

Hippocampal synaptic and membrane function in the DBA/2J-mdx mouse model of Duchenne muscular dystrophy

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Hippocampal synaptic and membrane function in 1 the DBA/2J-mdx mouse model of Duchenne 2 muscular dystrophy 3 4 5 6 Abbreviated title: Duchenne's associated hippocampal hypoexcitability 7 8 Key words: Membrane properties, Duchenne's muscle dystrophy, Hippocampus, Synaptic 9 function, after-hyperpolarization. 10 11 Authors: Riccardo Bianchi^{1,2}, Wouter Eilers³, Federica Pellati², Lorenzo Corsi², 12 Helen Foster³, Keith Foster³ & Francesco Tamagnini¹ 13 14 15 16 ¹ University of Reading School of Pharmacy, Hopkins Building, Whiteknight Campus, Reading, RG6 6LA, UK. 17 ² University of Modena and Reggio Emilia, Department of Life Sciences, Via G. Campi 103-18 19 287, 41125 Modena, Italy 20 ³ University of Reading School of Biological Sciences, Hopkins Building, Whiteknight 21 Campus, Reading, RG6 6LA, UK. 22 23 Corresponding author: Dr Francesco Tamagnini 24 Email: F.Tamagnini@reading.ac.uk 25 T: +44 (0)118 378 4745 26 27

- 29 Highlights
 - Duchenne's muscular dystrophy is often associated with mental retardation
 - The association between dystrophin and excitatory neuronal function in hippocampal physiology is yet largely unexplored
 - We show that hippocampal function is largely preserved in the dystrophinlacking DBA/2J-mdx mouse model
 - Medium after-hyperpolarization is increased in the DBA/2J-mdx mouse model
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37 Abstract

38 Dystrophin deficiency is associated with alterations in cell physiology. The functional 39 consequences of dystrophin deficiency are particularly severe for muscle physiology, as 40 observed in Duchenne muscle dystrophy (DMD). DMD is caused by the absence of a 427 kDa 41 isoform of dystrophin. However, in addition to muscular dystrophy symptoms, DMD is 42 frequently associated with memory and attention deficits and epilepsy. While this may be 43 associated with a role for dystrophin in neuronal physiology, it is not clear what neuronal 44 alterations are linked with DMD. Our work shows that CA1 pyramidal neurons from DBA/2J-45 mdx mice have increased afterhyperpolarization compared to WT controls. All the other 46 electrotonic and electrogenic membrane properties were unaffected by this genotype. Finally, 47 basal synaptic transmission, short-term and long-term synaptic plasticity at Schaffer collateral 48 to CA1 glutamatergic synapses were unchanged between mdx and WT controls. These data 49 show that the excitatory component of hippocampal activity is largely preserved in DBA/2J-50 mdx mice. Further studies, extending the investigation to the inhibitory GABAergic function, 51 may provide a more complete picture of the functional, network alterations underlying impaired 52 cognition in DMD. In addition, the investigation of changes in neuronal single conductance 53 biophysical properties associated with this genotype, is required to identify the functional 54 alterations associated with dystrophin deficiency and clarify its role in neuronal function.

56 Introduction

57 The 427 kDa isoform of dystrophin (Dp427)¹, is present in healthy striated/smooth muscle 58 and neurons, particularly in the hippocampus, prefrontal cortex and cerebellum ^{2,3}. The lack of 59 dystrophin causes Duchenne muscular dystrophy (DMD) which involves progressive wasting 60 of skeletal muscle and the replacement of muscle tissue with connective and adipose tissue, 61 as well as the development of dilated cardiomyopathy in later stages of the disease ^{4,5}. In 62 addition, DMD has been associated with altered brain function, including attention deficits, 63 memory impairments and increased risk of seizures, which suggests a role for dystrophin in 64 brain function⁶. In the brain, neurons expressing the Dp427 isoform include cerebellar Purkinje cells, hippocampal and cortical pyramidal cells and cells in the amygdala⁷⁻¹⁰; other brain 65 isoforms are Dp140 and the Dp71^{11,12}. 66

67 The Dp427 isoform is highly localized in synaptic spines and post-synaptic densities (PSD), 68 where it associates with a dystrophin-associated protein complex (DAPC) which spans the cell membrane and links the actin cytoskeleton with the extracellular matrix ¹³. Both spines 69 70 and PSDs have a critical role for neurotransmission and synaptic plasticity, which underlie 71 cognitive functions, including memory and learning¹⁴. The cognitive impairment associated 72 with DMD has been linked to the effects of the lack of dystrophin in brain structures associated 73 with memory, language and attention, including the hippocampus and the prefrontal cortex^{7,15}. 74 In addition, it should be noted that only one third of DMD patients show cognitive impairment¹⁶. 75 The cognitive impairment can appear, in the absence of dystrophin, even without the muscular 76 dystrophy symptoms, suggesting that the cognitive symptoms are a direct consequence of the 77 lack of protein, rather than an epiphenomenon paralleling and caused by the motor 78 impairment¹⁷. Finally, the lack of dystrophin has been associated with epilepsy, as the 79 prevalence of epilepsy in DMD patients is significantly higher in comparison to control 80 populations¹⁸⁻²¹.

However, it is not clear yet how the lack of the Dp427 isoform of dystrophin may affect the biophysical properties of membrane conductance and synaptic machinery, responsible for neuronal excitability and synaptic transmission and plasticity, respectively. For example, the 84 mdx mouse, the most commonly used mouse model for dystrophin deficiency and DMD, which 85 lacks the expression of the Dp427 kDa dystrophin isoform, has been reported to show impaired spatial memory and enhanced long-term potentiation (LTP) of synaptic transmission 86 87 in the hippocampus^{9,22}, while other research groups have not observed differences in either spatial learning or hippocampal LTP²³. Intriguingly, motivational differences between the *mdx* 88 89 mouse and its wild type control can give conflicting operant conditioning, dependent on the 90 type of behavioural characteristics being assessed²⁴. The reasons for all these discrepancies 91 are unclear; however, the absence of a mechanistic pathway leading from the absence of 92 dystrophin to cognitive dysfunction contributes to the lack of consensus on the causal 93 connection between DMD and neuronal function ²⁵. In addition, such a variety of hippocampal 94 function phenotypes may be associated with the unpredictability of mental retardation onset 95 in people with DMD (only 1/3 of DMD patients show cognitive deficit)¹⁶.

96 GABA-receptor density and clustering on post-synaptic membranes, and GABA dependent 97 inhibitory neurotransmission have been shown to be altered in *mdx* mice. The frequency of 98 GABAergic-dependent spontaneous inhibitory post-synaptic currents (IPSCs) is indeed 99 increased in the amygdala pyramidal neurons of *mdx* mice, revealing a possible effect of this 100 genotype in promoting the hyperexcitability of inhibitory interneurons in this brain area ⁹.

