Impaired *in vivo* binding of MeCP2 to chromatin in the absence of its DNA methyl-binding domain

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ABSTRACT

MeCP2 is a methyl-CpG-binding protein that is a main component of brain chromatin in vertebrates. In vitro studies have determined that in addition to its specific methyl-CpG-binding domain (MBD) MeCP2 also has several chromatin association domains. However, the specific interactions of MeCP2 with methylated or non-methylated chromatin regions and the structural characteristics of the resulting DNA associations in vivo remain poorly understood. We analysed the role of the MBD in MeCP2-chromatin associations in vivo using an MeCP2 mutant Rett syndrome mouse model $(Mecp2^{tm1.1Jae})$ in which exon 3 deletion results in an N-terminal truncation of the protein, including most of the MBD. Our results show that in mutant mice, the truncated form of MeCP2 (AMeCP2) is expressed in different regions of the brain and liver, albeit at 50% of its wild-type (wt) counterpart. In contrast to the punctate nuclear distribution characteristic of wt MeCP2, AMeCP2 exhibits both diffuse nuclear localization and a substantial retention in the cytoplasm, suggesting a dysfunction of nuclear transport. In mutant brain tissue, neuronal nuclei are smaller, and Δ MeCP2 chromatin is digested faster by nucleases, producing a characteristic nuclease-resistant dinucleosome. Although a fraction of Δ MeCP2 is found associated with nucleosomes, its interaction with chromatin is transient and weak. Thus, our results unequivocally demonstrate that in vivo the MBD of MeCP2 together with its adjacent region in the N-terminal

domain are critical for the proper interaction of the protein with chromatin, which cannot be replaced by any other of its protein domains.

INTRODUCTION

Methyl-CpG-binding protein 2 (MeCP2) is a basic chromosomal protein that binds to symmetrical methylated 5'CpG dinucleotide sequences (1). Although the protein consists of several well-defined structural domains (2) (Figure 1A), it is an excellent example of an intrinsically disordered protein (5) because of a very low content of secondary structure organization (40%) (5,6). Intrinsically disordered proteins contain low levels of secondary structure that can increase on interaction with other binding partners (7). In addition to DNA, MeCP2 has numerous protein interaction partners (8) and has been shown to interact with RNA (9). It should thus not come as a surprise that mutations throughout the whole protein have potentially deleterious consequences.

The protein is encoded by a single-copy gene located in the X-chromosome and contains 4 exons. Alternative splicing of exon 2 results in two isoforms, E1 and E2, that differ only in the first 24 N-terminal amino acids (10). For historical reasons, many of the structural features of MeCP2 are referred to the shorter E2 isoform, which was identified first in rat (11). Although discovered later, the longer E1 isoform is by far the most abundant (10-fold higher in brain tissue) (10,12,13) and will be used as a reference throughout this article (Figure 1A).

The structural organization resulting from interactions of MeCP2 with DNA and chromatin templates has been studied extensively *in vitro* (14–20) and to a lesser extent *in situ* (20–22). However, interpreting conclusions from

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in vitro studies is complicated in part by the difficulty in reproducing the conditions occurring within the native nuclear environment. In addition to the difficulty in replicating the exact ionic composition and concentrations in the nucleus, the methyl binding preference of MeCP2 by DNA and nucleosome templates can only be mimicked in the presence of competitor DNA (20,23). MeCP2 is a highly basic protein that will bind indiscriminately to DNA under conditions of sufficiently low ionic strength. As a result, the preference of MeCP2 for methylated DNA templates, as assessed from the *in vitro* work, has remained controversial [reviewed in (2)]. Although the interactions of MeCP2 with methylated and non-methylated regions of DNA in vivo remain an issue for discussion [reviewed in (8)], high-throughput DNA sequencing has demonstrated a strong correlation between DNA methylation and the MeCP2 distribution (24,25).

