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## Tyr682 in the APP intracellular domain regulates synaptic connectivity, cholinergic function and cognitive performance

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### SUMMARY

Processing of A $\beta$ -precursor protein (APP) plays an important role in Alzheimer's disease (AD) pathogenesis. The APP intracellular domain contains residues important in regulating APP function and processing, in particular the <sup>682</sup>YENPTY<sup>687</sup> motif. To dissect the functions of this sequence *in vivo*, we created an APP knock-in allele mutating Y<sup>682</sup> to Gly (*APP<sup>YG/YG</sup>* mice). This mutation alters processing of APP and TrkA signaling, and leads to postnatal lethality and neuromuscular synapse defects when expressed on an *APP-like protein 2* KO background. This evidence prompted us to characterize further the *APP<sup>YG/YG</sup>* mice. Here, we show that *APP<sup>YG/YG</sup>* mice develop aging-dependent decline in cognitive and neuromuscular functions, a progressive reduction in dendritic spines, cholinergic tone and TrkA levels in brain regions governing cognitive and motor functions. These data are consistent with our previous findings linking NGF and APP signaling and suggest a causal relationship between altered synaptic connectivity, cholinergic tone depression and TrkA signaling deficit, and cognitive and neuromuscular decline in *APP<sup>YG/YG</sup>* mice. The profound deficits caused by the Y<sup>682</sup> mutation underscore the biological importance of APP and indicate that *APP<sup>YG/YG</sup>* are a valuable mouse model to study APP functions in physiological and pathological processes.

### Keywords

Amyloid Precursor Protein; YENTP domain; Alzheimer's Disease; Dendritic spines; Cholinergic system; TrkA receptor; behavior

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### AUTHOR CONTRIBUTIONS

Author contributions: C.M. and L.D. designed research; C.M., S.L., C.N., R.T., A.P., M.A.-T., L.R.L.R., Y.L., A.P.M.B., F.B. and F.L. performed research; C.M., S.L., C.N., A.P., M.A.-T., L.R.L.R., H.Z., Y.L., R.T. and L.D. analyzed data; C.M., S.L. and L.D. wrote the paper.

## INTRODUCTION

APP is extensively studied because processing of APP is linked to AD pathogenesis. AD is characterized by extracellular brain deposition of amyloid plaques and intra-neuronal neurofibrillary lesions (Hardy & Selkoe, 2002). Amyloid plaques consist primarily of A $\beta$  peptides, which are produced by a sequential  $\beta$ - and  $\gamma$ -secretase cleavage of APP (Hardy & Selkoe, 2002). Some familial forms of dementias are caused by mutations in either *APP* or genes that regulate APP processing, *BRI2/ITM2B* and *PSEN1/PSEN2* (Hardy & Selkoe, 2002; Fotinopoulou et al., 2005; Matsuda et al., 2005, 2008, 2011; Vidal et al., 1999), underlying the relevance of APP processing to AD pathogenesis. The prevalent pathogenic model of AD posits that aggregates of A $\beta$  trigger dementia (Hardy & Selkoe, 2002). However, new evidence suggests that alterations of APP functions/processing contribute to AD pathogenesis (Tamayev et al., 2011; Tamayev et al., 2012), and that it is therefore important to understand the role of APP *in vivo*.

The intracellular region of APP is functionally important. Numerous proteins bind this region of APP and these protein-protein interactions regulate processing and functions of APP (Matsuda et al., 2003; Roncarati et al., 2002; Scheinfeld et al., 2003a, 2003b). Most interactions of APP involve the YENPTY sequence (amino acids 682–687, following the numbering of 695 amino acids long brain APP isoform). Phosphorylation of Y<sup>682</sup> is consequential. Some proteins, such as Grb2 (Russo et al., 2002; Zhou et al., 2004), Shc (Russo et al., 2002; Tarr et al., 2002), Grb7 and Crk (Tamayev et al., 2009) interact with APP only when Y<sup>682</sup> is phosphorylated; others, like Fe65, Fe65L1 and Fe65L2 only when this tyrosine is not phosphorylated (Zhou et al., 2009), suggesting that phosphorylation–dephosphorylation on Y<sup>682</sup> modulates APP functions.

To test the *in vivo* function of Y<sup>682</sup> we have created mice with Y<sup>682</sup> replaced by G (*APP<sup>YG/YG</sup>* mice which express the mutant APP<sup>YG</sup> protein). When this point mutation is introduced into an *APLP2-KO* background, *APP<sup>YG/YG</sup>/APLP2<sup>-/-</sup>* mutant mice exhibit neuromuscular synapse deficits and early lethality (Barbagallo et al., 2011), similar to *APP/APLP2* double KO mice (Li et al., 2010) suggesting an essential role of Y<sup>682</sup> in development and/or aging. *APP<sup>YG/YG</sup>* mice also show a decrease in APP processing by  $\beta$ -secretase, leading to a reduction in A $\beta$  peptides, as compared to wild-type littermates (Barbagallo et al., 2010), showing that Y<sup>682</sup> plays a pivotal role in regulating APP processing.

Several data have linked TrkA signaling to APP processing (Matrone et al., 2008, 2009). Conversely, *APP<sup>YG/YG</sup>* mice have impaired NGF/TrkA signaling (Matrone et al., 2011), showing that APP (and in particular Y<sup>682</sup>) regulates TrkA function. Altogether, these data underscore a close crosstalk between these two membrane proteins.

Here, we have further characterized the *APP<sup>YG/YG</sup>* mice and analyzed whether this Y<sup>682</sup>G mutation causes developmental and/or aging-dependent deficits.

## RESULTS

### Normal neurological functions in *APP<sup>YG/YG</sup>* mice

Visual inspection did not reveal obvious difference between *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* mice. *APP<sup>YG/YG</sup>* mice maintained a normal gait and posture and no abnormal stereotyped behaviors or signs of ataxia or dystonia were observed. Eyes were normally open and lacrimation was absent. Fur appeared tidy and well groomed, piloerection was absent, and whiskers were intact in all mice. Righting reflex showed comparable results for the two

genotypes (not shown). Mice were weighed monthly until 7M of age. No differences between *APP<sup>YG/YG</sup>* and *APP<sup>WT/WT</sup>* mice were detected in the growth curve (Fig. 1A).

### ***APP<sup>YG/YG</sup>* mice present altered neuromuscular and physical performances, reduced paired-pulse ratio and increased synaptic failure**

To search for possible abnormalities in muscular strength we performed the hang wire test in young and adult *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* mice. Latencies to fall in hanging wire test were similar in 2M old *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* mice, while it was significantly reduced in 5M old *APP<sup>YG/YG</sup>* mice (Fig. 1B).

Two-month old *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* mice showed similar locomotor activity, as measured by the number of crossings during 60 min of free exploration in shuttle-box cages (Fig. 1C). At 5M of age, *APP<sup>WT/WT</sup>* mice showed a significantly reduced locomotor activity than that at 2M of age, while 5M old *APP<sup>YG/YG</sup>* mice were significantly more active compared to age-matched WT mice (Fig. 1C).