101 Much less is known about the effect of the loss of dystrophin on basic electrotonic and 102 electrogenic properties of neuronal membranes. In the light of all the above, in this study we 103 investigated the effects of the absence of Dp427 on the excitability of hippocampal CA1 104 pyramidal neurons as well as the functionality of Schaffer Collaterals - CA1 synapses in 105 DBA/2J-mdx mice. This mouse line has been generated by backcrossing the more commonly 106 used BI/10-mdx mouse line onto a DBA/2J genetic background which results in a more severe 107 pathology of the muscle (ref 37, Coley et al). Although our study was not designed to determine 108 the effect of genetic background on neuronal parameters, we reasoned that the increased 109 muscle weakness in the DBA/2J-mdx mice compared to the BI/10-mdx mice might compound 110 any effects of the lack of dystrophin per se because of additional mobility restrictions in the

mice due to muscle weakness. To the best of our knowledge, no studies have investigatedneuronal function in this DMD model.

113

114 Materials and methods

115 Experimental animals

Male DBA/2J-*mdx* mice (The Jackson Laboratory, Bar Harbour, ME USA) and DBA2 wild-type mice (Envigo, UK) were group housed in animal facilities at the University of Reading on a 12:12 hour light/dark cycle with standard chow and water available at libitum. In all experiments, DBA/2J-*mdx* mice were compared against age- and sex-matched wild-type mice. All experiments were carried out at the University of Reading under a United Kingdom Home Office licence in compliance with the Animals (Scientific Procedures) Act 1986.

122 Preparation of brain slices.

123 Animals were sacrificed using cervical dislocation in accordance with schedule 1 of the 124 Animals (Scientific Procedures) Act (1986). The brain was rapidly removed and transferred to 125 an ice cold cutting solution consisting of (in mM): 189 Sucrose, 10 D-Glucose, 26 NaHCO₃, 3 KCl, 5 MgSO₄(7 H₂O), 0.1 CaCl₂, 1.25 NaH₂PO₄. Three hundred µm coronal sections were 126 127 cut using a Leica VT1200 microtome and immediately transferred to a holding chamber 128 containing artificial cerebrospinal fluid (aCSF) continuously perfused with carbogen. The 129 composition of the aCSF was as follows (in mM): 124 NaCl, 3 KCl, 24 NaHCO₃, 2 CaCl₂, 1.25 130 NaH₂PO₄, 1 MgSO₄, 10 D-glucose. The slices were then allowed to recover for 30 min at 37° 131 C and subsequently at room temperature for at least 1 hour prior to transfer into a recording 132 chamber.

133 Whole cell patch clamp recordings

Slices were transferred to a recording chamber where they were submerged in carbogenequilibrated aCSF and maintained at a temperature between 33-34 °C. The recording chamber was secured on the stage of an Olympus BX51 upright microscope and individual CA1 pyramidal neurons were visualised using infrared differential interference contrast optics. 138 Borosilicate glass microelectrodes with a resistance ranging from 3-7 M Ω were pulled, fire-139 polished and filled with a K-Gluconate based internal solution consisting of (in mM): 120 mM 140 K-gluconate, 10 mM Na2-phosphocreatine, 0.3 mM Na2-GTP, 10 mM HEPES, 4 mM KCl, 4 141 mM Mg-ATP (pH 7.2, 280-290 mOsm). Following entry into whole cell configuration, a junction 142 potential error of 15 mV arose due to the pairing of the pipette solution with the aCSF, which 143 was corrected for arithmetically during analysis. Signals were recorded using a Multiclamp 144 700A amplifier, digitised using a Digidata 1550B and stored for future analysis using pClamp 145 10 software.

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147 All recordings were made from a defined pre-stimulus membrane potential set by injecting a 148 continuous flow of bias current through the recording electrode. This facilitated the analysis of 149 passive neuronal properties and action potential generation from pre-stimulus membrane 150 potentials (V_m) of both -80 and -74 mV. In order to measure neuronal passive membrane 151 properties a 500 ms, -100 pA hyperpolarising current was injected across the membrane from 152 each V_m. The subsequent voltage deflection at the steady state of the hyperpolarisation was 153 used to calculate the input resistance (R_{in}) of the membrane using Ohm's law (V = IR). The 154 extrapolation of a single exponential curve at an infinite time, fitted to the membrane charging 155 response between 10 and 95% of the peak amplitude, was used to calculate the membrane 156 time constant (T). An approximation of capacitance was measured as the ratio between the T 157 and R_{in}.

Sag, measured as the difference between the negative peak and the steady state
hyperpolarisation, was expressed as a percentage of the peak hyperpolarisation in response
to a 500 ms, -100 pA hyperpolarising current injection.

161 Standard "Zap" protocols were used to measure subthreshold membrane resonance 162 properties. Briefly, the ratio of the Fast-Fourier transform of the voltage response versus the 163 current injection was calculated as a measure of the impedance profile of the pyramidal 164 neurons ($Z = V_{fft}/I_{fft}$). Subsequently, the impedance versus frequency profile was smoothed with a 35-point moving average function. The maximal impedance (Z max) frequency at which this maximal impedance occurred (Peak frequency), and the quality factor of the resonator (Q) were quantified to facilitate comparisons between the genotypes. The quality factor of the resonator "Q" was calculated as the ratio between the impedance at peak frequency and the impedance at 0.5 Hz ²⁶.

170 In order to quantify neuronal excitability, a series of incremental 500 ms depolarising square current injections, ranging from 50 to 350 pA, were injected across the membrane. The 171 172 number of action potentials (AP) generated for each current injection was used as a measure 173 of excitability. The first AP fired in response to a 350 pA depolarising current step was used to 174 compare action potential waveforms between the genotypes. AP threshold was quantified, 175 defined as the voltage at which the rate of rise (dV/dt) surpassed 20 V s⁻¹; the maximal rate of 176 rise (RoR) was also measured for each cell as the highest value of the first derivative of voltage 177 in time, within the AP duration; the AP width was measured at $V_m = -15$ mV; finally, the AP 178 peak was measured as the V_m at which, following the ascending phase of the AP, dV/dt=0 179 V*s⁻¹.

180 For the voltage-clamp experiments, outside-out, somatic, nucleated macropatches were 181 excised as previously described ²⁷. Pipette capacitance was neutralized and the series resistance was compensated for (10% – 80% correction). VC recordings were made for the 182 183 quantitative evaluation of outward-going plateau voltage-gated K⁺ currents, by applying 30 ms 184 voltage steps, growing in 10 mV increments and starting from a holding voltage of -90mV. 185 Each recorded current amplitude was normalized to the membrane capacitance of the 186 macropatch in order to account for differences in current amplitude arising from different sizes 187 of the macropatch. The specific current (pA/pF) was plotted against the value of the voltage 188 step.