MeCP2 is an X-linked gene, and mutations are associated with >85% of the cases of Rett syndrome (RTT) (26). RTT is a neurodevelopmental disease that affects mainly females, as males hemizygous for the mutation are severely affected (27). As a result of this association, a large amount of the initial research on this protein has focused on this disease. An important part of this effort has been directed towards the development of several mouse models that recapitulate phenotypic features of the human disorder (8). Two well-studied lines are $Mecp2^{tm1.1Bird}$ (28) and $Mecp2^{tm1.1Jae}$ (29), both originally described as protein-null mutations. In the $Mecp2^{tm1.1Bird}$ line, exons 3 and 4 were deleted, resulting in a complete knockout of protein expression. $Mecp2^{im1.1Jae}$ mice contain an in-frame deletion of exon 3. The status of $Mecp2^{im1.1Jae}$ mice as protein-null has remained contentious, although subsequent characterizations have confirmed the presence of stable mRNA (30) and suggested that a truncated protein may be present (31). Direct phenotypic comparisons of the two lines has supported the presence of a partially functional protein in Mecp2^{tm1.1Jae} mice, which have shown a different and often milder phenotype for many parameters, including brain weight, brain region volumes, dendritic spine morphology and cerebellar gene expression profiles (30, 32, 33).

As a result of the exon 3 deletion in the Mecp2^{tm1.1Jae} model (29), the truncated version of MeCP2 lacks 116 N-terminal amino acids, including the 48 N-terminal amino acids of the methyl-CpG-binding domain (MBD), as well as the first 'enriched in proline, glutamic, serine and threonine' (PEST) sequence. However, within exon 4, it retains the complete transcriptional repressor domain, the nuclear localization signal (NLS) (3) and the PEST 2 sequence (4) (Figure 1A). Consequently, this mouse model provides a valuable tool in trying to address the question about the effect of a disrupted MBD on MeCP2-chromatin interactions in vivo. Furthermore, several MeCP2 mutations in patients suffering from RTT occur within the MBD, including R106W, R1333C, P152R, F155S and T158M. Of those, R106W takes place within the region of MBD truncated in the $Mecp2^{tm1.IJae}$ mouse model. Our results show that the N-terminal truncated Δ MeCP2 is still capable of binding to DNA and chromatin, but that the binding specificity is significantly impaired, suggesting that interactions with methylated DNA are necessary for binding specificity. The trafficking of the protein from the cytoplasm to the nucleus is also altered, despite the retained NLS. Our results demonstrate that the RTT-like phenotypic characteristics of the *Mecp2*^{tm1.1Jae} mutant mouse model are a consequence of the truncated protein's inability to properly recognize features of methylated chromatin, impaired nuclear import and accelerated protein turnover.

MATERIALS AND METHODS

Animal breeding

Mecp2 mutant mice (*Mecp2*^{tm1.1Jae}/Mmcd) (MMRRC, UC Davis) (29) and yellow fluorescent protein (YFP)-H transgenic mice [B6.Cg-Tg(Thy1-YFPH)2Jrs/J, Jackson Laboratory Sacramento, CA, USA) (34) were bred and genotyped as described elsewhere (35).

Chromatin preparation and fractionation

Chromatin purification and fractionation was carried out as described elsewhere (22).

Time course micrococcal nuclease digestions of mouse liver and brain nuclei were carried out in 50 mM NaCl, 10 mM piperazine-N,N'-bis(2-ethanesulfonic acid) (PIPES), pH 6.8, 5 mM MgCl₂, 1 mM CaCl₂ buffer with 2 U of micrococcal nuclease per milligram of DNA for 8, 16, 24 and 32 min.

Sucrose gradient fractionation

Chromatin fractions S1 and SE were prepared as described previously (22) and were run on 5-20% sucrose gradients in 25 mM NaCl, 10 mM Tris, pH 7.5, and 5 mM EDTA buffer for 21 h at 96 000g and 4°C and collected thereafter in 0.5-ml aliquots.

Cytoplasmic and nuclear fractionation

Briefly, wild-type (wt) and mutant brain and liver tissues were homogenized in $1 \times$ phosphate-buffered saline (PBS) (137 mM NaCl, 2.7 mM KCl, 4.3 mM Na₂HPO₄ and 1.47 mM KH₂PO₄) and centrifuged at 4500g for 10 min at 4°C. The pellet was added to $1 \times$ PBS, and the sample was split in two aliquots. One half was kept (starting MeCP2), while the other half was centrifuged at 4500g for 10 min. The cellular pellet was homogenized in 0.25 M sucrose, 0.5% NP 40 buffer 1, incubated for 10 min on ice and centrifuged at 4500g to produce a supernatant cytoplasmic MeCP2 fraction and a pellet nuclear MeCP2 fraction. The samples were mixed with 2× SDS sample loading buffer and analysed by western blot.

