MECP2e1 isoform mutation affects the form and function of neurons derived from Rett syndrome patient iPS cells

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**Abstract**

MECP2 mutations cause the X-linked neurodevelopmental disorder Rett Syndrome (RTT) by consistently altering the protein encoded by the MECP2e1 alternative transcript. While mutations that simultaneously affect both MECP2e1 and MECP2e2 isoforms have been widely studied, the consequence of MECP2e1 deficiency on human neurons remains unknown. Here we report the first isoform-specific patient induced pluripotent stem cell (iPSC) model of RTT. RTT patient iPSC cell-derived neurons retain an inactive X-chromosome and express only the mutant allele. Single-cell mRNA analysis demonstrated they have a molecular signature of cortical neurons. Mutant neurons exhibited a decrease in soma size, reduced dendritic complexity and decreased cell capacitance, consistent with impaired neuronal maturation. The soma size phenotype was rescued cell-autonomously by MECP2e1 transduction in a level-dependent manner but not by MECP2e2 gene transfer. Importantly, MECP2e1 mutant neurons showed a dysfunction in action potential generation, voltage-gated Na+ currents, and miniature excitatory synaptic current frequency and amplitude. We conclude that MECP2e1 mutation affects soma size, information encoding properties and synaptic connectivity in human neurons that are defective in RTT.

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**Introduction**

Rett Syndrome (RTT) is a neurodevelopmental disorder [OMIM12750] characterized by repetitive hand motions and loss of acquired language (Chahrour and Zoghbi, 2007). Heterozygous loss-of-function mutation in the X-linked gene encoding Methyl-CpG Binding Protein 2 (MECP2) is the prime cause of RTT in girls (Amir et al., 1999). This gene is alternatively spliced into MECP2e1 and MECP2e2 isoforms that encode distinct proteins differing at the N-termini due to exclusion or inclusion of exon 2 respectively (Kriaucionis and Bird, 2004; Mnatzakanian et al., 2004). Mutations that affect both isoforms have been widely studied, and the role of MECP2 in binding methylated and hydroxy-methylated cytosine genome-wide (Mellen et al., 2012; Skene et al., 2010) to recruit chromatin-remodelling proteins that modulate global transcription is now well established (Chahrour et al., 2008; Lyst et al., 2013). TALEN-mediated mutagenesis of the MECP2 locus demonstrated that MECP2 ablation results in global decreases in gene transcription and translation in a human ES cell-based model of RTT (Li et al., 2013). These defects were manifest in abnormal neuronal morphology and function, including impaired mitochondrial function.

The majority of RTT patient mutations affect both isoforms but identification of individuals with a MECP2e1-specific mutation that does not alter MECP2e2 indicates that MECP2e1 isoform dysfunction is sufficient to cause RTT (Mnatzakanian et al., 2004). A recent report of a Mecp2e1-specific mouse model of RTT further suggests that this MECP2 isoform is responsible for RTT-related behavioural abnormalities (Yasu et al., 2014), but the effect of MECP2e1 deficiency on human neurons has not been evaluated. In contrast, no MECP2e2-specific mutations have been described in RTT patients, and Mecp2e2-mutant mouse lacks neurological phenotypes and instead exhibit placental defects (Itoh et al., 2012). However, expression of either Mecp2e1 or Mecp2e2 can improve a subset of RTT-related behavioural phenotypes in Mecp2-null mice (Kerr et al., 2012). These findings suggest that endogenous MECP2e1 is essential for normal brain function, but Mecp2e2 can ameliorate certain disease features in mouse models of RTT.
We, and others, reported the generation of human and mouse induced pluripotent stem cells (hiPSCs and miPSCs, respectively) from RTT patients and mouse models that carry pathogenic mutations in both MECP2 isoforms (Ananiev et al., 2011; Cheung et al., 2011; Kim et al., 2011). RTT iPSC-derived neurons exhibit maturation and electrophysiological defects reminiscent of those seen in RTT patients and mouse models (Farra et al., 2012) and are amenable to rescue by introduction of exogenous MECP2 or drugs such as IGF1 (Li et al., 2013; Marchetto et al., 2010). Generally, female RTT-hiPSCs retain an inactive X-chromosome (Xi) (Pomp et al., 2011; Tchieu et al., 2010). Here, we generated hiPSC-derived neurons that express mutant MECP2e1. Using this system, we find that MECP2e1 mutation affects the soma size and electrophysiological properties of human neurons.