To test for possible motor coordination impairments we performed rotarod test at accelerated rate. Rotarod test was preceded by few trials of training in which mice were habituated to stay in equilibrium on the rod at fixed rate. Fig. 1D show the mean rotarod performances, recorded in *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* mice at 2M (left panel) and 5M (right E) of age, expressed as mean time on the rod before falling during both the four trials of training (trials 1–4) with rotarod at fixed rate and the eight trials of testing (trials 5–12) with accelerated rotarod. Interestingly, during the training phase *APP<sup>YG/YG</sup>* mice learned less quickly than *APP<sup>WT/WT</sup>* mice to stay in equilibrium in the rod rotating at fixed speed. This is particular evident in 5M old *APP<sup>YG/YG</sup>* mice. RMANOVA showed no significant differences in mice at 2M of age ( $F_{1,14}=2.996$ ;  $p=0.1055$ ) and confirmed significant difference between the two genotypes at 5M of age during training phase (trials 1–4) with significant main effect for genotypes ( $F_{1,13}=8.806$ ;  $p=0.0109$ ), for trials ( $F_{3,33}=8.947$ ;  $p=0.0001$ ) and significant genotypes  $\times$  trials interaction ( $F_{3,33}=4.625$ ;  $p=0.0073$ ). Post-hoc comparison showed significant differences between *APP<sup>YG/YG</sup>* mice and *APP<sup>WT/WT</sup>* during training at trials 1–3. Although this initial impairment, at the end of training phase, i.e. at trial 4, *APP<sup>YG/YG</sup>* mice reached the same performance of *APP<sup>WT/WT</sup>* mice. During testing phase, with rod rotating at accelerated speed, *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* mice behaved almost identical, without significant differences both at 2M and 5M of age (RMANOVA; 2M:  $F_{1,14}=0.363$ ;  $p=0.5562$ ; 5M:  $F_{1,13}=0.009$ ;  $p=0.9254$ ).

By means of accelerated treadmill we tested the physical resistance of mice during an endurance exercise. Fig 1E–G show that 2M old mice behaved similarly both in total running time, total distance and work performed. However, at 5M of age *APP<sup>YG/YG</sup>* mice showed a significant reduction in total running time, total distance and vertical work performed.

To determine if abnormal neuromuscular transmission is correlated with altered motor function seen in *APP<sup>YG/YG</sup>* mice, we evaluated endplate potential (EPP) in diaphragms of 5M old *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* mice. We first compared single electrical stimulation induced EPP, the *APP<sup>YG/YG</sup>* EPP showed similar amplitude compared to controls (*APP<sup>WT/WT</sup>*:  $8.8 \pm 0.7$ ,  $N=19$ ; *APP<sup>YG/YG</sup>*:  $8.5 \pm 1.04$ ,  $N=20$ ). We next performed paired-pulse protocol by applying two stimulating pulses at 20, 50, 100 and 200 msec intervals in control and *APP<sup>YG/YG</sup>* NMJs. A significantly lower ( $p<0.05$ ) paired-pulse ratio was observed in *APP<sup>YG/YG</sup>* diaphragm at inter-pulse intervals of 20 and 50 msec (Fig. 1H), which is similar to what have been seen in *APP<sup>-/-</sup>* mouse model though the later demonstrated more severe deficits (Yang et al., 2007). We then tested the response of *APP<sup>YG/YG</sup>* NMJs to 1 sec stimulus train at 50, 100 and 200 Hz. Synaptic failure was observed at high frequency (200

Hz) in 5 out of 10 recordings in *APP<sup>YG/YG</sup>* mice, which was 2 of 13 in *APP<sup>WT/WT</sup>* mice (Fig. 1I).

Altogether, the results show that, compared to WT mice, *APP<sup>YG/YG</sup>* mice are normal in locomotor activity and motor coordination but develop impairment in physical resistance and a reduced neuromuscular strength.

### Altered cognitive performances of *APP<sup>YG/YG</sup>* mice

To test ability of learning and memory in *APP<sup>YG/YG</sup>* mice during learning of conditioned task we performed the active avoidance (AA) test. AA represents a behavioral paradigm to evaluate associative learning and the retention of conditioning events. Fig. 2A and B shows the avoidance responses during 5 days of 100-trial of AA sessions in *APP<sup>YG/YG</sup>* and *APP<sup>WT/WT</sup>* mice at 4M and 7M of age. As previously reported (Bovet et al., 1969) for C57BL/6J mice learned this task quite poorly. In fact, only 30% and 45% of AA were recorded at 4M and 7M of age, respectively. A completely different behavior was observed in *APP<sup>YG/YG</sup>* mice. At 4 months of age, these mice appeared to learn more rapidly to avoid electric shock than *APP<sup>WT/WT</sup>* mice during the fifth session (Fig. 2A). In contrast, at 7M of age the *APP<sup>YG/YG</sup>* mice were significantly impaired during sessions 4 and 5, compared to WT mice (Fig. 2B). RMANOVA of the avoidance responses of 4M-old mice showed that there was not significant effect for genotype ( $F_{1,20}=1.911$ ;  $p=0.1821$ ), but a significant effect of daily session ( $F_{4,40}=39.415$ ;  $p<0.0001$ ) and a significant genotype  $\times$  session interaction ( $F_{4,80}=2.818$ ;  $p=0.0305$ ). In 7M old mice, we found a main effect for genotype ( $F_{1,20}=14.810$ ;  $p=0.001$ ), for avoidance session ( $F_{4,40}=69,103$ ;  $p<0.0001$ ) and for genotype  $\times$  avoidance session interaction ( $F_{4,80}=11.952$ ;  $p<0.0001$ ). Post-hoc comparison showed significant differences between *APP<sup>YG/YG</sup>* and *APP<sup>WT/WT</sup>* mice only during sessions 5 at 4M of age (improved learning) and during sessions 4 and 5M at 7M of age (impaired learning).

To further test memory, we performed novel object recognition (NOR) and radial-arm water maze (RAWM) tests. NOR is a non-aversive task that relies on the mouse's natural exploratory behavior. Open field studies showed that *APP<sup>WT/WT</sup>*, *APP<sup>YG/WT</sup>* and *APP<sup>YG/YG</sup>* mice have no defects in habituation, locomotor and anxiety-like behavior (Fig. 2C and D). Mice were analyzed at 4, 5, 8 and 13M of age. At all ages, mice of all genotypes spent the same amount of time exploring the two identical objects during the training session, (Fig. 2E, G, I and K). At 4M of age, when a novel object was introduced the following day, *APP<sup>WT/WT</sup>*, *APP<sup>YG/WT</sup>* and *APP<sup>YG/YG</sup>* mice spent more time exploring the novel object (Fig. 2F). However, 5M-old *APP<sup>YG/YG</sup>* mice spent the same amount of time exploring the two objects as if they were both novel to them, while *APP<sup>YG/WT</sup>* mice behaved like the *APP<sup>WT/WT</sup>* and explored preferentially the novel object (Fig. 2H). At 8 and 13M of age, both *APP<sup>YG/WT</sup>* and *APP<sup>YG/YG</sup>* mice showed novel object recognition memory deficits (Fig. 2J and L), showing that the *APP<sup>YG</sup>* mutation causes severe aging- and dose of mutation-dependent memory deficits.

We performed the spatial working memory test RAWM, which depends upon hippocampal function and tests short-term memory. Mice were required to learn and memorize the location of a hidden platform in one of the arms of a maze with respect to spatial cues. At 4 and 6M of age *APP<sup>WT/WT</sup>*, *APP<sup>YG/WT</sup>* and *APP<sup>YG/YG</sup>* mice were able to acquire (A) and retain (R) memory of the task (Fig. 2M and N). Seven-eight months old *APP<sup>YG/YG</sup>* mice showed abnormalities during both acquisition and retention of the task (Fig. 2O), and at 13M of age both *APP<sup>YG/WT</sup>* and *APP<sup>YG/YG</sup>* mice showed severe impairment in short-term spatial memory for platform location during both acquisition and retention of the task (Fig. 2P) This defect was due to a deficit in memory *per se* and not to deficits in vision, motor coordination, or motivation because testing with the visible platform showed no difference

in the time needed to find the platform and swimming speed between the *APP<sup>WT/WT</sup>*, *APP<sup>YG/WT</sup>* and *APP<sup>YG/YG</sup>* mice (Fig. Q and R). Taken together, these findings provide compelling genetic evidence that Y<sup>682</sup> of APP plays an important function in normal memory formation.