Gel electrophoresis

SDS-(15%) PAGE (36), acetic acid-urea (AU)-PAGE (37), acetic acid-urea-Triton X-100 (AUT)-(10%) PAGE (38), native-(4%) PAGE (39) and 1% agarose gels were performed as described earlier (40,41).

Histological preparation

Animals were anaesthetized using urethane (2 g/kg i.p.) (Sigma-Aldrich, St. Louis, MO, USA) and transcardially perfused with 10 ml of 0.1 M PBS (pH 7.4), 10 ml of 4% formaldehyde at room temperature in 0.1 M phosphate buffer (PFA), pH 6.5, followed by 10 ml of PFA, pH 11, with a 2-h post-fix in PFA, pH 11. Brain tissue was dissected out and fixed in 4% PFA at 4°C overnight, washed three times in PBS and sunk in 30% sucrose-PBS. Agar-embedded 200-µm coronal sections were cut on a Pelco Vibratome 1000 and coverslipped with Shandon Immunomount (Thermo Scientific Rockford, IL, USA). Animals used for MeCP2 immunofluorescence were processed in the same manner except for fixation conditions: 10 ml of PFA, pH 6.5, followed by 10 ml of PFA, pH 11, with a 2-h post-fix in PFA, pH 11. Cryopreserved brains were frozen on dry ice, and 50um coronal sections were cut on a Leica CM1850 UV cryostat.

Western blotting

Whole brain homogenates from 8-week-old wt and MeCP2 mutant littermates were prepared in 4 volumes of solution A [0.32 M sucrose, 1 mM MgCl2, 0.5% (w/v) NP-40], incubated for 15 min on ice and centrifuged for 10 min at 5000g at 4°C. Pellets were resuspended in solution A and recentrifuged as described above. The pellets were then taken through two rounds of resuspension and centrifugation in 4 volumes of solution B [50 mM NaCl, 5 mM MgCl₂, 1 mm CaCl₂, 10 mM PIPES (pH 6.8)] at 8000g for 10 min at 4°C. The final pellets were resuspended in equal volumes of solution B and $2 \times$ SDS-PAGE buffer to a DNA concentration of 2 mg/ml, boiled for 3 min at 100°C, loaded on a 15% SDS-PAGE gel and blotted onto a polyvinylidene difluoride (PVDF) membrane. All samples contained Complete Protease Cocktail Inhibitor (Roche Molecular Biochemicals, Laval, Quebec, Canada) at a 1:100 dilution. The primary antibodies used were a polyclonal rabbit α -MeCP2 (M9317) targeting C-terminal amino acids 465-478 (Sigma-Aldrich St. Louis, MO, USA) at 1:2000 dilution, and monoclonal mouse α -MeCP2 against N-terminal amino acids 81-170 (Clone 4B6) (Sigma-Aldrich St. Louis, MO) at 1:5000 dilution. Secondary antibodies used were horseradish peroxidase-conjugated anti-rabbit (Jackson ImmunoResearch Inc., West Grove, PA, USA) at $1:10\,000$ dilution and anti-mouse (Rockland Immunochemicals, Gilbertsville, PA, USA) at 1:3000 dilution. Visualization was performed using enhanced chemiluminescence (ECL) (PerkinElmer, Santa Clara, CA, USA).

For quantitative western analysis, the proteins of the SDS–PAGE were blotted onto a 0.2- μ m nitrocellulose membrane, using 25 mM Tris base with 192 mM glycine as transfer buffer. The antibodies used were SIGMA M 9317 α -MeCP2 at a 1:2000 dilution, mouse polyclonal anti-histone H1 [0.T.60] Cat no. ab4269 (Abcam, Cambridge, MA, USA) at a 1:500 dilution and an anti-histone H4 (prepared in-house) at a 1:10 000 dilution. The secondary antibodies used were as follows: anti-rabbit

IRDye[®] 800-conjugated antibody, cat no. 611-132-122 (Rockland, Gilbertsville, PA, USA) at a 1:5000 dilution and goat anti-mouse IRDye[®] 680, Cat no.926-32220 (LI-COR Biosciences, Lincoln, NE, USA) at a 1:10000 dilution. Membrane was incubated with secondary antibodies in dark conditions for 1 h and visualized using Odyssey Licor Developer v.2.1. Band intensity was quantified using Licor software.