Materials and methods

Cell culture

RTTe1-fibroblasts were obtained from Dr. Patrick Macleod at the Victoria General Hospital, Victoria, BC, Canada, and cultured under the approval of the SickKids Research Ethics Board and Canadian Institutes of Health Research Stem Cell Oversight Committee. Fibroblasts were maintained in fibroblast medium: Dulbecco’s Modified Eagle Medium (DMEM) containing 10% Fetal Bovine Serum, and 100× penicillin and streptomycin (all from Invitrogen). RTTe1-hiPSCs were generated from fibroblasts and maintained in hiPSC medium as previously described (Hotta et al., 2009).

Androgen receptor assay

To identify the methylated Xi, 200 ng of DNA was digested overnight at 37 °C with methylation-sensitive enzymes HpaII and HhaI (Invitrogen). To discriminate between the two parental X-chromosomes, 20 ng of digested and undigested DNA was amplified with primers spanning the heterozygous polymorphic trinucleotide repeat in the first exon of the AR gene for 32 cycles. The 5’ end of the forward primer was labelled with FAM fluorescein (Invitrogen). PCR products were separated on an ABI3100 Genetic Analyzer with 500 LIZ size standard and analysed by Peak Scanner software (all from Applied Biosystems). XCI ratio (Table S1) was calculated as previously described (Cheung et al., 2011).

In vitro and in vivo differentiation

For in vitro differentiation, hiPSCs were detached and grown in suspension in hiPSC medium (Hotta et al., 2009) without FGF2 for eight days to form embryoid bodies. Embryoid bodies were adhered and allowed to further differentiate for eight days. Differentiated derivatives were analysed via immunocytochemistry with appropriate antibodies (Table S4). For in vivo differentiation, one 10 cm dish of hiPSCs was detached and suspended in a mixture of KNOCKOUT DMEM (Invitrogen), Matrigel (BD Biosciences), Collagen (STEMCELL Technologies) (ratio 2:1:2), and 10 μM ROCK Inhibitor (Sigma) and injected intramuscularly into immunodeficient mice. Fixed tumours were embedded in paraffin, sectioned, and stained with haematoxylin and eosin for pathological analysis. All procedures using animals were approved by the SickKids Animal Care Committee under the auspices of The Canadian Council on Animal Care, and conducted with the approval of the Canadian Institutes of Health Research Stem Cell Oversight Committee.

Immunocytochemistry

Cells were fixed with 4% formaldehyde (EMD Biosciences) for 10 min at room temperature (RT), permeabilized with 0.1% Nonidet P-40 (Sigma). Blocking was performed for 3 h at RT, primary antibodies diluted in block solution and incubated overnight at 4 °C (See Table S4 for antibodies used). Images were captured using a Leica DMI4000B microscope equipped with Leica DFC340FX camera and Leica Application Suite software for hiPSCs or Zeiss Axiovert 200M microscope equipped with a Hamamatsu C9100-13 EMCCD camera and Improvion Velocity software for neurons. Soma size of neurons was scored using Improvion Velocity software on 40× images blinded to the observer.

Neuronal differentiation

The Brennand protocol with slight modifications (Brennand et al., 2011; Chambers et al., 2009) was used for neuronal differentiation of the RTTe1#27 line (Table S2). The Kim protocol with slight modifications (Kim et al., 2012) or the Brennand protocol with the addition of DAPT in the medium was used for the remaining lines (Table S2). Analysis was compared to WT-neurons generated with the same protocol. Soma-size measurements were performed as described (Cheung et al., 2011). See supplemental material for more details.

Generation and transduction of lentivirus

MECP2 isoform-specific lentiviruses were previously reported (Rastegar et al., 2009). The generation and transduction of lentivirus were performed as previously described (Hotta et al., 2009). In brief, plasmids containing cDNA of interest were transfected into 293T cells using Lipofectamine 2000 (Invitrogen). The supernatant containing the virus was collected two days post-transfection and concentrated by ultracentrifugation. 293T cells were maintained in Dulbecco’s Modified Eagle Medium (DMEM) containing 10% Fetal Bovine Serum, and 100× penicillin and streptomycin, MEM NEAA (all from Invitrogen).