### Region-specific and age-dependent dendritic spine loss in *APP<sup>YG/YG</sup>* mice

Synaptic loss correlates to cognitive impairment (Arendt, 2009). Dendritic spines, the protrusions emerging from the dendritic shaft of neurons, are the primary sites of excitatory synapses and their density reflects the synaptic connectivity of the neural circuits they belong to. Accordingly, the impact of the Y<sup>682</sup>G mutation on dendritic spine density was analyzed in Golgi-stained CA1 pyramidal hippocampal and spiny dorsolateral striatal (DSL) neurons involved in cognitive and motor learning, respectively. The ANOVA performed on hippocampal spines (Fig. 3A) counted in the apical dendrite compartment revealed a significant main effect of genotype ( $F_{1,101}=60.63$ ;  $p<0.001$ ), of age ( $F_{1,101}=5.126$ ;  $p<0.05$ ), and a significant genotype  $\times$  age interaction ( $F_{1,101}=9.30$ ;  $p<0.01$ ). Post-hoc comparisons showed that spine density was significantly lower in *APP<sup>YG/YG</sup>* mice than in *APP<sup>WT/WT</sup>* mice at both age-points ( $p<0.01$ ). Importantly, although spine density was stable across age-points in the *APP<sup>WT/WT</sup>* mice ( $p>0.05$ ), it was significantly lower in 7M old than in 2M old mutants ( $p<0.001$ ) thus indicating a mutation-specific age-dependent spine loss. The ANOVA performed on hippocampal spines counted in the basal dendrite compartment also revealed a significant main effect of genotype ( $F_{1,162}=162.6$ ;  $p<0.001$ ), of age ( $F_{1,162}=28.61$ ;  $p<0.001$ ), but no significant genotype  $\times$  age interaction ( $F_{1,162}=1.47$ ;  $p>0.1$ ). Thus, differently from what observed for apical dendrites, age-dependent spine loss was evident in both genotypes, although it was considerably stronger in *APP<sup>YG/YG</sup>* mutants. The ANOVA performed on DSL spines (Fig. 3B) also showed a main effect of genotype ( $F_{1,118}=60.23$ ;  $p<0.001$ ), of age ( $F_{1,118}=6.270$ ;  $p<0.05$ ) but no significant genotype  $\times$  age interaction. As for hippocampal basal dendrites, less spines were counted on DSL neurons of *APP<sup>YG/YG</sup>* at each age-point therefore revealing that spine density decreased with age in both genotypes, this effect being more marked in *APP<sup>YG/YG</sup>* mice than in *APP<sup>WT/WT</sup>* mice. Altogether, our data provide the evidence that the Y<sup>682</sup> mutation severely disrupts connectivity in hippocampal and DLS circuits.

### Reduced expression levels of ChAT and TrkA proteins in brain from *APP<sup>YG/YG</sup>* mice

Several data indicate a correlation between muscular functions and cholinergic tone (Schliebs & Arendt, 2011). Cholinergic neurons in the CNS undergo complex changes during normal ageing, leading to a reduced support of NGF via TrkA/p75 receptors (Schliebs & Arendt, 2011; Niewiadomska et al., 2011), and deficit in NGF signaling causes atrophy and loss of cholinergic neurons (Ruberti et al., 2000). These data and the evidence showing impaired NGF/TrkA signaling in *APP<sup>YG/YG</sup>* mice (Matrone et al., 2011), prompted us to analyze the expression levels of choline acetyltransferase (ChAT) and TrkA proteins in different brain structures, such as hippocampus (Hp), striatum (Str), cortex (Cx) and septum (Spt). *APP<sup>YG/YG</sup>* mice show a progressive reduction of the expression levels of ChAT and TrkA in Hp and Spt. TrkA, but not ChAT expression, was decreased also in striatum of *APP<sup>YG/YG</sup>* mice. Neither protein was reduced in the Cx. Such reductions reached the peak at 6M of age (Fig 4A, B and E). Consistent with the WB analysis, a significant reduction in the number ChAT positive neurons was observed in the hippocampus, septum as and the caudatum of 6M-old *APP<sup>YG/YG</sup>* mice (4C and D). Furthermore, the reduction of the expression of ChAT protein and the significant loss of the cholinergic stained nuclei are associated to the decrease of the septum volume (4D), indicative of cholinergic tone impairment in such area. Conversely the hippocampus and caudate volumes are not affected.

## DISCUSSION

A role of Y<sup>682</sup> of APP in development was previously reported. Mutation of Y<sup>682</sup> into a G alters APP trafficking and processing leading to a redistribution of APP towards the non-amyloidogenic pathway (Barbagallo et al., 2010). Moreover, Y<sup>682</sup> plays a fundamental role in the functions of APP necessary for normal development as shown by the evidence that *APP<sup>YG/YG</sup>/APLP2<sup>-/-</sup>* mice present with postnatal lethality and neuromuscular synapse defects similar to doubly deficient *APP/APLP2* mice (Barbagallo et al., 2011). Thus, Y<sup>682</sup> is indispensable for the essential function of APP in developmental regulation. Additionally, an intimate crosstalk between APP and NGF pathways has been described, and Y<sup>682</sup> plays a central role in this functional relationship (Matrone et al., 2008, 2009 and 2011; Calissano et al., 2010).

Here, we show that APP regulates neuronal homeostasis and ageing via Y<sup>682</sup>. The *APP<sup>Y682G</sup>* mutation causes an ageing-dependent neuronal decline, leading to ageing-dependent reduction in: (i) muscular strength; (ii) physical performance during treadmill protocol of endurance exercise; (iii) motor learning performance; (iv) AA learning (v) object recognition and short-term spatial working memory; (vi) hippocampal and DSL synaptic connectivity; (vii) cholinergic tone and TrkA levels in hippocampus and septum.

*APP<sup>YG/YG</sup>* mutant mice do not have motor coordination deficit but are less vigorous and more prone to fatigue than *APP<sup>WT/WT</sup>* mice already at 5M of age. This is probably due to reduced neuromuscular strength as suggested by EPP data. Interestingly, *APP<sup>YG/YG</sup>* mice also develop ageing-dependent learning and memory deficits. Active avoidance test shows a cognitive decline in 7M-old *APP<sup>YG/YG</sup>* mice, which is similar to that observed in 10M-old *APP<sup>-/-</sup>* mice (Dawson et al., 1999). *APP<sup>YG/YG</sup>* and *APP<sup>YG/WT</sup>* mice also develop ageing-dependent deficits in object recognition and spatial working memory. Consistent with the poor cognitive and motor performance, *APP<sup>YG/YG</sup>* mice showed massive synaptic loss in the hippocampus which controls spatial working memory and long term object recognition (D'Amato et al., 2011) and in the DSL which, apart from its role in motor functions, is selectively involved in AA learning (Ammassari-Teule et al., 2009). Interestingly, spine loss was already evident at 2M of age, when *APP<sup>YG/YG</sup>* mice do not yet show behavioral impairments, suggesting that a critical synaptic loss threshold has to be reached before cognitive and motor functions are altered and that synaptic loss precedes behavioral deficits.

APP transgenic mice, which are widely used as animal models of AD, develop memory deficits similar to those seen in *APP<sup>YG/YG</sup>* animals. The memory deficits of APP transgenic mice are attributed to increased production of A $\beta$  and amyloid burden. However, *APP<sup>YG/YG</sup>* mice show a significant decrease in A $\beta$  levels (Barbagallo et al., 2010), suggesting that alterations of APP function/processing can lead to memory deficits that are not linked to A $\beta$ . Whether similar mechanisms can also lead to aging-dependent forms of dementias without amyloidosis in humans it remains to be determined.