Immunohistochemistry

Tissue sections were permeabilized in 1% PBST (PBS, pH 7.4, with 1% Triton X-100) and blocked overnight in 5% goat serum. All antibody incubations were done for 24 h at room temperature in 0.1% PBST. A chicken α-MeCP2 antibody targeting C-terminal amino acids 466-484 (gift from Janine LaSalle, UC Davis) and mouse α -MeCP2 4B6 were used as primary antibodies at a 1:5000 dilution. Alexa 555-conjugated goat antichicken and Alexa 647-conjugated goat anti-mouse (Invitrogen, Carlsbad, CA, USA) were used as secondantibodies (1:500).YO-PRO-1 $(1:10\,000)$ ary (Invitrogen, Carlsbad, CA, USA) was used as a nuclear counter-stain. Sections were mounted on Vectashield Hard Set medium (Vector Laboratories, Burlingame, CA, USA), coverslipped and allowed to set for 24h before imaging.

Plasmid constructs

Wild-type and mutant MeCP2 cDNAs (20) were used in conjunction with the following primers: 5'-GCGCGAAT TCAGTAGCTGGGATGTTAGGGCT-3'/5'-GCGCGG ATCCTCAGCT AACTCTCTCGGTCAC-3' and cloned into the pEGFP-C1 plasmid (Clontech, Mountain View, CA, USA) to allow expression of C-terminal EGFPtagged wt and mutant MeCP2 under the control of the hCMV IE promoter.

Cell culture and plasmid transfection

The Neuro-2A (N2A) mouse neuroblastoma cell line (ATCC) was used for live-cell plasmid transfection experiments. N2A cells were cultured at 37°C and 5% CO₂ in equal parts high-glucose Dulbecco's modified Eagle's medium and minimal essential medium (MEM)+ glutamax (GIBCO/Invitrogen, Carlsbad, CA, USA), supplemented with 10% fetal bovine serum (Invitrogen, Carlsbad, CA, USA), 2 mM L-glutamine (Sigma-Aldrich, St. Louis, MO, USA), 100 U/ml of penicillin (Sigma-Aldrich, St. Louis, MO, USA) and 100 µg/ml of streptomycin (Sigma-Aldrich, St. Louis, MO, USA). The day before plasmid transfections, 1.0×10^6 N2A cells were seeded into 35-mm glass bottom culture dishes (No. 1.5 coverglass; MatTek). Coverslips were coated with 1% gelatin for 2.5h before seeding. Cells were transfected with 2µg of plasmid DNA/plate using Lipofectamine 2000 (Invitrogen, Carlsbad, CA, USA) following manufacturer's instructions, incubated for 18 h in serum- and antibiotic-free culture medium and then cultured for 48 h in complete medium. Hoechst 33342 TriHCl (Molecular Probes/Invitrogen, Carlsbad, CA, USA) was added to culture media 30 min before imaging. Immediately prior to imaging, cells were washed twice with pre-warmed HEPES-buffered extracellular solution, pH 7.4 (150 mM NaCl, 4 mM KCl, 2 mM MgCl₂, 2 mM CaCl₂, 10 mM HEPES, 10 mM glucose and 2 mM ascorbic acid). Cells were maintained in extracellular solution at 37°C during imaging using a TC-344B Dual Automatic Temperature Controller (Warner Instrument Corp., Hamden, CT, USA).

Imaging and image analysis

For MeCP2 immunofluorescence, 1024×1024 pixel 3D confocal fluorescence image stacks were obtained using an Olympus Fluoview FV1000 with Fluoview software (v. 1.7 c) with 488-, 543- and 635-nm lasers, an Olympus Plan SApo NA 0.95 40× oil objective with 10×2 -µm z-steps and pinhole size set automatically (85–120 µm). Multi-channel imaging was done sequentially. All images were averaged twice with scan speeds of 1.68–3.94 µs/pixel (Olympus, Center Valley, PA, USA).