Titration of lentivirus was performed as previously described (Hotta et al., 2009). In brief, 293T cells were transduced with EF1α-EGFP lentivirus. Titer (infectious units, IU) of lentivirus was calculated with the following formula: Viral titer (IU ml⁻¹) = [Infected cell number] × [EGFP %/100]% [Amount of virus used (ml)]. Transduction of NPCs was performed in the presence of 0.6 μg ml⁻¹ Hexadimethrine bromide (Sigma) for six hours with a multiplicity of infection (MOI) of one calculated for 293T cells.

RNA isolation, RT-PCR and cDNA sequencing

RNA was isolated using Trizol extraction method (Invitrogen) and SSII RT (Invitrogen) was used for the reverse transcription following manufacturer’s instructions. Primers for the Real-Time PCR assays were designed using Primer3 online primer design software and qPCR was carried out using SYBR green (Applied Biosystems) on an ABI 7900HT PCR System (Applied Biosystems). To sequence RTTe1-hiPSC cDNA, RT-PCR was performed using V1 primers (Table S3). The MECP2e1 amplicon (382 bp or 371 bp for WT or mutant allele, respectively) was gel purified from the MECP2e2 amplicon (506 bp) using QIAquick PCR Purification Kit following manufacturer’s instructions and sequenced using the V1-f for amplification.

Bisulfite sequencing

Bisulfite conversion was performed as previously described (Fussner et al., 2011). Briefly 1 μg of DNA was subjected to conversion using the DNA Methylation Gold Kit (Zymo Research). 50 ng of converted DNA was subjected to PCR with appropriate primers (Table S3).

Single-cell fluidigm array

Neurons were subjected to single cell sorting and Fluidigm analysis as described (Pasca et al., 2011). Single cell sorting was performed by The Flow Cytometry Facility at The Hospital for Sick Children (Toronto, Ontario, Canada) using MoFlo BRU cell sorter (Beckman Coulter). Cells
were stained by propidium iodide and live cells were sorted into CellsDirect amplification master mix (Invitrogen) into 96-well plates. RT reaction and 15 rounds of cDNA amplification were performed using primers to genes of interest (Table S3). Fluidigm Biomark qPCR assays were performed by Janet Rossant lab at The Hospital for Sick Children (Toronto, Ontario, Canada). Data analyses and generation of figures were performed using R-Statistical analysis software (http://cran.r-project.org). For neuronal identity quantifications, cells were deemed neurons if expressing NCAM or MAP2 with: FOXP1 and ETV1 double positive (lower layer); FOXP1-/ETV1- /CUX1+/SATB2+/Ctip2+/REELIN+ (upper layer); CAMK2+, VGLUT1+, VGLUT2+ or VGLUT3+ (glutamatergic); GAD67+ or VGAT+ (GABAergic) (Pasca et al., 2011).

Western blot analysis

Nuclear proteins were collected using nuclear extraction protocol, samples were aliquoted and snap frozen in liquid nitrogen for storage until western blots were performed. 5 μg of total protein was loaded for western blots, transferred to nitrocellulose membranes overnight at 4 °C Membranes were blocked in 5% Milk PBS-T and incubated

![Western Blot Analysis](image)

**Fig. 1.** Generation and pluripotency characterization of RTTe1-hiPSCs. (A) Androgen receptor (AR) methylation screen of RTTe1-hiPSCs demonstrates skewed XCI ratios in all examined lines. Bar graph depicts XCI ratio of two X chromosomes (X1 and X2). Fib., fibroblasts. (B) qRT-PCR analyses of pMXs reprogramming retroviral transgenes. (C) qRT-PCR of endogenous pluripotency genes in RTTe1-hiPSC lines. Data are expressed as mean ± SEM. (D) Bisulfite sequencing of retroviral pMXs-LTR reprogramming vectors in RTTe1-hiPSC lines. Open and closed Cpg sites indicate unmethylated and methylated Cpg sites, respectively. (E) RTTe1-hiPSCs differentiate into the three germ layers in vitro and in vivo (representative images of RTTe1-hiPSC#39 teratomas). Scale bars, 50 μm (immunocytochemistry) and 100 μm (histology).
overnight in primary antibodies (see Table S4), washed 5 times in PBS-T and incubated in appropriate HRP-conjugated secondary antibodies (Invitrogen) for 1 h at room temperature. Densitometry measurements and normalization (to Histone H3 signal) was performed using ImageJ software.

**Karyotyping**

Standard G-banding chromosome analysis with a 400–500 banding resolution was performed at The Centre of Applied Genomics (TCAG [The Hospital for Sick Children, Toronto, Canada]).

