can reverse LTD and short-term memory impairments if treatment is begun at 30 d of age, a time point after the deficits appear. These treatments should function to counteract decreased cAMP signaling and may counteract in an age-onset cognitive decline.

**Genetic reduction of the InsP$_3$R pathway also prevents age-onset cognitive decline**

We wanted to further elucidate the pathways downstream of DmGluRA that might be responsible for the rescue observed when the DmGluRA activity is reduced, genetically or pharmacologically. Lithium treatment and decreases in mGluR activity result in increased InsP$_3$R-mediated calcium signaling in vertebrates. In our genetic tests, we found that genetic reduction of InsP$_3$R-mediated calcium signaling provides similar rescue to that observed when DmGluRA activity is reduced genetically. By testing multiple alleles of IPPase as well as InsP$_3$R, we found that, in general, loss-of-function mutants of both genes provided rescue of age-onset deficits of LTD and short-term memory in the psn-hets.

The InsP$_3$R is an endoplasmic reticulum localized Ca$^{2+}$ channel that releases Ca$^{2+}$ from the endoplasmic reticulum into the cytoplasm in response to InsP$_3$ binding. It was demonstrated recently that FAD mutations of presenilin expressed in cells resulted in increased sensitivity to InsP$_3$-mediated Ca$^{2+}$ release, leading to enhanced cytoplasmic concentrations of Ca$^{2+}$ (Cheung et al., 2008). These results fit with previous studies in which both FAD mutations and loss of functional activity of wild-type presenilin have been linked to increases in InsP$_3$-mediated calcium signaling in cell culture and transgenic mouse models of AD (Peterson et al., 1988; Huang et al., 1991; Ito et al., 1994; Takebayashi et al., 1995; Hirashima et al., 1996; Etcheberrigaray et al., 1998; Stutzmann et al., 2004). Our results demonstrating that pharmacological treatment with drugs that should reduce DmGluRA activity, as well as genetic reduction of DmGluRA, IPPase, and the InsP$_3$R, are consistent with the hypothesis that reduction in this Ca$^{2+}$ release mechanism might ameliorate the age-onset cognitive deficits associated with AD. Moreover, these results provide the first test of the pathogenic nature of enhanced InsP$_3$R-mediated calcium signaling in AD. Previous studies have speculated on the pathogenic nature of this calcium signaling but could not rule out the possibility that these changes were compensatory in AD. Herein we were able to demonstrate a rescue of...
cognitive deficits by decreasing this signaling, indicating that up-regulation of this pathway leads to cognitive loss over time.

In conclusion, our study of Drosophila psn haploinsufficiency contributes to the understanding of age-onset cognitive loss and provides a new model to study aspects of Alzheimer’s disease. We examined several phases of learning and memory in young and older Drosophila adults and identified age-dependent cognitive impairments in learning during training and short-term memory. We have demonstrated that lithium treatment can rescue cognition in an animal model of AD. These results are in contrast to a previous attempt with lithium treatment in a mouse model of AD that failed to rescue working memory impairments (Caccamo et al., 2007). We have additionally identified and demonstrated the efficacy of mGluR antagonists as a novel therapeutic target for the treatment of the cognitive deficits associated with AD. Our results also indicate that lowering IP3ase activity or lowering InsP3, R-mediated calcium signaling can rescue cognitive loss. Finally, this study demonstrates that reversal of memory impairment after onset can be obtained by treatment with either mGluR antagonists or lithium. These results suggest novel therapeutic targets that may have relevance for treatment of AD to be explored in other animal models.

References