For Neuro-2A live-cell imaging, monochromatic 2560×1920 pixel wide-field fluorescence images were obtained using a Nikon $100 \times / 1.49$ NA Apo TIRF objective (Nikon, Melville, NY, USA), 1.5× zoom, with a Nikon DS-U1 camera and ACT-2U software. Images were converted to 8-bit .tif files, processed and analysed using ImageJ software (NIH). Green fluorescent protein (GFP)/Hoechst image pairs were converted to stacks, and noise pixels were smoothed using a 2-pixel median filter. The means and standard deviations of nuclear greyscale intensity values were obtained over the same outlined area for both channels. Nuclei from 27 wt and 22 mutant cells were outlined based on the Hoechststaining pattern. Coefficients of variation were generated for MeCP2-GFP and Hoechst staining by dividing the greyscale standard deviation by the mean intensity for each cell. A ratio of the coefficients of variations (MeCP2-GFP/Hoechst) was generated for each cell. Genotypes were compared using the Mann-Whitney Utest. Images used for the pixel intensity line plots through the soma and nucleus were converted to 32-bit .tif files, and greyscale values were squared to enhance contrast. Images were adjusted for brightness and contrast and tiled using Adobe Photoshop CS4.

Fluorescence recovery after photobleaching

WtMeCP2-GFP and Δ MeCP2-GFP plasmids with C-terminal GFP were constructed (42) and analysed by fluorescence recovery after photobleaching (FRAP) as described elsewhere (43).

Preparation of tissue samples for real-time quantitative polymerase chain reaction

Brain tissue samples from 8-week-old animals (wt n = 5, Δ MeCP2 n = 4) were rapidly dissected, flash-frozen in isopentane chilled to -20° C and stored at -80° C. Tissues were transferred to chilled RNAlater[®] ice (Life Technologies) at -20° C for at least 16 h before RNA extraction. Approximately 100–150 mg of brain tissue was homogenized in 1 ml of TRIzol[®] (Invitrogen, Carlsbad, CA, USA) using 3-mm stainless steel beads and 1.5 ml Safe-Lock Eppendorf microcentrifuge tubes in a Retsch MM301 Mixer Mill (Fisher Scientific Ltd., Ottawa, Ontario, Canada) at 25 Hz for 9 min. RNA was isolated according to the manufacturer's instructions. Extracted RNA was resuspended in 30 μ l of RNase-free dH₂O and stored at -80° C. QuantiTect Reverse Transcription cDNA Synthesis Kit (Qiagen, Germantown, MD, USA) was used to synthesize cDNA from 1 μ g of total RNA as described by the manufacturer (Qiagen, Germantown, MD, USA). The cDNA reactions were diluted 20-fold for use in PCR amplification. Total RNA and cDNA quality was confirmed by 1.5% agarose gel electrophoresis.

The RT–PCR primer sequences used were: *MECP2* primers: forward 5'-CCCAGGTTTTGTGCTTTGTT and reverse 5'-GGGAGTGCTCTTGTCTCCTG (104-bp amplicon); *rp L8* primers: forward 5'-AAAGGCATCGT AAAGGACAT and reverse 5'-ACCAGCCACAACACC AAC (376-bp amplicon). *rp L8* was used to normalize the target gene (*MECP2*) and to account for any reference gene expression variations that may be present between different samples (44). The suitability of the target and reference gene primers for RT–PCR was assessed by serial dilution validation assay. The slope of the ΔC_t (*MeCP2* gene primer minus *rpL8* normalizer gene primer), for the *MeCP2* primer, versus log cDNA concentration dilution was <0.1, and efficiency of the primer set was >99%.