**Electrophysiology**

Electrophysiology was performed as described (Farra et al., 2012). See supplemental information for detailed protocol.

**Results**

**Isolation of mutant RTTe1-hiPSCs through XCI**

Fibroblasts acquired from an RTT patient with an 11 base-pair (bp) deletion in exon 1 of MECP2 (RTTe1) were used to generate RTTe1-hiPSCs via retroviral transduction of human OCT4, SOX2, KLF4, and c-MYC. This exon 1 specific NM_00110792.1:c.47_57del; p. (Gly16Glufs*22) mutation causes a frameshift resulting in severely truncated MECP2e1 but intact MECP2e2 (Mnatzakanian et al., 2004). We used the Androgen Receptor (AR) assay to determine the X-inactivation status in RTTe1-hiPSCs, revealing that all of the 22 RTTe1-hiPSC lines exhibited XCI skewing (80:20—100:0) towards the same parental X-chromosome (Fig. 1A, Table S1). We selected four RTTe1-hiPSC lines for detailed pluripotency characterization (Table S2). All four hiPSC lines silenced and methylated the reprogramming retroviral transgenes, activated the pluripotency-related genes (Figs. 1B–D) and had normal karyotypes (Fig. S1). Embryoid body-mediated spontaneous in vitro and teratoma-based in vivo differentiation assays demonstrate that RTTe1-hiPSC lines give rise to cells of all three germ layers (Fig. 1E). Collectively these data demonstrate that the generated hiPSC lines are of high quality appropriate for RTT disease modelling. To determine the identity of the active X-chromosome, we performed MECP2 cDNA sequencing with primers spanning the 11 bp deletion revealing that the RTTe1-hiPSCs solely expressed the mutant allele (Fig. 2A). To ensure that neither a change in the Xi nor Xi erosion (Mekhoubad et al., 2012) occurred during differentiation, we confirmed persistent expression of the mutant MECP2e1-containing X-chromosome using cDNA sequencing and the AR assay on hiPSC-derived neurons (Fig. 2B) (see below for neuronal characterization).

**Directed differentiation into cortical neurons expressing mutant MECP2e1**

To generate neurons in vitro, we found that the Brennand protocol efficiently differentiated the RTTe1-hiPSC lines into neuronal progenitor cells (NPCs) and neurons while the remaining three RTTe1-hiPSC lines required optimization using previously described protocol variations (Table S2) (Brennand et al., 2011; Kim et al., 2012). Cumulative results using neurons derived from all four RTTe1-hiPSC lines are shown for subsequent experiments, and the RTTe1#27 line was utilized for all genetic rescue studies. We observed a 10-fold increase in MECP2 mRNA upon transition of RTTe1-NPCs into neurons (Fig. 3A). Since we were unable to establish RTTe1-hiPSCs expressing the wild-type allele, we compared RTTe1-neurons to those transduced by a lentiviral vector expressing MECP2e1-MYC under the control of the Mecp2 promoter (MeP) or the ubiquitously expressed EF1α promoter (Rastegar et al., 2009). Vector transduction resulted in specific increase of MECP2e1 mRNA in differentiated MeP neurons (Fig. 3B). Total MECP2 protein moderately increased in MeP transduced cells, and was higher in the EF1α transduced cells while MECP2e2 protein levels remained unaffected (Fig. 3C). The relative levels of exogenous MECP2 are most easily quantified by densitometry using the MYC antibody which has a cleaner signal than MECP2, indicating that the EF1α vector expresses roughly twice as much MECP2e1-MYC protein as MeP in neurons similar to the qRT-PCR data shown in Fig. 3B. These results demonstrate that RTTe1 cells continue to express MECP2e2 while the vector transduced cells express different levels of exogenous MECP2e1 protein.