189 Field excitatory postsynaptic potentials

Following the initial post-slicing recovery period, the slices were then transferred to a recording
chamber in which they were submerged and continually perfused with aCSF, pre-equilibrated

with carbogen and maintained at room temperature. Field excitatory postsynaptic potentials (fEPSPs) were elicited by delivering a short pulse of electrical current (0.1 ms) through tungsten bipolar stimulating electrodes. These stimulating electrodes were placed such as to stimulate axons of the Schaffer Collateral (SC) pathway. Borosilicate glass recording microelectrodes with a resistance ranging from 3-5 M Ω were pulled and filled with aCSF. Recorded signals were collected with a Multiclamp 700A amplifier, digitised with a Digidata 1550B and stored for future analysis using pClamp 10 software.

199 Input-output curves were constructed by incrementally increasing the current passing through 200 the stimulating electrode and recording the evoked response (0 - 300μ A in 50μ A). For the 201 rest of the experiment, the stimulus intensity was then set to induce 50% c.a. of the maximal 202 response.

203 Short-term plasticity was tested with a paired-pulse profile, consisting in 2 subsequent 204 stimulating pulses, with inter-pulse intervals of increasing value (in ms): 10, 17, 32, 56, 100, 205 170, 320, 560, and 1000.

206 After a period of at least 20 minutes recording baseline responses at low frequency (0.033 207 Hz), induction of LTP was attempted in the SC-CA1 pathway using a theta-patterned (5 Hz) 208 burst stimulation (TBS) protocol. This consisted of 5 bursts of 10 stimuli at 100 Hz applied with 209 an inter-burst interval of 200 ms. This was repeated 4 times with an inter-repeat interval of 20 210 s. The fEPSPs were then followed for 1 hour before the TBS protocol was delivered to the 211 other pathway and the fEPSPs were followed up for another hour. We have decided to use 212 TBS over other existing LTP-induction protocols, such as high frequency stimulation ¹⁴, as 213 TBS mimics the firing patterns observed in vivo in rodents performing a task leading to longterm memory encoding ²⁸⁻³⁰. In addition, TBS results in the most effective NMDA- and voltage-214 215 gated Calcium channel-dependent form of long-lasting LTP ³¹⁻³⁴.

216 Data analysis

217 Data were analysed using Clampfit 11. Statistical assessments of differences between 218 genotypes were made using unpaired two-tailed students t-tests and two-way analysis of variance (ANOVA) as appropriate. Figures were prepared with Origin Pro 2018. All results were expressed as mean ± standard error of the mean (SEM). The experimental "n", refers to the number of slices and to the number of cells we have recorded from in the field potential and patch-clamp experiments, respectively.

223

224 Results

225 Most of the studies investigating the functional correlation between the lack of dystrophin and 226 neuronal and brain dysfunction have focused on the alteration of GABAergic 227 neurotransmission. Our study aimed to characterize, in DBA/2J-mdx mice, the effects of the 228 lack of dystrophin on the electrogenic and electrotonic membrane properties of CA1 pyramidal 229 neurons (CA1-PCs). In addition, we investigated whether basal synaptic transmission, short-230 term synaptic plasticity and long-term potentiation were affected in DBA/2J-mdx mice. Field 231 potential, extracellular recordings were carried out in 9 m/o mice, while single cell, whole-cell 232 patch-clamp recordings were carried out in 7 m/o mice. At the chosen age the mice show 233 significant muscle atrophy and weakness (data not shown). In addition, we have observed that 234 DBA/2J-mdx mice show less rearing activity but no changes in other activity related 235 parameters, when tested in activity cages (unpublished observations). Whether this is due to 236 the significant reduction in muscle strength in these mice, or due to reduced motivation to act 237 or explore is not known. Previous studies have shown that the mdx genotype is associated 238 with cognitive impairment³⁵. However, to our knowledge, there are no available data on the 239 memory performance in DBA/2J-mdx mice.

First, we tested the effect of the DBA/2J-*mdx* genotype on synaptic transmission (Figure 1A; two-way ANOVA; source of variability: genotype; F=3.622, P=0.06), short-term plasticity (Figure 1B; two-way ANOVA; source of variability: genotype; F=0.037, P=0.847) and LTP (Figure 1C; unpaired two-tailed t-test; P=0.97). None of these parameters were affected by the DBA/2J-*mdx* genotype.

Synaptic transmission and plasticity are often regarded as the neuronal correlates of cognitive
 functions, such as memory and learning; hence a change/impairment in these functional

outcomes would be expected to underlie cognitive dysfunction. However, changes in single cell membrane excitability properties are also fundamental functional correlates underlying cognition, even in the absence of synaptic input/output (I/O), (paired-pulse profile) PPP and long-term potentiation (LTP) alterations. For this reason, we measured the electrotonic and electrogenic membrane properties in CA1-PCs from DBA/2J-*mdx* and age-matched WT controls.



Figure 1. DMD-associated DBA/2J-*mdx* **genotype is not associated with impairments of synaptic transmission and plasticity.** A. Basal synaptic transmission was measured as the relationship between fEPSP amplitude and stimulus intensities. Input/output curves were built by plotting the amplitude of SC-CA1 fEPSPs amplitude vs stimulus intensity. Two-way ANOVA does not reveal significant differences between genotypes (F=3.622 p=0.06) nor interactions between genotype and stimulus intensity (F=0.104 p=0.991). B. Paired pulse ratios of SC-CA1 fEPSPs, tested at increasing inter-pulse intervals, were not different between genotypes (two-way ANOVA; F=0.037, p=0.847) and no interaction was observed between genotype and ISI (two-way ANOVA; F=0.499, p=0.855). C. LTP induction in SC-CA1 synapses was not affected by genotype (Unpaired T-Test on the last 10 minutes of follow-up: p=0.68). The reported "n" refers to the number of slices.

- 254 First, no differences were observed in the resting membrane potential (RMP) between
- 255 genotypes. To avoid biases arising from cell-to-cell variability in RMP, all the other properties
- were measured from a pre-stimulus potential of -80 mV, obtained with a constant current



Figure 2. DMD-associated DBA/2J-mdx genotype does not affect the electrotonic, subthreshold membrane properties in hippocampal CA1-PCs. A. V_m average \pm SEM hyperpolarization, upon the injection of a -50 pA, 500 ms current step injection, for each genotype. This approach was used to measure properties like R_{in} (C), Tau (E) and capacitance (G), which were not affected by the *mdx* genotype (unpaired t-test, two tailed). B. Average \pm SEM smoothed impedance (sZ), measured upon the injection of a current oscillating at subthreshold values, with frequencies comprised between 0.5 – 20 Hz, is plotted here. Resonance properties were measured between genotypes and no effect was observed between *mdx* mice and WT controls on peak Z (D), peak frequency (F) and quality factor of the resonator Q (H). The reported "n" refers to the number of cells.

258 injection. No differences were observed in electrotonic passive properties (Figure 1), neither 259 under the injection of a continuous current (Figure 2A, C, E and G) nor the injection of a 260 subthreshold oscillating current, with a frequency linearly increasing between 0.5 Hz and 20 261 Hz within a 30 s time window (Figure 2B, D, F, and G). This latter approach was used to 262 investigate the effects of DBA/2J-mdx genotype on the resonance properties of CA1-PCs, represented as Peak Z, Peak Frequency and quality factor of the resonator – Q, as previously 263 264 described ³⁶. Mean, SEM and P values for subthreshold properties in DBA/2J-mdx and control 265 mice are reported in Table 1.