RT-PCR was conducted using the First Strand Universal SYBR Green Master Mix with Rox (Roche Molecular Biochemicals, Laval, Quebec, Canada) with an Agilent Technologies Stratagene Mx3005P thermocycler. Transcript levels were analysed using MxPro-Mx3005P v4.10 Build 389, Schema 85 software (Stratagene, La Jolla, CA, USA). Cycling parameters were Segment 1 (1 cycle): 9 min. denaturation at 94°C; Segment 2 (40 cycles): 15s denaturation at 95°C, 30 s annealing at 62°C, 45 s extension at 72°C; Segment 3 (1 cycle-for dissociation profile): 1 min at 95°C, 30s at 55°C, 30s at 95°C. Each cDNA 1/20 dilution was run in quadruplicates as were cDNA template-free controls to test for any DNA contamination. Quadruplicate Ct values were averaged for mutant and wt samples, and average Ct values were normalized to the rp L8 control using the comparative Ct method $(\Delta\Delta C_t)$. Data analysis was done using comparative C_t (2- $\Delta\Delta C_t$) method (45) with one wt sample as the comparator for the mutant samples to determine the relative fold changes in Mecp2 expression levels (45-47). The specificity of the amplification reactions was determined by sharpness of dissociation curves and matching of melting temperature (Tm) between samples.

Statistical analyses were conducted using Prism 5 v.5.0.b (Graphpad Software). All data were analysed using the two-tailed unpaired student's *t*-test unless otherwise indicated. Results were considered significant at P < 0.05 and reported as the mean \pm SEM. Data set distributions were tested for normality using the D'Agostino and Pearson omnibus normality test.

RESULTS

MeCP2 protein missing the sequence corresponding to exon 3 of $Mecp2^{tm1.IJae}$ mutant mice is expressed at reduced levels in different regions of the brain and liver

The expression of Δ MeCP2 in the $Mecp2^{tm1.1Jae}$ mouse model of RTT has been controversial. Although the first report of the mutant phenotype showed expression of a faster electrophoretic band by western blot analysis (29), a later report found stable mRNA but was unable to detect the protein by western blot (30).

We therefore decided to revisit this issue. Nuclear fractions from 8- to 11-week-old $Mecp2^{tm1.1Jae}$ mutant and wt mice were probed by western blot using two antibodies, a mouse α -MeCP2 targeting the N-terminus and a rabbit α -MeCP2 targeting the C-terminus (Figure 1B). The first 45 of 89 amino acids of the N-terminal antibody epitope are deleted in the mutant protein. Consistent with the predicted labelling pattern, the C-terminal α -MeCP2 generated both a high MW band in the wt mouse and a fainter, faster band (corresponding to a lower molecular mass) in the mutant, whereas the N-terminal α -MeCP2 only labelled an identical high-molecular mass band in the wt mice. These results are in agreement with the results from the group that created the mouse model (29).

Western blot quantification of the amount of protein expressed in whole brain homogenates carried out as described previously (22) and using IRDye green fluorescence indicated that $35 \pm 3\%$ of the truncated protein is present in the $Mecp2^{tm1.1Jae}$ mice, a result that closely resembles the western shown in Figure 1E of Chen *et al.* (29). As seen in Figure 1C, the RT–PCR graph shows a trend towards a decrease in the relative abundance of Δ MeCP2 mRNA in agreement with Chen *et al.* (29), who reported a reduced Δ MeCP2 transcript in mutant brain using northern-blot analysis. However, statistical analysis of the data in Figure 1C (see 'Materials and Methods' section) revealed that, despite the trend, there was no significant difference (t₇ = 0.9716, P = 0.3636) between wt MeCP2 and Δ MeCP2.



Figure 1. Expression of \triangle MeCP2 in *Mecp2*^{tm1.1Jac} mutant mice. (A) Schematic representation of mouse MeCP2 E1. The N-terminal domain (NTD), MBD, the ID, the C-terminal domains (2), NLS (32) and the two PEST sequences (33) are indicated. (B) SDS–PAGE and western blot of nuclear extracts from *Mecp2*^{tm1.1Jac} mutant (\triangle) and wt mouse brain. CM, chicken erythrocyte histone marker. Top: SDS–PAGE loadings were normalized based on histone H4 contents. Bottom: Western blots were performed using C-terminal MeCP2 (left) or N-terminal MeCP2 (right) antibodies. Both of them label a higher MW band in wt nuclear extracts but only the C-terminal antibody labels a lower MW band in the mutant (\triangle), consistent with the presence of a truncated MeCP2 protein with an N-terminal deletion in the MBD. (C) RT–PCR analysis of wtMeCP2 and \triangle MeCP2 expression in brain.