Although it is tempting to speculate that the compromised crosstalk between NGF/TrkA signaling and APP processing is mechanistically linked to the decline in cholinergic tone, cognitive and neuromuscular performance of *APP<sup>YG/YG</sup>* mice, further experiments will be needed to establish direct cause-effect relationships among these deficits. Nevertheless, the profound deficits caused by this single point mutation of APP indicate that *APP<sup>YG/YG</sup>* mice can be useful to dissect the mechanistic aspects of APP function in physiological and pathological processes.

## EXPERIMENTAL PROCEDURES

### Preliminary observation

The animals used for these studies were backcrossed to C57Bl6/J mice for at least 14 generations. Mice were handled according to NIH and Italian ethical regulations. *APP<sup>YG/YG</sup>* mice were used in all the experiment with C57BL/6J mice serving as wt controls. For NOR and RAWM, *APP<sup>YG/YG</sup>*, *APP<sup>YG/WT</sup>* and *APP<sup>WT/WT</sup>* mice used were littermates derived from crossings of male and females *APP<sup>YG/WT</sup>* mice. For preliminary observations and behavioral assessment, 8–10 *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* were tested. Sequence of behavioral examinations was as follow: at day 1, mice were subjected to preliminary neurological screening and to tail-suspension and righting reflex tests; after 1 day of rest, at day 3, mice were subjected to locomotor activity followed again by 1 day of rest; at day 5, mice begun treadmill running test which was protracted for 9 days with 1 day of rest between each running sessions; finally, starting from day 23, mice performed 7 days of rotarod test. All testing were carried out blindly. Gait and posture, tremors, palpebral closure and lacrimation, piloerection and whisker appearance, grooming and defecation were taken as indexes of general health status. Visual inspection was performed by putting mouse into a white plastic sheet surrounded by a clear acrylic cylindrical viewing jar (14 cm diameter; 18 cm height) and observing mouse behavior for 5 min. After 5 min observation, mice were subjected to righting reflex. Turning the mouse on its back and recording the latency to regain the footing position from the back position evaluated righting reflex.

### Assays of neuromotor behavior

**Hang wire test**—Neuromuscular strength was tested by wire hang test. For this purpose, mice were placed on a steel bar (55 cm length; 2-mm thick) and the ability to hold in equilibrium on the bar with their forelimb/hind limb was measured as latency to fall off a wire after exhaustion. A cut off time of 60 s was considered as the end of test.

**Locomotor activity test**—Locomotor activity was tested in eight toggle-floor cages. Each cage consisted of a rectangular white 40×10 cm Plexiglas box divided into two equal 20×10 cm compartments by a 3×3 cm opening through a black partition. The number of crossings from one side of the box to the other was recorded automatically by means of a micro-switch connected to the tilting floor of the box. Mice were subjected to a 60 min activity test and the number of crossings between the two compartments represented the score for each mouse (Luvisetto et al., 2004).

**Rotarod**, which tests motor coordination, balance and motor learning ability, was performed in three phases, as reported (Luvisetto et al. 2008): habituation (1 day), training (2 days) and testing (4 days). The mice were subjected to two daily trials separated by a 45 min interval. Mice were accustomed to the stationary rod (habituation) and to the rod rotating at 6 rpm (training) for 3 min. During the four consecutive days of testing, mice were placed on the rod rotating at 4 rpm for 30 s, after which acceleration was started and gradually increased to 40 rpm over a time course of 5 min. Testing sessions were terminated after 5 min or when mice fell off the rod.

**Treadmill running exercise**—Exercise studies were performed on five-lane motorized treadmill equipped with an electronic control unit and an electric shock grid at one end of the treadmill. Shock intensity was set at 0.4 mA. Inclination of treadmill was set at 8.5° (15%). Mice were subjected to an endurance protocol with treadmill belt running at incremental speed. Exercise was repeated for 9 days with 1 day of rest between each trial. At each day, mice were first acclimated with treadmill for 1 min, followed by a running session with belt speed initially set at 15 m/min. After 10 min, belt speed was increased by 3 m/min

every 5 min and exercise continued until exhaustion. Exhaustion was considered as inability to maintain running speed despite repeated contact with the electric grid. The time for removal of mice from the treadmill was 5 s on the shocker plate without attempting to reengage the treadmill. The time to exhaustion and whole travelled distance were automatically recorded from the beginning of the running session. Endurance exercise performances were estimated by the time of running (in min), the travelled distance (in meters) and the vertical work (in kgxm) performed by each mouse. Then, for each animal, daily parameters were summed and the cumulative values after nine trials were averaged for the different genotype-age mice groups. According to Massett and Berk (2005), vertical work was calculated as the product of body weight (in Kg) and vertical distance (in meters), where vertical distance was the product between travelled distance and  $\sin(\alpha)$ , where  $\alpha$  is equal to angle of treadmill inclination.

### Cognitive tests

**AA acquisition**—The same apparatus employed to measure locomotor activity was used for measurement of AA acquisition. The apparatus was computer controlled and consisted of eight shuttle-boxes; each one divided into two-20×10 cm compartments, connected by a 3×3 cm opening. A light (10W) was switched on alternately in the two compartments and used as a conditioned stimulus (CS). The CS preceded the onset of the unconditioned stimulus (US) by 5 s, and overlapped it for 25 s. Using this procedure, the light was present in the compartment for 30 s: 5 s alone and 25 s together with the US. At the end of the 30 s period, both CS and US were terminated automatically, and the cycle began in the other compartment. The US was an electric shock (0.2 mA) applied continuously to the grid floor. An avoidance response was recorded when the animal avoided the US by running into the dark compartment within 5 s after the onset of the CS. If animals failed to avoid the shock they could escape it by running from the US. Failure of either avoidance or escape response seldom occurred. Mice were subjected to five daily 100-trial avoidance sessions.

**Spatial working memory**—The task studied with the RAWM test has been described previously (Tamayev et al., 2011). The scores for each mouse on the last 3 days of testing were averaged and used for statistical analysis. Briefly, a six-armed maze was placed into white tank filled with water (24–25°C) and made opaque by the addition of nontoxic white paint. Spatial cues were presented on the walls of the testing room. At the end of one of the arms was positioned a clear 10 cm submerged platform that remained in the same location for every trial in 1 d but was moved approximately randomly from day to day. On each trial, the mouse started the task from a different randomly chosen arm. Each trial lasted 1 min, and errors were counted each time the mouse entered the wrong arm or needed more than 10 s to reach the platform. After each error, the mouse was pulled back to its starting position. After four consecutive acquisition trials, the mouse was placed in its home cage for 30 min, then returned to the maze and administered a fifth retention trial.

**Visible platform testing**—Visible platform training to test visual and motor deficits was performed in the same pool as in the RAWM; however, the arms of the maze were removed. The platform was marked with a black flag and positioned randomly from trial to trial. Time to reach the platform and speed were recorded with a video tracking system (HVS 2020; HVS Image).

**Open Field and NOR**—After 30 min to acclimate to the testing room, each mouse was placed into a 40×40 cm open field chamber with 2 ft high opaque walls. Each mouse was allowed to habituate to the normal open field box for 10 min, and repeated again 24 hours later, in which the video tracking system (HVS 2020; HVS Image) quantifies the number of entries into and time spent in the center of the locomotor arena. NOR was performed as



previously described (Taghialatela et al., 2009). Results were recorded as discrimination ratio, which is calculated by dividing the time the mice spent exploring the novel object, divided by the total amount of time exploring the two objects.