- 266
- Table 1. Subthreshold passive and resonance membrane properties of CA1
- 268 pyramidal neurons in hippocampal slices from DBA/2J-mdx mice and age-
- 269 matched WT controls.

	WT n=29		<i>mdx</i> n=	<i>idx</i> n=25 cells	
Property	Average	SEM	Average	SEM	Р
RMP (mV)	-71.4 (n=39)	2.6	-73.5 (n=33)	3.0	0.6
Rin (MΩ)	126.0	8.0	146.0	11.2	0.1
tau (ms)	15.3	0.8	17.0	1.3	0.3
Capacitance (pF)	110.0	5.3	105.4	8.3	0.6
Peak frequency (Hz)	3.1	0.3	3.2	0.2	0.7
Q	1.11	0.01	1.10	0.02	0.9
Peak Z (MΩ)	196.5	31.1	180.4	18.8	0.7

- 270
- 271

We tested the firing rate properties of CA1-PCs in DBA/2J-*mdx* and WT controls upon the injection of square current injections of progressively increasing intensity. The average firing rate was not affected by genotype (Figure 3B; two-way ANOVA, source of Variability: genotype, F=0.291, P=0.590).

276 In addition, the AP waverform properties (Figure 4), namely width (Figure 4C), maximal rate 277 of rise (dV_m/dT) of the AP (RoR) (Figure 4D), peak (Figure 4E) and threshold were not affected

278 by genotype. For averages \pm SEM and P values see Table 2.



Figure 3. DMD-associated DBA/2J-*mdx* genotype does not affect the AP firing rates upon depolarization of the V_m, in hippocampal CA1-PCs. A. Example traces of CA1-PCs firing APs upon the injection of 100 - 200 - 300 pA, 500 ms, depolarizing current steps. B. The number of APs fired at each current intensity (50 - 350 pA, with 50 pA increases), within a 500 ms time window, was plotted as a measure of average firing excitability. No effect of the genotype (two-way ANOVA; F=0.291, p=0.590) nor interaction between genotype and current intensity (two-way ANOVA; F=0.745, p=0.590) was observed on CA1-PCs average firing properties. The reported "n" refers to the number of cells.

- 280 Post-firing properties, such as the medium component of after-hyperpolarisation (mAHP),
- were tested to investigate the effect of DBA/2J-mdx genotype on the relative refractory
- 282 period, as an indirect measure of neuronal excitability. mAHPs were evoked by the
- subsequent application of 5-10-15-20-25, 2 nA-2ms depolarising current steps. The
- amplitude of the mAHP, comprised of the repolarization of the last evoked AP and the
- subsequent 500 ms, was measured across number-of-pulses conditions and between
- 286 genotypes. Figure 5A shows



Figure 4. DMD-associated DBA/2J-mdx genotype does not affect the AP waveform properties in hippocampal CA1-PCs. A. Average \pm SEM traces of the first action potential evoked by a 350 pA, 500 ms current step, grouped between genotypes. B. Average \pm SEM phase-plots, grouped between genotypes. CA1-PCs action potential waveform properties, such as width (C), maximal rate of rise (D), peak value (E) and threshold (F) are not different between mdx and WT controls. The reported "n" refers to the number of cells.

the average ± SEM boundaries of the medium and slow components of the AHP evoked by

289 20 pulses, while Figure 5B expands the mAHP component, which is visibly bigger in

- 290 DBA/2J- mdx CA1-PCs. Two-way ANOVA showed a significant effect of genotype in
- 291 increasing the mAHP in DBA/2J-mdx mice, compared to WT controls (Figure 5C; two-way
- ANOVA, source of variability: genotype, F=9.687, P=0.002).



Figure 5. DBA/2J-*mdx* genotype is associated with the increase of the amplitude of post-burst medium but not slow afterhyperpolarizations in hippocampal CA1-PCs. A. Average \pm SEM traces of the mAHP evoked by 20 pulses, 2 nA – 2 ms, delivered at 50 Hz, grouped by genotype. An increased mAHP is observed in CA1-PCs from *mdx* mice, but no genotype effect was apparent on slow AHP (B). C. The overall mAHP, evoked by 5-10-15-20-25 2 nA / 2 ms pulses, delivered at 50 Hz, was bigger in *mdx* in comparison to WT controls (two-way ANOVA, F=9.687, p=0.002); however, no interaction was observed between genotype and number of pulses. The reported "n" refers to the number of cells.

295Table 2. Action potential properties of CA1 pyramidal neurons in hippocampal

	WT n=29 cells		<i>mdx</i> n=25 cells		
Property	Average	SEM	Average	SEM	Р
AP_peak (mV)	18.1	2.7	18.2	2.6	1
AP_width (ms)	0.99	0.05	1.01	0.05	0.7
AP_thres (mV)	-56.0	1.4	-57.5	2.1	0.5
AP_max_dvdt (Vs ⁻¹)	279.5	23.5	278.1	23.6	1

slices from DBA/2J-*mdx* mice and age-matched WT controls.

The generation of mAHP has been ascribed to the gating properties of non-inactivating, 298 299 voltage-gated K⁺ currents, such as I_{K} ³⁶. For this reason, we decided to measure the 300 biophysical properties of voltage-gated non-inactivating I_K. Voltage-clamp recordings on 301 nucleated, somatic, outside-out macropatches were performed and cell-to-cell Boltzmann fits 302 were performed to calculate the maximal current density (I_{max}) and the half-activation potential 303 $(V_{1/2})$. However, no significant differences were observed between genotypes for both the I_{max} 304 (Figure 6; DBA/2J-mdx I_{max} = 99.7 pA/pF ± 54.4 pA/pF vs WT I_{max} = 136.9 pA/pF ± 62.3 pA/pF; 305 P=0.7) and V_{1/2} (DBA/2J-mdx V_{1/2} = -42.6 mV \pm 1.9 mV vs WT V_{1/2} = -43.3 mV \pm 1.02; P=0.7).

306 **Discussion**

307 We investigated the effects of the lack of dystrophin on both CA1-PCs intrinsic membrane 308 excitability and SC-CA1 glutamatergic synaptic function in a mouse model of muscular 309 dystrophy, the DBA/2J-mdx^{37,38}. The DBA/2J-mdx mouse model, while carrying the same 310 mutation, shows a more severe phenotype, in comparison to mdx mice on a BI/10 genetic 311 background ^{37,38}. In fact, the DBA/2J background contains genetic modifier loci, which are responsible for the increased severity of symptoms observed in the DBA/2J-mdx mice ³⁹. To 312 313 our best knowledge, this is the first-time brain function has been tested in this model. Our main 314 observation was that while synaptic function, electrotonic and firing properties were unaffected 315 by the genotype, the mAHP was bigger in DBA/2J-mdx mice than in controls.