Wild-type MeCP2 has been shown to be differentially expressed in various brain regions, with the highest expression in cerebellum and cortex (48–50). Therefore, and because of the reduced expression of the protein, we decided to check its expression in different regions of the brain. As seen in Figure 2, Δ MeCP2 is expressed throughout different sections of the brain. However, although a noticeable increase in the expression of wtMeCP2 was detected in wt cortex as previously described, no significant difference could be observed for the expression of Δ MeCP2 among different regions of the *Mecp2^{tm1.1Jae}* mutant brain.

To assess whether the mutant protein is expressed in peripheral tissues, we also measured MeCP2 levels in the liver. Δ MeCP2 is expressed in $Mecp2^{tm1.1Jae}$ mutant mouse liver, but the amount of protein expressed is only $12 \pm 1\%$ of that of wtMeCP2 expressed in wt mice (results not shown). The reason for the further decrease of Δ MeCP2 in this tissue when compared with brain is not clear, but it could be related to the faster turnover rate of proteins in liver when compared with brain (51,52).

Δ MeCP2 exhibits an unusual cellular distribution

Previous immunohistochemical data from the $Mecp2^{tm1.1Jae}$ line have been contradictory, with the original study showing a loss of immunoreactivity in histological sections of hippocampus and cerebellum [see (29), Figure 1F], while a subsequent study found diffuse Δ MeCP2 staining in the hippocampus [see (31), Figure 2G]. We re-investigated this issue by



Figure 2. Distribution of wtMeCP2 and Δ MeCP2 in different sections of the brain. The upper and lower part show a western blot analysis using an MeCP2 and histone H3 antibody, respectively, of the SDS–PAGE analysis of the nuclear protein composition shown in the middle. CB: cerebellum; CT: cortex and MB: midbrain.

double-immunolabelling cortical brain slices using the mouse N-terminal α-MeCP2 and a chicken C-terminal α-MeCP2 to compare wt males, mutant males and heterozygous $Mecp2^{+/-}$ females (Figure 3A). We found that successful double-immunolabelling was highly sensitive to tissue fixation conditions, which may account for the discrepant results in previous reports. The immunoreactivity of the C-terminal antibody was robust under most fixation conditions, but the N-terminal antibody immunoreactivity was extremely sensitive to overfixation, and was only reliably obtained after a brief formaldehyde fixation using a pH-shift protocol (53). In wt males, both antibodies labelled all neurons with a typical staining pattern showing bright puncta against a diffuse nuclear background (Figure 3A, top). Immunolabelled puncta co-localized with heterochromatin regions, as previously described (3,54). In hemizygous $Mecp2^{-/y}$ mutant males, only the C-terminal antibody was immunoreactive, and nuclear labelling was faint and diffuse, with no puncta observed (Figure 3A, bottom). To facilitate comparison of the subcellular distribution patterns of wt with mutant MeCP2 in vivo, heterozygous females were obtained from a cross of the $Mecp2^{tm1.IJae}$ line with the B6.Cg-Tg(Thy1-YFPH)2Jrs/J line, which express YFP in a subset of cortical neurons. Heterozygous $Mecp2^{+/-}$ females showed a mixture of the two patterns of immunoreactivity observed in males. When imaged under high-exposure conditions to amplify the fluorescence signal, wt MeCP2 labelled with both antibodies was strictly localized to the nucleus. In neurons labelled by the C-terminal antibody alone, immunoreactivity was also observed in the cytoplasm, extending up into the primary apical dendrite, albeit at much lower levels. The marked difference in nuclear staining at the border of the nucleus relative to adjacent cytoplasm indicated that Δ MeCP2 still retained a predominantly nuclear localization.