### Statistical analysis of behavioral data

All data are shown as mean  $\pm$  s.e.m. Data for growth curves, hanging wire and locomotor activity, as well as for total time, total distance and work performed during treadmill test, were analyzed with Student's t-test. Statistics for NOR and RAWM tests were performed by one-way analysis of variance (ANOVA). Statistics for rotarod, daily treadmill running, and active avoidance tests were performed by two-factor analysis of variance for repeated measures (RMANOVA), considering genotype-age group as between-subject factor and daily sessions, as repeated measures (within-subject factor). When appropriate, pairwise comparisons between genotype-age groups were carried out using Bonferroni/Dunn test.

### Spine density analyses

**Golgi Cox staining**—*APP<sup>YG/YG</sup>* mice (2 months, N = 5; 7 months, N=5) and wt mice (2 month old n = 4; 7 month old n=5) were sacrificed via transcardially perfusion with 200 ml 0.9% saline solution under deep anesthesia (chloral hydrate 400 mg/kg). Brains were dissected and the two hemispheres were impregnated separately in a Golgi-Cox solution (1% potassium dichromate, 1% mercuric chloride, 0.8% potassium chromate) at room temperature. Samples were then placed in sucrose (30%) for three days, sectioned coronally (100  $\mu$ m) using a vibratome and mounted. The color reaction consists of consecutive steps in water (10 minutes), ammonium hydroxide (30 minutes), water (10 minutes), the developer solution (Kodak fix 100%, 30 minutes) and water (10 minutes). Sections were then dehydrated through successive steps in alcohol at increasing concentrations (50%, 75%, 95% and 100%, 100%) before being closed with slide cover slips. All successive measurements were performed by an investigator blind to the experimental condition of the sample under examination.

**Quantification of spine density**—Hippocampal CA1 pyramidal or DLS spiny neurons were identified under low magnification (20X 0.5 NA) using an optical microscope DMLB Leika. Subsequently, the quantification of dendritic spines was done online at a magnification of  $100 \times 1.5$  NA using a camera (Qimaging Qicam Fast1394) connected to the microscope. Spine density was measured on the apical and basal dendrites of hippocampal CA1 pyramidal neurons. Measurements were performed on segments of CA1 basal dendrites (secondary, tertiary and quaternary branches), of CA1 apical dendrites (secondary and tertiary branches) and of DLS spiny neuron dendrites (secondary and tertiary branches); all data were analyzed using the software Neurolucida. Data collected in each brain region were analyzed by means of two-way ANOVAs with genotype and age-point as main factors. Bonferroni post-hoc tests were then used for pair comparisons where necessary.

### Immunofluorescence analysis

Tissue processing and IHC were performed in free-floating sections. Briefly, mice were anesthetized with 400 mg/kg-1 chloral hydrate (Sigma Aldrich) and transcardially perfused with phosphate buffered saline (PBS) followed by 4% paraformaldehyde in PBS; their brains removed, left in the fixative overnight, equilibrated in sucrose 30% and cryopreserved at  $-80^{\circ}\text{C}$ . IHC was performed on serial sections cut transversely at 30–40- $\mu$ m thickness at  $-25^{\circ}\text{C}$  in a cryomicrotome (Leica Camera) from brains embedded in Tissue-Tek OCT (Sakura). Sections were stained with an  $\alpha$ -ChaT antibody (Santa Cruz, 1:100) overnight. Primary antibody staining was revealed using Cy2-conjugated donkey anti-rabbit (Jackson ImmunoResearch). Nuclei were stained with Hoechst.

## Quantification of cell number and volumes

For medial septum (MS)-diagonal band of Broca (DB), stereological analysis of the number of cells expressing ChAT was performed on one-in-four series of 30- $\mu\text{m}$  free floating coronal sections (120- $\mu\text{m}$  apart), which were analyzed by confocal microscopy throughout the whole rostrocaudal extent of MS-DB (+1.18 to -0.34 from bregma). A total of nine sections were used for data analysis. The same sections and procedure were used to measure ChAT cells in caudate, as well as caudate volume. The reference space of the medial septum encompassed the triangular area that contains 95% or more of the basal forebrain cholinergic neurons dorsal to a line drawn across the tops of the anterior commissures. For hippocampus stereological analysis of the number of cells was performed on one-in-six series of 40- $\mu\text{m}$  free-floating coronal sections (240  $\mu\text{m}$  apart) throughout the whole rostrocaudal extent of hippocampus (-0.94 to -3.08 from bregma). Total cell number was obtained according to the optical disector principle, by systematic sampling of counting frames of 100- $\mu\text{m}$  sides in each section. ChAT positive neurons that appeared in the different focal planes of the frame were included in the count; while ChAT positive neurons in the uppermost focal plane of each section and intersecting the exclusion boundaries of the counting frame were excluded, as defined by the optical disector principle. Total cell number (N) was calculated using the formula  $N = N_v \times V_{\text{ref}}$ , where  $N_v$  is the average cell number per disector volume [corresponding to  $100 \times 100 \times 30 \mu\text{m}^3$  (for MS-DB, MS and caudate) and  $100 \times 100 \times 40 \mu\text{m}^3$  (for hippocampus)] and  $V_{\text{ref}}$  (reference volume) is the total volume of the MS, MS-DB, caudate and hippocampus. The reference volume was obtained multiplying the sum of the traced areas of MS, MS-DB, caudate or hippocampus by distance between sections analyzed. Labeled cells and areas were measured by computer-assisted analysis using the I.A.S. software (Delta Systems, Rome, Italy).

## Western blotting

Equal amounts (10–30 $\mu\text{g}$ ) of proteins were separated on 4–12% Bis-Tris SDS-PAGE gels (Invitrogen), blotted onto PVDF membranes (Millipore) and incubated overnight with the appropriate primary antibody. The antibodies used were: anti- $\beta$ -actin (Sigma),  $\alpha$ -ChAT and  $\alpha$ -TrkA (Santa Cruz).

## Electrophysiology

Mice were anesthetized with isoflurane. Diaphragm preparations with the phrenic nerve supply intact were isolated and gently pinned flat in a sylgard-coated recording chamber. After the diaphragms had equilibrated for about an hour, the nerve was taken up into a suction electrode and the muscle fibers along the main intra-muscular branches nerve were impaled with 3 M KCl – filled glass micropipettes (15 – 25 M $\Omega$ ). Intracellular sharp-electrode recording was performed to record evoked endplate potentials (EPPs) in normal Ringer's solution (30 °C) containing 2.3- $\mu\text{M}$   $\mu$ -Conotoxin GIIIB (Bachem). The Ringer's solution was of the following composition (in mM): NaCl 116, KCl 4.5, MgSO<sub>4</sub> 1, NaHCO<sub>3</sub> 23, NaH<sub>2</sub>PO<sub>4</sub> 1, Dextrose 11, CaCl<sub>2</sub> 2. The stimulation pulses (0.01 msec) were applied via an AMPI Master-8 pulse generator and an AMPI Iso-Flex stimulus isolator. Potentials were amplified via a MultiClamp 700B amplifier (Axon Instruments), digitized at 10 KHz and recorded to a computer using pClamp 9 software (Axon Instruments). Offline data analysis was performed using Clamfit 9, (Axon Instruments) and OriginPro 7.5 (OriginLab). In paired-pulse experiments, EPPs were elicited by nerve stimulation with supramaximal double pulses with interpulse intervals range from 20 to 200 ms. Paired-pulse ration was evaluated by calculating the ratio of P2/P1: where P2 is the EPP amplitude elicited by the second stimulus and P1 is the EPP amplitude elicited by the first stimulus.