316 We have chosen to carry out the recordings on adult mice, aged 7 and 9-month-old. These 317 ages have been chosen because, while DMD is a disorder mostly affecting people during 318 development, we were interested to assess the long-term effects of dystrophin deficiency on 319 hippocampal function. In fact, understanding and predicting brain dysfunction associated with 320 DMD, may have a critical role on people's quality of life, should a novel treatment for DMD be 321 developed. While investigating basal synaptic transmission and short-term synaptic plasticity 322 in dystrophin deficient models is novel, LTP has been tested by other groups, showing 323 contrasting results: while some have observed enhanced LTP in dystrophin deficient mice ^{9,22}, others have not found any effect ²³. Our results fit with the latter. Part of the explanation for 324 325 the discrepancy



Figure 6. The DBA/2J-*mdx* genotype does not affect non-inactivating K⁺ currents in CA1-PCs. A. Example of an outside-out nucleated somatic macropatch, pulled from a CA1-PCs at the end of current-clamp recordings. B. Example trace of inward and outward currents evoked by a +30 mV voltage step: because of the fast inactivating kinetics of I_{Na} and I_A, we focused our measures on the inactivating component of voltage-gated outward currents. C. I_K current densities from *mdx* and WT controls plotted versus the intensity of the voltage step. Cell-to-cell fitting of a Boltzmann function was performed to measure somatic I_K maximal current density and V_{1/2}. Cell-to-cell analysis of the biophysical properties of CA1-PCs' I_K, revealed no effect of the *mdx* genotype on somatic I_K V_{1/2} (C) and maximal current density (D). Scale bar: 5 µm.

327 between the results obtained by different groups may be the different conditions used in each 328 study. First, the LTP-inducing stimulation protocol used in each study is different. In fact, both 329 repeated, strong (2x1s at 100Hz every 20 s) and single, weak (1x1s at 30Hz) high frequency 330 stimulation resulted in a genotype-dependent LTP enhancement in mdx but not in mdx^{3cv} mice ^{9,22}: however, *in vivo* LTP experiments on *mdx* mice, using 3x1s 100 Hz every 20 s HFS, did 331 not show any genotype-related change in *mdx* mice ²³; finally, our experiments were carried 332 333 out in DBA/2J-mdx mice, using theta-patterned burst (100 Hz) stimulation. Although experiments were carried out in three different DMD mouse models (mdx, mdx^{3cv} & DBA/2J-334 335 mdx) which differ in the severity of the dystrophic pathology, it is currently not possible to 336 conclude whether this influences synaptic plasticity in the brain, because of the use of different 337 protocols by different groups to induce LTP and the fact that dystrophin isoform expression 338 differs between mdx, mdx^{3cv} and DBA/2J-mdx mice. Given that the muscular pathology of 339 DMD patients is much more severe than that of the *mdx* mouse, and that muscle-secreted 340 and physical activity-induced factors have an influence on brain neurogenesis and synaptic plasticity ⁴⁰ it would be of interest to determine whether severity of muscle pathology affects 341 342 synaptic plasticity in DMD mouse models. Our data, obtained from the DBA/2J-mdx mouse 343 model, which displays significant muscle atrophy and fibrosis, show no deficit in hippocampal 344 LTP (Figure 1).

345 Previous work reported that the lack of dystrophin leads to decreased GABAergic 346 neurotransmission ^{41,42}, due to decreased GABA-A clustering on post-synaptic densities ^{7,43,44}. 347 Reduced GABAergic neurotransmission has been proposed to be the possible mechanism underlying the increased prevalence of epilepsy in boys with DMD ^{45,46} and enhanced 348 hippocampal LTP in DMD mouse models ^{9,47}. In addition, the Dp71 isoform-dependent 349 proposed as 350 dysregulation of the potassium Kir4.1 channel has been an 351 alternative/complementary mechanism of action responsible for epilepsy in DMD patients.

However, it is known that altered synaptic function and epileptiform hypersynchronous network activity also rely on the presence of single cell membrane hyperexcitability ^{27,48-52}. As we did not observe any change in basal synaptic transmission, short- and long-term synaptic 355 plasticity in DBA/2J-mdx mice, we decided to further investigate the effects of lack of 356 dystrophin on the intrinsic physiological properties of the CA1-PC's plasma membrane, as a possible neuronal correlate of DMD-associated brain dysfunction. While electrotonic and firing 357 358 properties were unchanged between genotypes, the after-hyperpolarization was significantly 359 bigger in DBA/2J-mdx mice. mAHP is mediated by voltage-gated K⁺ channels (VGKC) ⁵³. To 360 assess the possible role of VGKC for the alteration of mAHP in mdx mice, we performed 361 outside-out nucleated macropatch recordings to assess the biophysical properties of outward voltage-gated currents, as previously reported ^{27,54}. We did not observe any significant effect 362 363 of genotype on such current. However, a more detailed investigation of DBA/2J-mdx-364 dependent alterations of biophysical properties is due, as in our native system we could not 365 tease apart the role of each conductance responsible for voltage-gated K⁺ currents. In fact, 366 dystrophin deficiency has been linked to an array of changes in membrane K⁺ conductances 367 in cardiomyocytes, mediated by different signalling mechanisms. For example, inward rectifier K⁺ (K_{ir}) current densities 2.1 were decreased in conditions of dystrophin deficiency ⁵⁵ but no 368 369 changes in the protein channel levels were observed, suggesting a change in trafficking rather 370 than expression. In fact, K_{ir} 2.1 is functionally and structurally connected to dystrophin via syntrophin ⁵⁶ so it has been hypothesized that dystrophin deficiency may result in reduced 371 372 presence of this conductance on the plasma membrane. However, Ky 2.1 and 1.5 currents, 373 mediating the slow inactivating component of I_k , are also impaired in cardiomyocytes; they are 374 connected to dystrophin via the F-actin and α -actinin-2 network. For this reason, we 375 hypothesize that, in neurons, the lack of dystrophin may result in increased mAHP via 376 syntrophin mediated altered docking of M-current-generating voltage-gated conductances, such as $K_v 7$. In fact, $K_v 7.x$ channels have been reported to mediate the mAHP ³⁶. 377

While previous work has been carried out on the intrinsic excitability of cerebellar Purkinje cells, showing reduced excitability due to hyperpolarized RMP in dystrophin-deficient mice ⁵⁷, such measures have not been performed before in the hippocampus.