To investigate neuronal identity of the differentiated RTTe1-hiPSC lines, we characterized single cell mRNA profiles using Fluidigm arrays (Pasca et al., 2011). More RTTe1-MeP cells were expressing the mature neuronal markers DCX, NCAM and MAP2 compared to RTTe1 mock infected cells but, importantly, both RTTe1 mock and RTTe1-MeP cells were of comparable neuronal types (Fig. 3D). These included cells with a dorsal forebrain identity indicated by PAX6 expression, a roughly 60:30% distribution of lower and upper cortical layer neurons, respectively; and an equal (~35:45%) mixture of glutamatergic and GABAergic neurons (Fig. 3E), based on concurrent expression of...
region- or neurotransmitter-specific genes from single-cell Fluidigm arrays (see Materials and Methods). To determine the consequences on neurons that lack both isoforms, we compared neurons derived from previously generated MECP2 null iPSCs (Δ3-4#20) to their isogenic (Δ3-4#37) MECP2 WT-neurons (Cheung et al., 2011). The Fluidigm array results showed that complete absence of MECP2 also did not dramatically alter the type of neurons generated (Figs. S2A and S2B). Collectively, these results indicate that lack of MECP2e1 has minimal effects on neuronal differentiation fate in vitro and that cortical neurons relevant to RTT phenotypes were generated.

**MECP2e1 is a level-dependent cell-autonomous regulator of soma size**

RTT is thought to be a neurodevelopmental disorder with defects in neuronal maturation and/or maintenance (Kishi and Macklis, 2004; Nguyen et al., 2012). To determine whether the loss of MECP2e1 alone results in a neuronal phenotype, we measured soma size in RTTe1-neurons. All 4 RTTe1-hiPSCs lines were differentiated into MAP2-positive neurons, with negligible MECP2, consistent with continued low-level expression of MECP2e2 and detectable levels of the MYC tagged MECP2e1 with unchanged levels of MECP2e2 protein. Transduction with EF1α-MECP2e1 construct leads to overexpression of total MECP2 protein and higher levels of MYC tagged MECP2e1. Densitometry bar graph is shown with normalization to loading control, histone H3. (D) Bar graphs show comparable percentage of cells expressing majority of neuronal markers as determined by Fluidigm arrays in RTTe1-mock (cumulative data from all 4 RTTe1 lines) and MeP rescued RTTe1#27 cells. Data are expressed as mean ± SEM; *P < 0.05. (E) Similar percentages of cortical layer and neurotransmitter neurons are produced from both RTTe1-mock and MeP rescue cells. Data are expressed as mean ± SEM; *P < 0.05.
To assess whether the soma size phenotype is due to the specific absence of MECP2e1, we performed additional transductions of RTTe1#27-NPCs with MECP2e2 vectors. MECP2e2 was poorly expressed by the MeP promoter in neurons, likely due to its shorter protein half-life (Yasui et al., 2014). Therefore, higher MeP-MECP2e2 multiplicity of infection (MOI) was used to obtain MAP2 positive neurons with detectable MYC signals (Fig. 5A). As expected, the control MYC-positive MeP-MECP2e1 transduced neurons rescued the soma size equivalent to WT-neurons and the EF1α-MECP2e2 transduced neurons showed no statistically significant differences in soma size (Fig. 5B). Strikingly, the MeP-MECP2e2 vector was unable to rescue (Fig. 5B). We observed that the mRNA levels in MeP-MECP2e2 transduced cells were greater than that in MeP-MECP2e1 (Fig. 5C) but the MYC intensity in MeP-MECP2e2 transduced neurons was not (Fig. 5B). Because soma size can only be measured in single cells, the most accurate comparison of the two rescue vectors was to identify single neurons expressing similar protein levels by binning intracellular immunofluorescence intensity levels of the MYC signal. MYC signal intensity covered the same range (3600–30,000 arbitrary units) in both the MeP-MECP2e1 and MeP-MECP2e2 transduced neurons. We found that low expressing MeP-MECP2e1 RTTe1 neurons (3600–5000 arbitrary units) have a soma size rescue that is statistically significantly different from the high MECP2e1 expressing and mock infected cells, confirming that MECP2e1 rescue is level-dependent (Fig. 5D). On the other hand, neither high (greater than 8000 arbitrary units) nor the similarly low MeP-MECP2e2 expressing RTTe1 cells had a soma size rescue. Since both isoforms contained the same C terminal MYC tag and were expressed from the same vector, the rescue is not due to tag or vector effects. We conclude that MECP2e2 is unable to efficiently rescue soma size in our system when comparing single cells expressing similar levels of the MECP2 isoform transgenes. These data indicate that MECP2e1 is a cell-autonomous regulator of soma size.