## Acknowledgments

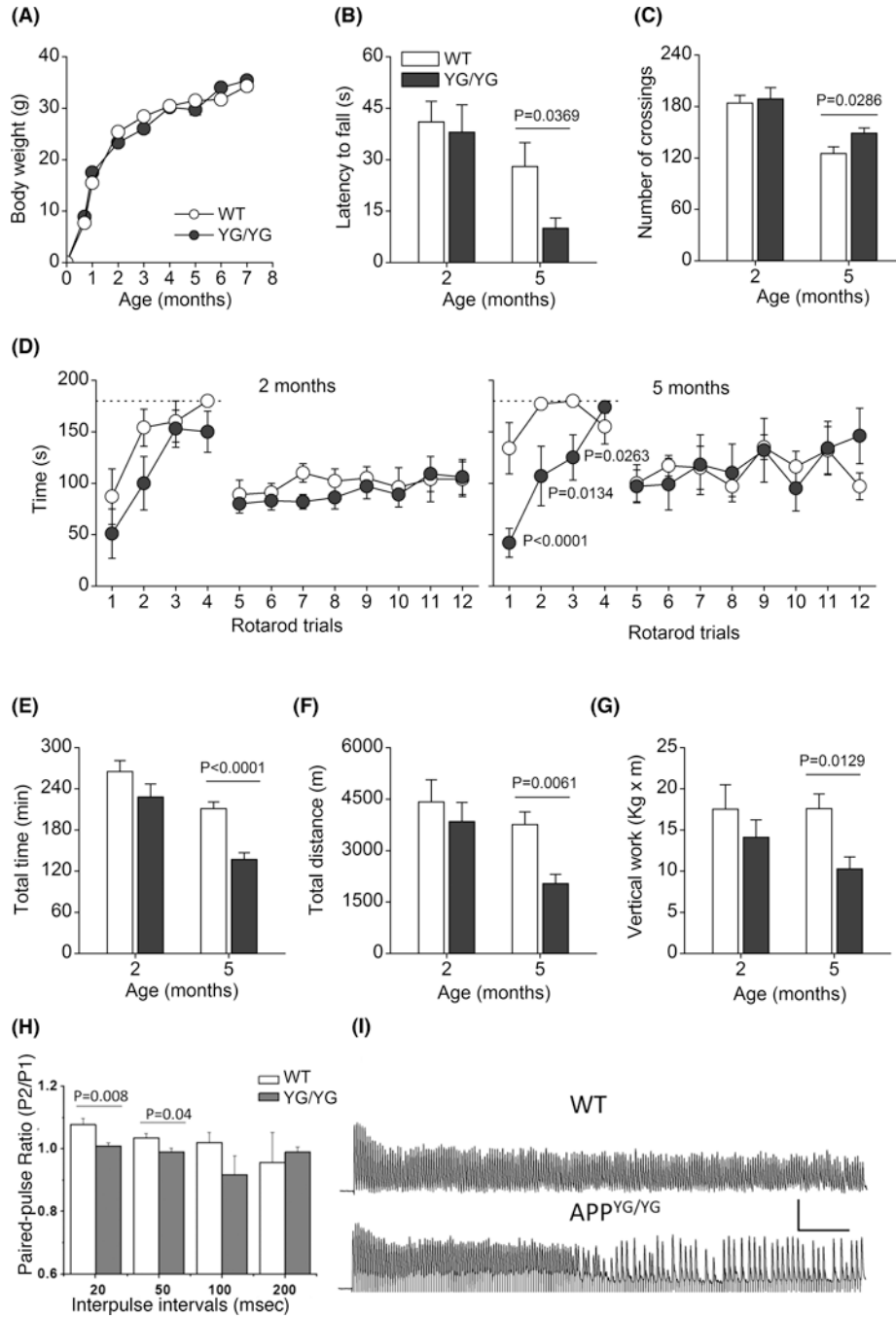
This research was supported by: European Molecular Biology Organization (ASTF264-2010 to C.M.); Alzheimer Association (IIRG-09-129984 and ZEN-11-201425 to L.D.); NIH (AG033007 and AG041577 to L.D) and Thome Foundation (to L.D.); Italian Minister of Health (Project N 263/RF-2009-1536072 to M.A.-T.); the Regione Lazio, as part of the “Distretto Tecnologico delle Bioscienze”; by research grant from FILAS Regione Lazio Funds for “Sviluppo della Ricerca sul Cervello”. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. We greatly appreciate the technical contribution of Maria Teresa Ciotti and the administrative support of Pamela Papa.

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**Figure 1. Behavioral characterization and paired-pulse and 200 Hz train stimulation induced EPPs in NMJs of APP<sup>Y<sup>G</sup>/Y<sup>G</sup></sup> and APP<sup>WT/WT</sup>**

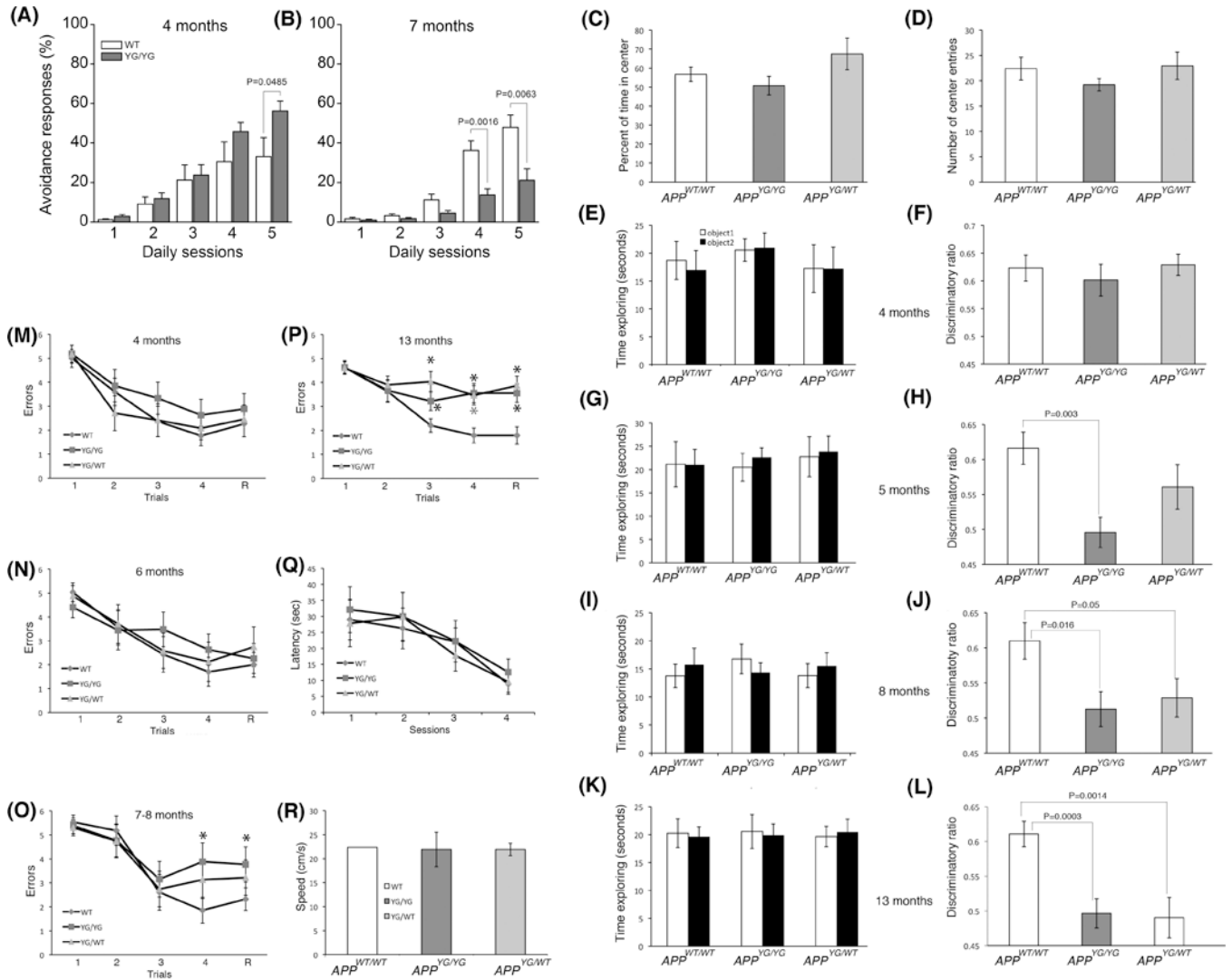
The different panels represent: (A), growth curves from weaning to 7M of age; (B), latency to fall in hanging wire; (C), locomotor activity, measured as number of crossings in shuttle-box cages; (D), time to fall from rotarod during training and testing trials; (E), total time of running during the whole treadmill running; (F), total distance travelled during treadmill running; and (G), vertical work performed during treadmill running. Dashed line in panel D indicates cut-off time, set at 180 sec, during the rotarod training trials. The number of mice was 8–10 for each genotype, except for panel A where the number of mice was 15–17. Data were analyzed by Student’s t-test, except for data in panel D which were analyzed by two-