These observations reinforce the idea that dystrophin may play a key role in brain physiology,
 regulating both motor and cognitive functions. Several lines of evidence show that dystrophin

383 expression is altered in non DMD patients: for example, dystrophin expression is reduced in people with different forms of epilepsy, such as temporal mesial lobe epilepsy ^{58,59}, sclerotic 384 hippocampus ⁶⁰ and in focal cortical dysplasia ⁶¹. However, while the same phenotype has 385 386 been observed in the piriform cortex of pilocarpine-induced status epilepticus rats rodent models of epilepsy 62,63, no changes in dystrophin expression have been found in the 387 388 hippocampus of these animals. On the other hand, dystrophin almost disappeared from the 389 granule cells of the dentate gyrus, following the induction of seizures in mice treated with kainic 390 acid ⁴⁶. Such diversity of experimental outcomes somehow reflects the discrepancies 391 observed in the effects of dystrophin deficiency on excitatory synaptic function and it probably 392 reveals the generalized lack of knowledge about the role of this protein on brain physiology.

393 Our results showed increased mAHP in CA1-PCs: this observation may have consequences 394 on network function and cognitive decline observed in people with dystrophin deficiency. In 395 fact, the mAHP corresponds to the refractory period following the AP and has a key role in determining the ability of a single neuron to sustainably fire ³⁶. An increased mAHP amplitude 396 397 may result in reduced excitability of the neural network. This, in turn, may account for the 398 cognitive impairment paralleling dystrophin deficiency and be an adaptive mechanism due to 399 increased network excitability. However, the consequences of the increased mAHP in 400 dystrophin deficient mice, both at network and behavioural level, require a more detailed 401 investigation.

402 It should also be highlighted that while our investigation provides important insights into the 403 alterations of hippocampal neuronal function associated with the lack of Dp427 isoforms, 404 cognitive deficits are more significant in DMD patients which lack expression of shorter 405 dystrophin isoforms derived from downstream promoters on the dystrophin gene, in addition 406 to the full length Dp427 isoform ⁶⁴. Whether this is because of a specific role for the shorter 407 dystrophin isoforms or whether this is simply due to the further reduction of the presence of 408 C-terminal dystrophin protein domains in the cell is unknown. However, it is currently not 409 possible to study the role of the shorter dystrophin isoforms in isolation due to the mechanism

410 by which these different isoforms are generated, but it is possible to isolate the role of the411 Dp427 isoform.

For this reason, we should be cautious in translating these data to all DMD patients. Nonetheless, we believe the results presented here provide insight into the general role of Dp427 in hippocampal function and suggest that the severe muscle dysfunction observed in these mice does not affect most of the CA1 PCs membrane properties.

416 In addition, the analysis of inhibitory interneurons' intrinsic properties in dystrophin deficient 417 mice is required to better clarify the single cell alterations, which may mechanistically explain 418 the effects of dystrophin deficiency on brain function. Finally, clarifying the role of dystrophin 419 in cognition and the pathological effects of its deficiency in brain function may provide evidence 420 of common pathogenic mechanisms between different disorders resulting in cognitive deficit, 421 including DMD, Alzheimer's disease and epilepsy. This may eventually inform the 422 development of a unified model for the onset different brain diseases resulting in impaired 423 cognition.

424

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429 Author contributions

RB: extracellular recordings and relative data analysis; WE: experimental design, manuscript
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experimental design and manuscript revision; FT: scientific direction, intracellular recordings,
data analysis, experimental design and manuscript preparation.

435	Competing interests statement					
436	No competing interest					
437	Data availability					
438	The da	atasets generated during and/or analysed during the current study are available from				
439	the co	responding author on reasonable request.				
440 441		References				
442	1	Hoffman, E. P., Brown, R. H., Jr. & Kunkel, L. M. Dystrophin: the protein product of the Duchenne muscular ductrophy locus. <i>Call</i> 51 , 010, 028 (1087).				
445 444 445	2	Chamberlain, J. S. <i>et al.</i> Expression of the murine Duchenne muscular dystrophy gene				
445 446 447	3	Lidov, H. G., Byers, T. J. & Kunkel, L. M. The distribution of dystrophin in the murine central nervous system: an immunocytochemical study. <i>Neuroscience</i> 54 , 167-				
448 449	1	187 (1993). D'Amario D. <i>et al.</i> Dystrophin Cardiomyopathies: Clinical Management. Molecular				
450 451	-	Pathogenesis and Evolution towards Precision Medicine. <i>Journal of clinical medicine</i> 7 doi:10.3390/icm7090291 (2018)				
452 453	5	Yiu, E. M. & Kornberg, A. J. Duchenne muscular dystrophy. <i>Journal of paediatrics</i> and child health 51 , 759-764, doi:10.1111/jpc.12868 (2015)				
454 455	6	Duchenne, G. B. A. Reserches sur la paralysie musculaire pseudeohyper-trofique, ou paralysie myo-sclerosique. <i>Arch. Gen. Med.</i> 11 , 5–25, 179–209,305–121, 421–143,				
456 457	7	552–188. (1868). Anderson, J. L., Head, S. I., Rae, C. & Morley, J. W. Brain function in Duchenne				
458 459	8	muscular dystrophy. <i>Brain : a journal of neurology</i> 125 , 4-13 (2002). Cyrulnik, S. E. & Hinton, V. J. Duchenne muscular dystrophy: a cerebellar disorder?				
460 461 462	9	<i>Neurosci Biobehav Rev</i> 32 , 486-496, doi:10.1016/j.neubiorev.2007.09.001 (2008). Vaillend, C., Billard, J. M. & Laroche, S. Impaired long-term spatial and recognition				
462	10	mouse. <i>Neurobiol Dis</i> 17 , 10-20, doi:10.1016/j.nbd.2004.05.004 (2004).				
464 465 466	10	expression provide insights into the cognitive phenotype of Duchenne muscular dystrophy. <i>Scientific reports</i> 7 , 12575-12575, doi:10.1038/s41598-017-12981-5				
467 468	11	(2017). Lidov, H. G., Selig, S. & Kunkel, L. M. Dp140: a novel 140 kDa CNS transcript from				
469 470	12	the dystrophin locus. <i>Hum Mol Genet</i> 4 , 329-335 (1995). de Brouwer, A. P. <i>et al.</i> A 3-base pair deletion, c.9711 9713del, in DMD results in				
471 472		intellectual disability without muscular dystrophy. <i>European journal of human</i> genetics : <i>EJHG</i> 22 , 480-485, doi:10.1038/ejhg.2013.169 (2014).				
473 474	13	Kim, T. W., Wu, K., Xu, J. L. & Black, I. B. Detection of dystrophin in the postsynaptic density of rat brain and deficiency in a mouse model of Duchenne				
475 476 477	14	muscular dystrophy. <i>Proc Natl Acad Sci U S A</i> 89 , 11642-11644 (1992). Bliss, T. V. & Collingridge, G. L. A synaptic model of memory: long-term potentiation in the hippocampus. <i>Nature</i> 361 , 31-39, doi:10.1038/361031a0 (1993).				