**MECP2e1 controls action potentials and excitatory synaptic responses**

We next determined whether glutamatergic RTTe1-neurons exhibit any electrophysiological defects. Whole-cell patch-clamp recordings were performed on over 220 hiPSC-derived neurons from two WT (characterization of SK0019_002#7 hiPSCs is in Fig. S5) and four RTTe1 lines. Electrophysiological studies demonstrated RTTe1-neurons compared with WT-neurons had higher input resistance (Fig. 6A), and lower cell capacitance (Fig. 6B) dependent on the membrane area of the cells (Golowasch et al., 2009; Limon et al., 2005), corresponding to the decrease in soma size in RTTe1-neurons. There is no significant difference in the resting membrane potentials between WT- and RTTe1-neurons (Fig. S6A). These hiPSCs-derived cells had functional neuronal properties and generated tetrodotoxin-sensitive Na + channel-mediated spontaneous and/or evoked action potentials (Fig. 6C, Fig. S6C and S6D). However, evoked action potentials in RTTe1-neurons exhibited smaller amplitude, longer time course (increased rise time, half-duration, and decay time), and fewer numbers as a series of current steps were injected (Fig. 6D, Fig. S6E-I). The defects in generating action potentials in RTTe1-neurons could be attributed to a decrease in voltage-gated Na + currents. Indeed, significant decreases in amplitude and density of voltage-gated Na + currents were observed in RTTe1-neurons (Fig. 6E, Fig. S6B). There was no significant difference in voltage-gated K + currents between WT- and RTTe1-neurons. These findings indicate that RTTe1-neurons have deficiencies in intrinsic membrane properties similar to neurons with MECP2 null mutation (Calfa et al., 2011; Farra et al., 2012; Zhang et al., 2010).

The other evidence demonstrating the functional neuronal properties of these hiPSCderived cells is that WT- and RTTe1-neurons displayed spontaneous synaptic activity (Fig. 7A and Fig. 57). Studies from us (Farra et al., 2012) and other investigators (Chao et al., 2007, Dani et al., 2005; Marchetto et al., 2010; Nelson et al., 2006) have shown that neurons with MECP2 null mutation have synaptic defects.

**Fig. 4.** RTTe1-neurons exhibit a soma size defect that can be rescued by MECP2e1 in a cell-autonomous manner. (A) Immunocytochemistry for MAP2 and MECP2 in WT Δ3–Δ4#37 and RTTe1#27 neurons. Bar graph represents soma size analysis of neurons derived from individual RTTe1 hiPSC cell lines compared to WT neurons derived by three differentiation protocols utilized in the study. (B) Immunocytochemistry for MAP2 and MYC in RTTe1#27-neurons with or without MECP2e1-vectors. Bar graph shows cumulative soma size analysis of all four RTTe1-neurons, and RTTe1#27 neurons with MECP2e1-vectors compared to WT-neurons. Total number of measured neurons for each analysed genotype is indicated within the appropriate bar. Data are expressed as mean ± SEM (P < 0.001; Student’s t-test; n = at least 3 independent differentiations). Scale bars, 44 μm for large image, 10 μm for inset. Arrows, MYC-positive neurons. Arrowheads, MYC-negative neurons.

with MAP2 and MYC revealed that roughly half of the RTTe1-neurons expressed lentiviral transgenes (Fig. 4B). Therefore, soma size was scored in MYC-positive relative to MYC-negative neurons. No statistically significant differences were observed in the EF1α-transduced neurons (Fig. 4B, bars 3 and 4). These neurons express twice the level of MYC tagged MECP2e1 protein (Fig. 3C), which may be incompatible with rescue of neuronal morphology phenotypes similar to the finding that MECP2 duplication causes neurological phenotypes. Only the MeP-transduced MYC positive neurons showed a statistically significant soma size increase in comparison to the adjacent MYC negative neurons (bars 5 and 6). Mock infected cells were all MYC-negative with small soma size (bars 7 and 8). These results reveal that soma size is rescued by MeP-MECP2e1 in a cell-autonomous manner.
Thus, we recorded miniature excitatory postsynaptic currents (mEPSCs) to examine whether RTTe1-neurons have any defects in synaptic function. Estimates of mEPSCs uncovered that RTTe1-neurons displayed dysfunctional synaptic activities with significant decreases in both amplitude (Fig. 7B) and frequency (Fig. 7C) of mEPSCs. Collectively, these findings demonstrate that MECP2e1 controls physiological properties of neurons and its loss triggers immature neuronal phenotypes that are related to reduced soma size and synaptic connectivity.