factors RMANOVA followed by post-hoc pair wise comparison using Bonferroni-Dunn test. (H) Summary of paired-pulse ratio (p2/p1) in *APP<sup>YG/YG</sup>* (N=16) and *APP<sup>WT/WT</sup>* (N=21) at various interpulse intervals. (B) Representative traces showing 200 Hz train stimulation induced responses in *APP<sup>YG/YG</sup>* and *APP<sup>WT/WT</sup>* NMJs. Scale: 5 mV, 100 msec. \*p<0.05.

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**Figure 2. *APP*<sup>YG/WT</sup> and *APP*<sup>YG/YG</sup> mice develop age-dependent memory impairments** (A and B) AA behavioral responses, calculated as cumulative number of shock avoidance during 100 trials over 5 days in *APP*<sup>WT/WT</sup> (WT) and *APP*<sup>YG/YG</sup> (YG/YG) mice at 4M and 7M of age. Number of mice was 8 for each genotype–age group. Statistical analysis: RMANOVA followed by post-hoc pairwise comparison between WT and YG/YG mice, at 4M and 7M months of age, using Bonferroni-Dunn test. (C and D) Open field is a sensorimotor test for habituation, exploratory, emotional behavior, and anxiety-like behavior, in novel environments. The percent of time in center (C) and the number of entries into the center (D) are indicators of anxiety levels. The more the mouse enters the center and explores it, the lower the level of anxiety-like behavior. Since the *APP*<sup>YG/WT</sup> and *APP*<sup>YG/YG</sup> mice are similar to the WT mice, there is no deficit or excess of anxiety. (E and F) Four month-old *APP*<sup>YG/WT</sup> and *APP*<sup>YG/YG</sup> mice have normal object recognition memory. After spending the same amount of time exploring the two identical objects on day 1 (E), the 4M old *APP*<sup>WT/WT</sup>, *APP*<sup>YG/WT</sup> and *APP*<sup>YG/YG</sup> male mice spent more time exploring the novel object 24 hours later (F), showing normal object recognition. (G–L) *APP*<sup>YG/WT</sup> and *APP*<sup>YG/YG</sup> mice show a progressive loss of object recognition memory as they age, which becomes first detectable at 5M of age for *APP*<sup>YG/YG</sup> mice and at 8M of age for *APP*<sup>YG/WT</sup>