- Waite, A., Tinsley, C. L., Locke, M. & Blake, D. J. The neurobiology of the
 dystrophin-associated glycoprotein complex. *Annals of medicine* 41, 344-359,
 doi:10.1080/07853890802668522 (2009).
- 481 16 Anand, A. *et al.* Dystrophin induced cognitive impairment: mechanisms, models and
 482 therapeutic strategies. *Annals of neurosciences* 22, 108-118,
 483 doi:10.5214/ans.0972.7531.221210 (2015).
- de Brouwer, A. P. M. *et al.* A 3-base pair deletion, c.9711_9713del, in DMD results
 in intellectual disability without muscular dystrophy. *European journal of human genetics : EJHG* 22, 480-485, doi:10.1038/ejhg.2013.169 (2014).
- 487 18 Goodwin, F., Muntoni, F. & Dubowitz, V. Epilepsy in Duchenne and Becker
 488 muscular dystrophies. *Eur J Paediatr Neurol* 1, 115-119 (1997).
- 489 19 Etemadifar, M. & Molaei, S. Epilepsy in Boys with Duchenne Muscular Dystrophy
 490 Journal of Research in Medical Sciences 3, 116-119 (2004).
- 491 20 Pane, M. *et al.* Duchenne muscular dystrophy and epilepsy. *Neuromuscul Disord* 23, 313-315, doi:10.1016/j.nmd.2013.01.011 (2013).
- Hendriksen, R. G. *et al.* A possible role of dystrophin in neuronal excitability: a
 review of the current literature. *Neurosci Biobehav Rev* 51, 255-262,
 doi:10.1016/j.neubiorev.2015.01.023 (2015).
- 496 22 Vaillend, C. *et al.* Spatial discrimination learning and CA1 hippocampal synaptic
 497 plasticity in mdx and mdx3cv mice lacking dystrophin gene products. *Neuroscience*498 86, 53-66 (1998).
- 499 23 Sesay, A. K., Errington, M. L., Levita, L. & Bliss, T. V. Spatial learning and
 hippocampal long-term potentiation are not impaired in mdx mice. *Neuroscience letters* 211, 207-210 (1996).
- 50224Lewon, M. *et al.* Evaluation of the behavioral characteristics of the mdx mouse model503of duchenne muscular dystrophy through operant conditioning procedures.504505
- 504 Behavioural processes **142**, 8-20, doi:10.1016/j.beproc.2017.05.012 (2017).
- Wikiera, B., Jakubiak, A., Zimowski, J., Noczynska, A. & Smigiel, R. Complex
 glycerol kinase deficiency X-linked contiguous gene syndrome involving congenital
 adrenal hypoplasia, glycerol kinase deficiency, muscular Duchenne dystrophy and
 intellectual disability (IL1RAPL gene deletion). *Pediatric endocrinology, diabetes, and metabolism* 18, 153-157 (2012).
- Vazifehkhah Ghaffari, B., Kouhnavard, M., Aihara, T. & Kitajima, T. Mathematical
 modeling of subthreshold resonant properties in pyloric dilator neurons. *BioMed research international* 2015, 135787, doi:10.1155/2015/135787 (2015).
- 513 27 Brown, J. T., Chin, J., Leiser, S. C., Pangalos, M. N. & Randall, A. D. Altered
 514 intrinsic neuronal excitability and reduced Na+ currents in a mouse model of
 515 Alzheimer's disease. *Neurobiology of aging* 32, 2109 e2101-2114,
 516 doi:10.1016/j.neurobiolaging.2011.05.025 (2011).
- Ranck, J. B., Jr. Studies on single neurons in dorsal hippocampal formation and
 septum in unrestrained rats. I. Behavioral correlates and firing repertoires. *Experimental neurology* 41, 461-531 (1973).
- 520 29 Otto, T., Eichenbaum, H., Wiener, S. I. & Wible, C. G. Learning-related patterns of
 521 CA1 spike trains parallel stimulation parameters optimal for inducing hippocampal
 522 long-term potentiation. *Hippocampus* 1, 181-192, doi:10.1002/hipo.450010206
 523 (1991).
- 52430Hill, A. J. First occurrence of hippocampal spatial firing in a new environment.525*Experimental neurology* 62, 282-297, doi:https://doi.org/10.1016/0014-4886(78)90058-4 (1978).

- 527 31 Grover, L. M. & Teyler, T. J. Two components of long-term potentiation induced by
 528 different patterns of afferent activation. *Nature* 347, 477-479, doi:10.1038/347477a0
 529 (1990).
- Grover, L. M. & Teyler, T. J. Activation of NMDA receptors in hippocampal area
 CA1 by low and high frequency orthodromic stimulation and their contribution to
 induction of long-term potentiation. *Synapse (New York, N.Y.)* 16, 66-75,
 doi:10.1002/syn.890160108 (1994).
- 33 O'Connor, J. J., Rowan, M. J. & Anwyl, R. Long-lasting enhancement of NMDA
 receptor-mediated synaptic transmission by metabotropic glutamate receptor
 activation. *Nature* 367, 557-559, doi:10.1038/367557a0 (1994).
- 537 34 O'Leary, D. M. & O'Connor, J. J. Potentiation of synaptic transmission by (S)-3,5538 dihydroxy phenylglycine in the rat dentate gyrus in vitro: a role for voltage dependent
 539 calcium channels and protein kinase C. *Progress in neuro-psychopharmacology & biological psychiatry* 23, 133-147 (1999).
- 541
 35
 Comim, C. M. *et al.* Neurocognitive Impairment in mdx Mice. *Molecular*

 542
 Neurobiology 56, 7608-7616, doi:10.1007/s12035-019-1573-7 (2019).
- Gu, N., Vervaeke, K., Hu, H. & Storm, J. F. Kv7/KCNQ/M and HCN/h, but not
 KCa2/SK channels, contribute to the somatic medium after-hyperpolarization and
 excitability control in CA1 hippocampal pyramidal cells. *J Physiol* 566, 689-715,
 doi:10.1113/jphysiol.2005.086835 (2005).
- 54737Coley, W. D. *et al.* Effect of genetic background on the dystrophic phenotype in mdx548mice. *Hum Mol Genet* 25, 130-145, doi:10.1093/hmg/ddv460 (2016).
- 549 38 Hakim, C. H. *et al.* A Five-Repeat Micro-Dystrophin Gene Ameliorated Dystrophic
 550 Phenotype in the Severe DBA/2J-mdx Model of Duchenne Muscular Dystrophy. *Mol*551 *Ther Methods Clin Dev* 6, 216-230, doi:10.1016/j.omtm.2017.06.006 (2017).
- 55239Coley, W. D. *et al.* Effect of genetic background on the dystrophic phenotype in mdx553mice. *Hum Mol Genet* 25, 130-145, doi:10.1093/hmg/ddv460 (2016).
- 55440Lourenco, M. V. *et al.* Exercise-linked FNDC5/irisin rescues synaptic plasticity and555memory defects in Alzheimer's models. *Nat Med* 25, 165-175, doi:10.1038/s41591-556018-0275-4 (2019).
- Kueh, S. L., Head, S. I. & Morley, J. W. GABA(A) receptor expression and inhibitory
 post-synaptic currents in cerebellar Purkinje cells in dystrophin-deficient mdx mice. *Clin Exp Pharmacol Physiol* 35, 207-210, doi:10.1111/j.1440-1681.2007.04816.x
 (2008).
- 56142Nusser, Z., Cull-Candy, S. & Farrant, M. Differences in synaptic GABA(A) receptor562number underlie variation in GABA mini amplitude. Neuron 19, 697-709 (1997).
- Vaillend, C. *et al.* Rescue of a dystrophin-like protein by exon skipping in vivo
 restores GABAA-receptor clustering in the hippocampus of the mdx mouse. *Mol Ther*18, 1683-1688, doi:10.1038/mt.2010.134 (2010).
- Waite, A., Brown, S. C. & Blake, D. J. The dystrophin-glycoprotein complex in brain
 development and disease. *Trends Neurosci* 35, 487-496,
 doi:10.1016/j.tins.2012.04.004 (2012).
- 569 45 Nakao, K., Kito, S., Muro, T., Tomonaga, M. & Mozai, T. Nervous system
 570 involvement in progressive muscular dystrophy. *Proc Aust Assoc Neurol* 5, 557-564
 571 (1968).
- Knuesel, I., Zuellig, R. A., Schaub, M. C. & Fritschy, J. M. Alterations in dystrophin
 and utrophin expression parallel the reorganization of GABAergic synapses in a
 mouse model of temporal lobe epilepsy. *Eur J Neurosci* 13, 1113-1124 (2001).
- 575 47 Vaillend, C., Ungerer, A. & Billard, J. M. Facilitated NMDA receptor-mediated
 576 synaptic plasticity in the hippocampal CA1 area of dystrophin-deficient mice.