Discussion and conclusions

We developed RTTe1 patient iPSCs to discover the effect of MECP2e1 mutation on human neurons. RTTe1-hiPSCs retained an Xi allowing the generation of mutant MECP2e1 neurons upon differentiation. This feature of X-inactivation during reprogramming with a preferential retention of one specific Xi in human iPS cell lines has been previously observed by us and others (Cheung et al., 2011; Pomp et al., 2011). Single cell Fluidigm arrays demonstrated that the majority of neurons were cortical in nature with an equal mixture of glutamatergic and GABAergic neurons, and the relative proportion of cell types was unaffected by MECP2e1 or MECP2 null mutations.

We next investigated the effect of MECP2e1 mutation on neuronal form. RTTe1-neurons displayed a soma size defect and reduced dendritic complexity in comparison to WT-neurons. The soma size was rescued with a MECP2e1 transgene. The rescue effect was cell-autonomous as only RTTe1-neurons that received the vector and not their uninfected neighbours exhibited a soma size increase. These results are consistent with the finding that nuclear size is cell-autonomously regulated by MeCP2 in mouse ES cell derived neurons (Yazdani et al., 2012). The heterogeneous expression of the MECP2e1 transgene in our system allowed us to determine that transduced single cells expressing low levels of MECP2e1 had a soma size rescue comparable to WT neurons. Thus MECP2e1 rescue is level-dependent, in agreement with findings that mild overexpression of MECP2 results in neurological phenotypes (Collins et al., 2004). In contrast MECP2e2 transgenes regulated by the same promoter in single neurons with similar low MYC staining intensity did not rescue the soma size of RTTe1 neurons. Since Mecp2e2 transgenes can rescue certain behavioural RTT phenotypes in mice (Kerr et al., 2012), we cannot exclude the possibility that a particular level of MECP2e2 expression during neurodevelopment may rescue some function in neurons. Taking the rescue experiments together with the reproducibility of the soma size defect in neurons derived from all 4 RTTe1-hiPSC lines, we conclude that MECP2e1 mutation reduces soma size in human neurons.

Finally, we evaluated the effect of MECP2e1 on neuronal function by identifying electrophysiological defects in RTTe1-neurons. Our findings of alterations in action potential characteristics, Na+ channel function and synaptic responses in RTTe1-neurons resemble those in RTT-miPSC-derived neurons (Farra et al., 2012) and extend the results from other in vivo RTT mouse model systems (Chao et al., 2007; Dani et al., 2005; Nelson et al., 2006; Zhang et al., 2010). Similarly, defects in synapse formation and function have been previously reported in human ES cell (Li et al., 2013) and hiPSC (Marchetto et al., 2010) models of RTT with loss of both MECP2 isoforms. Reduced expression level of sodium channels has been observed in several RTT models (Kim et al., 2005; Taneja et al., 2009), is reduced in RTTe1-neurons suggests that manipulation of the AKT/mTOR pathway was shown to increase soma size and neurite complexity of MECP2 null human ES cell-derived neurons (Li et al., 2013) although effects on electrophysiological phenotypes were not assessed. In conclusion, our RTTe1 patient iPSC cell model demonstrates that MECP2e1 isoform mutation affects the form and function of human neurons,
and shows that the cellular consequences of disease-causing alternatively spliced transcripts can be defined using patient iPS cells.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.nbd.2015.01.001.

References

Amir, R.E., et al., 1999. Rett syndrome is caused by mutations in X-linked MECP2, encoding methyl-CpG-binding protein 2. Nat. Genet. 23, 185–188.

Ananikov, G., et al., 2011. Logenic pairs of wild type and mutant induced pluripotent stem cell (iPSC) lines from Rett syndrome patients as in vitro disease model. PLoS ONE 6, e25255.

Brennand, K.J., et al., 2011. Modelling schizophrenia using human induced pluripotent stem cells. Nature 473, 221–225.

Calder, G., et al., 2011. Experimental models of Rett syndrome based on Mecp2 dysfunction. Exp. Biol. Med. 236, 3–19.

Chahrour, M., Zoghbi, H.Y., 2007. The story of Rett syndrome: from clinic to neurobiology. Neuron 56, 422–437.

Chahrour, M., et al., 2008. MeCP2, a key contributor to neurological disease, activates and represses transcription. Science 320, 1224–1229.

Chambers, S.M., et al., 2009. Highly efficient neural conversion of human ES and iPS cells by dual inhibition of SMAD signaling. Nat. Biotechnol. 27, 275–280.