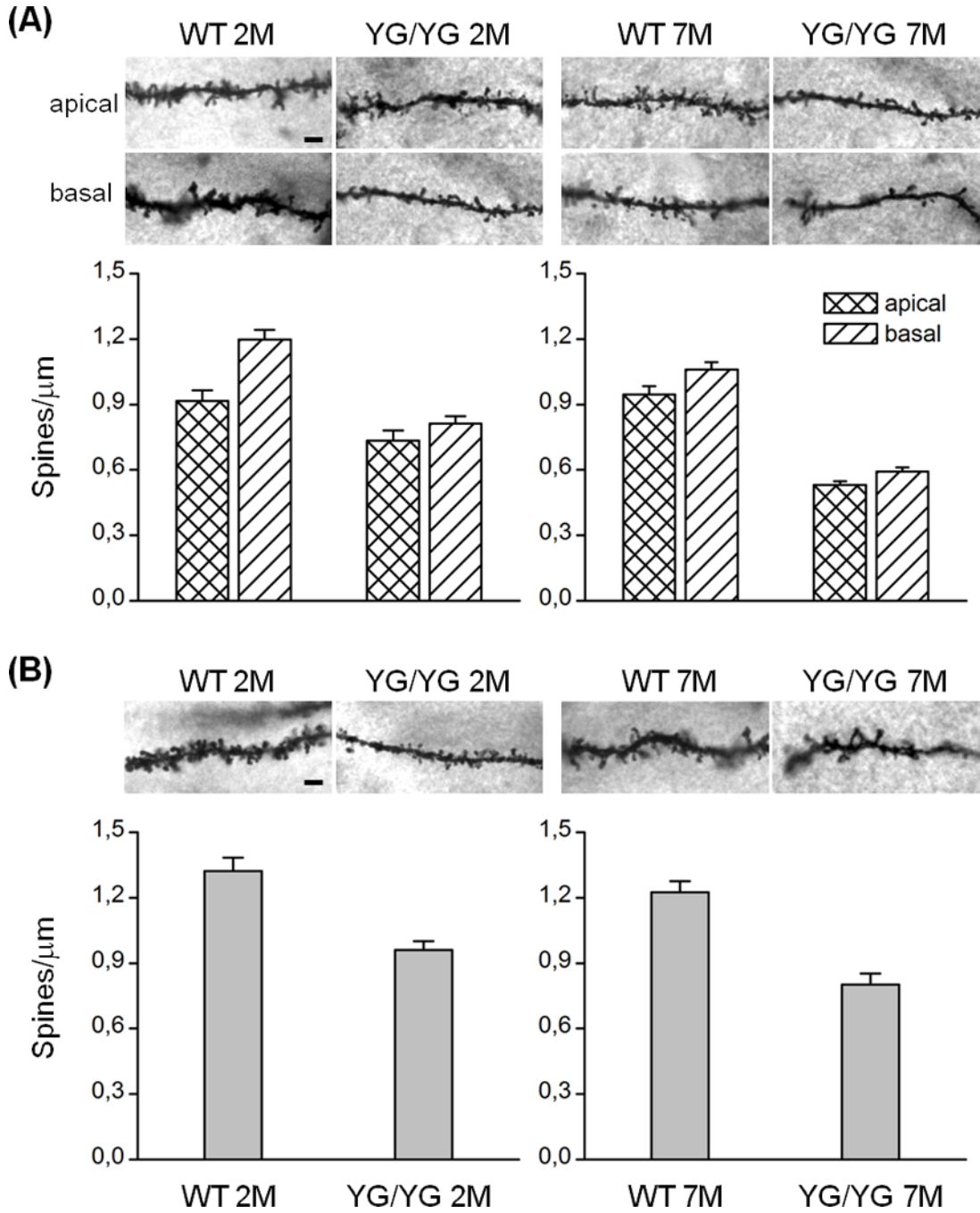


mice. (*M–P*)  $APP^{WT/WT}$ ,  $APP^{YG/WT}$  and  $APP^{YG/YG}$  male mice were run through a RAWM task, testing spatial working memory. At 4M and 6M of age,  $APP^{YG/WT}$  and  $APP^{YG/YG}$  mice made a similar number of mistakes compared to their WT littermates (*M* and *N*). (*O*) At 7–8M,  $APP^{YG/YG}$  mice showed spatial working memory deficits in the fourth acquisition (A4,  $APP^{WT/WT}$  vs.  $APP^{YG/YG}$ ,  $P=0.0004$ ) and the retention (R,  $APP^{WT/WT}$  vs.  $APP^{YG/YG}$ ,  $P=0.010$ ) trials. (*P*) At 13M of age, both  $APP^{YG/WT}$  and  $APP^{YG/YG}$  mice showed short memory deficits in both acquisition and retention trial (A3,  $APP^{WT/WT}$  vs.  $APP^{YG/YG}$ ,  $P=0.034$ ; A4,  $APP^{WT/WT}$  vs.  $APP^{YG/YG}$ ,  $P=0.0008$ ; R,  $APP^{WT/WT}$  vs.  $APP^{YG/YG}$ ,  $P=0.0012$ ; A3,  $APP^{WT/WT}$  vs.  $APP^{YG/WT}$ ,  $P=0.0004$ ; A4,  $APP^{WT/WT}$  vs.  $APP^{YG/WT}$ ,  $P=0.0005$ ; R,  $APP^{WT/WT}$  vs.  $APP^{YG/WT}$ ,  $P=0.0002$ ). (*Q and R*)  $APP^{WT/WT}$ ,  $APP^{YG/WT}$  and  $APP^{YG/YG}$  mice need similar time (latency) and have similar speed (Speed) to reach a visible platform.

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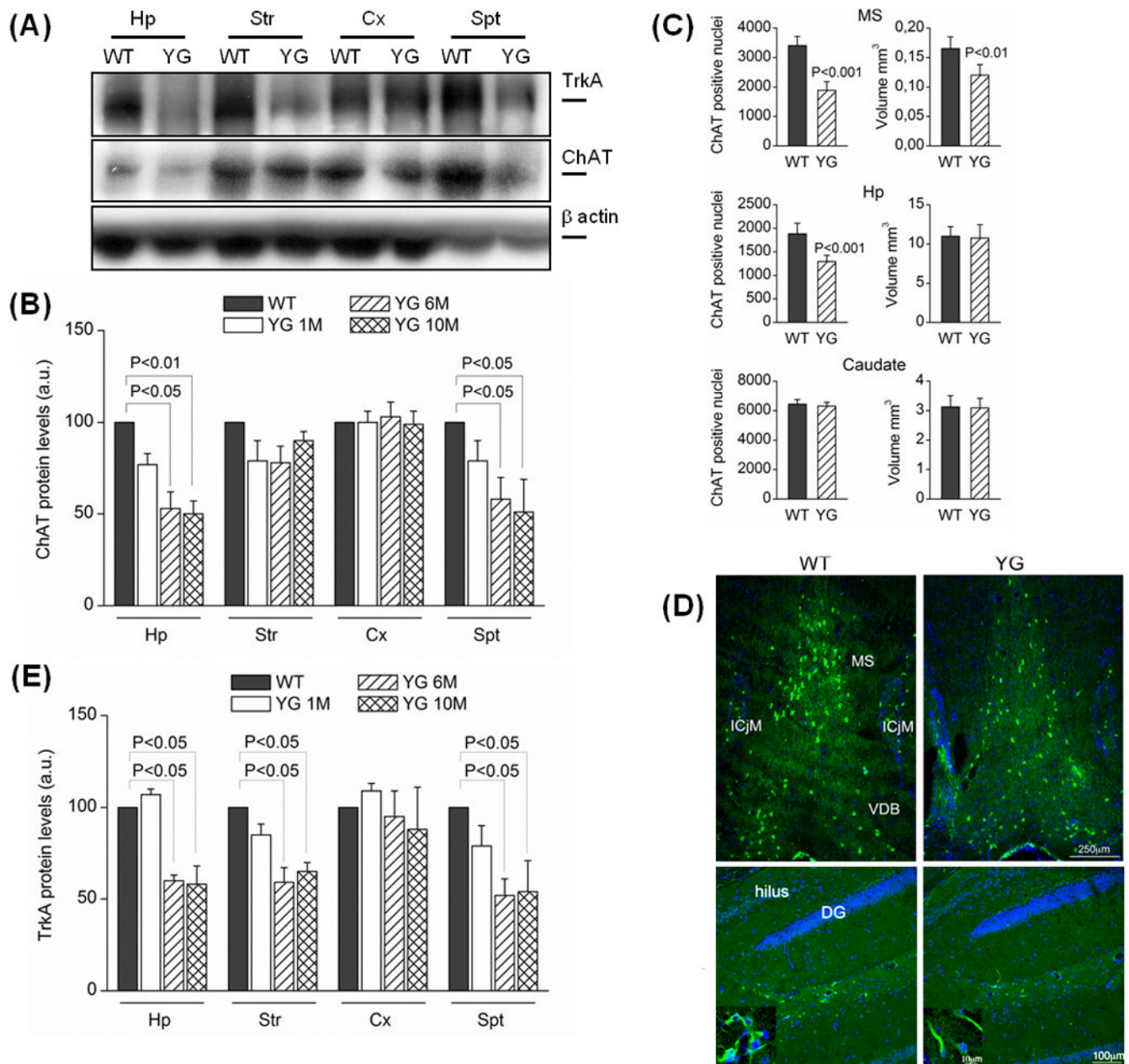
**Figure 3. Aging-dependent reduction in dendritic spine density in  $APP^{YG/YG}$  mice**  
*(A-top)* Representative examples of photomicrographs of Golgi-Cox stained dendritic segments of CA1 pyramidal neurons (apical and basal compartment) in 2M and 7M old  $APP^{WT/WT}$  (WT 2M; WT 7M) and  $APP^{YG/YG}$  (YG/YG 2M; YG/YG 7M) mice, respectively. Magnification: 100 $\times$ 1.25 NA. Scale bar: 3  $\mu\text{m}$ . *(A-bottom)* Effects of genotype on dendritic spine density measured on CA1 pyramidal neuron in basal and apical compartment in 2M and 7M old  $APP^{WT/WT}$  and  $APP^{YG/YG}$  mice. *(B-top)* Representative examples of photomicrographs of Golgi-Cox stained dendritic segments of dorsolateral striatum spiny neuron dendrites in 2M and 7M old  $APP^{WT/WT}$  and  $APP^{YG/YG}$  mice. Magnification: 100 $\times$  1.25 NA, Scale bar: 3  $\mu\text{m}$ . *(B-bottom)* Effects of genotype on dendritic

spine density measured on dorsolateral striatum spiny neuron dendrites 2M and 7M old *APP<sup>WT/WT</sup>* and *APP<sup>YG/YG</sup>* mice.

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**Figure 4. Aging-dependent ChAT and TrkA expression levels in dissected brain areas from *APP<sup>YG/YG</sup>* and *APP<sup>WT/WT</sup>* mice**

(A) Western blots performed in cortex (Cx), hippocampus (Hp), septum (Spt) and striatum (Str) from 1, 6M and 10M old mice. Optical densities (OD) analysis for ChAT (N=4) and for TrkA (N=6) proteins were reported in panel (B) and (E), respectively. Data were normalized on the basis of the correspondent  $\beta$ -actin values and expressed as % of WT (N=7; Test Newman Keuls) (\*)  $p < 0.05$  vs. WT. a.u. (arbitrary unit). (C) Number of ChAT positive nuclei and volume of different brain areas from 6M old mice. Representative pictures of ChAT positive nuclei (green) in septum and hippocampus were reported in panel (D). Symbols: MS, Medial Septum; ICjM, Islands of Calleja-Major; VDB, Vertical Diagonal Band, DG Dentate Gyrus.