577 578		<i>Synapse</i> 33 , 59-70, doi:10.1002/(sici)1098-2396(199907)33:1<59::aid-syn6>3.0.co;2-k (1999).
579	48	Palop, J. J. et al. Aberrant excitatory neuronal activity and compensatory remodeling
580		of inhibitory hippocampal circuits in mouse models of Alzheimer's disease. <i>Neuron</i>
581		55 , 697-711, doi:10.1016/j.neuron.2007.07.025 (2007).
582	49	Tamagnini, F. et al. Altered intrinsic excitability of hippocampal CA1 pyramidal
583		neurons in aged PDAPP mice. Frontiers in cellular neuroscience 9, 372,
584		doi:10.3389/fncel.2015.00372 (2015).
585	50	Tamagnini, F., Scullion, S., Brown, J. T. & Randall, A. D. Intrinsic excitability
586		changes induced by acute treatment of hippocampal CA1 pyramidal neurons with
587		exogenous amyloid beta peptide. <i>Hippocampus</i> 25, 786-797, doi:10.1002/hipo.22403
588		(2015).
589	51	Kerrigan, T. L. & Randall, A. D. A new player in the "synaptopathy" of Alzheimer's
590		disease - arc/arg 3.1. Front Neurol 4, 9, doi:10.3389/fneur.2013.00009 (2013).
591	52	Randall, A. D., Witton, J., Booth, C., Hynes-Allen, A. & Brown, J. T. The functional
592		neurophysiology of the amyloid precursor protein (APP) processing pathway.
593		<i>Neuropharmacology</i> 59 , 243-267, doi:10.1016/j.neuropharm.2010.02.011 (2010).
594	53	Gu, N., Vervaeke, K., Hu, H. & Storm, J. F. Kv7/KCNQ/M and HCN/h, but not
595		KCa2/SK channels, contribute to the somatic medium after-hyperpolarization and
596		excitability control in CA1 hippocampal pyramidal cells. <i>The Journal of physiology</i>
597		566 , 689-715, doi:10.1113/jphysiol.2005.086835 (2005).
598	54	Tamagnini, F. <i>et al.</i> Hippocampal neurophysiology is modified by a disease-
599		associated C-terminal fragment of tau protein. <i>Neurobiology of aging</i> 60 , 44-56,
600		doi:10.1016/j.neurobiolaging.2017.07.005 (2017).
601	55	Rubi, L., Koenig, X., Kubista, H., Todt, H. & Hilber, K. Decreased inward rectifier
602		potassium current IK1 in dystrophin-deficient ventricular cardiomyocytes. <i>Channels</i>
603		(Austin, Tex.) 11, 101-108, doi:10.1080/19336950.2016.1228498 (2017).
604	56	Willis, B. C., Ponce-Balbuena, D. & Jalife, J. Protein assemblies of sodium and
605		inward rectifier potassium channels control cardiac excitability and arrhythmogenesis.
606		American journal of physiology. Heart and circulatory physiology 308 , H1463-1473,
607		doi:10.1152/ajpheart.00176.2015 (2015).
608	57	Snow, W. M., Anderson, J. E. & Fry, M. Regional and genotypic differences in
609		intrinsic electrophysiological properties of cerebellar Purkinje neurons from wild-type
610		and dystrophin-deficient mdx mice. Neurobiol Learn Mem 107, 19-31,
611		doi:10.1016/j.nlm.2013.10.017 (2014).
612	58	Lee, T. S. et al. Aquaporin-4 is increased in the sclerotic hippocampus in human
613		temporal lobe epilepsy. Acta Neuropathol 108, 493-502, doi:10.1007/s00401-004-
614		0910-7 (2004).
615	59	Das, A. et al. Hippocampal tissue of patients with refractory temporal lobe epilepsy is
616		associated with astrocyte activation, inflammation, and altered expression of channels
617		and receptors. <i>Neuroscience</i> 220 , 237-246, doi:10.1016/j.neuroscience.2012.06.002
618		(2012).
619	60	Eid, T. et al. Loss of perivascular aquaporin 4 may underlie deficient water and K+
620		homeostasis in the human epileptogenic hippocampus. Proc Natl Acad Sci USA 102,
621		1193-1198, doi:10.1073/pnas.0409308102 (2005).
622	61	Medici, V., Frassoni, C., Tassi, L., Spreafico, R. & Garbelli, R. Aquaporin 4
623		expression in control and epileptic human cerebral cortex. Brain Res 1367, 330-339,
624		doi:10.1016/j.brainres.2010.10.005 (2011).

- 62 Kim, J. E. *et al.* Astroglial loss and edema formation in the rat piriform cortex and
 hippocampus following pilocarpine-induced status epilepticus. *J Comp Neurol* 518,
 627 4612-4628, doi:10.1002/cne.22482 (2010).
- 63 Sheen, S. H. *et al.* Decrease in dystrophin expression prior to disruption of brainblood barrier within the rat piriform cortex following status epilepticus. *Brain Res*630 **1369**, 173-183, doi:10.1016/j.brainres.2010.10.080 (2011).
- 631 64 Taylor, P. J. *et al.* Dystrophin gene mutation location and the risk of cognitive
 632 impairment in Duchenne muscular dystrophy. *PLoS One* 5, e8803,
 633 driv10.1271 (impact and prove 0008202 (2010))
- 633 doi:10.1371/journal.pone.0008803 (2010).
- 634 635