To better characterize the cytoplasmic versus nuclear localization of MeCP2, the cytoplasmic and nuclear fractions of brain cells were obtained and analysed by western blot (Figure 3B). The quantification was carried out using anti-rabbit IRDye-conjugated fluorescent antibodies. In mutant brain tissue, $27 \pm 3\%$ of the Δ MeCP2 is detected in the cytoplasmic fraction as opposed to $\sim 3\%$ of MeCP2, in agreement with the imunohistochemical data shown in Figure 3A. Interestingly, although the NLS is retained, the presence of mutant MeCP2 in the cytoplasm at low levels also suggests either a dysfunction or reversibility of nuclear transport.

Δ MeCP2 exhibits even distribution throughout nuclei and is associated with a significantly reduced nuclear size in the neuronal nuclei of $Mecp2^{tm1.1Jae}$ mutant mice

To look in more detail at the nuclear distribution of Δ MeCP2 with respect to wt MeCP2 in live cells, particularly in light of potential fixation artifacts, Neuro-2A cells were transfected with GFP-MeCP2 and GFP- Δ MeCP2 plasmid expression constructs. As seen in Figure 4A and B, the wt GFP-MeCP2 exhibits enrichment in the pericentric heterochomatin. By contrast, GFP- Δ MeCP2 exhibits



Figure 3. Δ MeCP2 has altered nuclear distribution in fixed brain tissue and is present in the cytoplasm. (A) Confocal fluorescence images of MeCP2immunolabelled coronal sections of motor cortex layer 5. Top eight panels (males): Nuclei are counter-stained with YO-PRO-1 (blue). Neuronal nuclei are larger, typically with bright punctate staining corresponding to dense heterochromatin, whereas glial nuclei are smaller and sometimes irregularly shaped. In wt (*Mecp2^{+/y}*) males, both α -N-terminal (green) and α -C-terminal (red) MeCP2 immunolabelling show bright puncta against a diffuse nuclear background. In MeCP2J mutant (*Mecp2^{-/y}*) males, α -N-terminal MeCP2 labelling is absent, whereas faint diffuse nuclear α -C-terminal MeCP2 labelling is still observed, consistent with the N-terminal deletion and absence of the MBD. Glial MeCP2 signal is not evident in either genotype. Scale bar = 40 µm. Bottom six (female): Heterozygous YFP⁺ *Mecp2*^{+/-} females have a mixture of both MeCP2 immunolabelling patterns (α -N-terminal, blue; α -C-terminal, red). α -C-terminal-MeCP2 labelling is comparatively faint in mutant cells, but can be seen in a high-exposure image (bottom left, greyscale). Under high exposure, immunolabelling in wt neurons for both antibodies is bright to oversaturated, but it remains exclusively nuclear (arrow #3). In mutant neurons, faint diffuse α -C-terminal MeCP2 immunolabelling is predominantly nuclear, but it also occurs in the cytoplasm (arrow #3). Scale bar = 40 µm. (B) SDS-PAGE (1) and western blot (2) analysis of overall MeCP2 present in wt and mutant brain in comparison with the cytoplasmic (C) and nuclear (N) fractions. The red asterisk indicates a degradation product of wt MeCP2.

pan-cellular distribution and seems to be homogeneous throughout the nucleoplasm unlike either wt protein or the distribution of DNA as revealed by Hoechst staining (Figure 4A). This is similar to what was observed immunohistochemically with the native Δ MeCP2 in brain tissue sections (Figure 3A). The absence of correlation between DNA distribution (Hoechst staining) and GFP- Δ MeCP2 is illustrated quantitatively in the line-scan (Figure 4B). Furthermore, the ratio of the variability of GFP- Δ MeCP2 staining relative to Hoechst staining is the inverse of that observed in the wt (Figure 4C). Moreover, in agreement with what was observed in cortical neurons in Figure 3, a substantial component of GFP- Δ MeCP2 is distributed in the cytoplasm.

The nuclear distribution alteration that we observed for GFP- Δ MeCP2 is similar to initial observations by Nan *et al.* (3) using mouse L cells. Similarly, Matsumura *et al.* (55) found that in mouse NIH 3T3 cells transfected with a truncated MeCP2 lacking the complete N-terminal region (including the MBD), the mutant protein also failed to efficiently retain in chromocenters. Moreover, this construct exhibited a diffuse nuclear distribution albeit with subtle enrichment in chromocenters (55). It is important to mention here that in both instances, transfections were carried out in the background of the native wt MeCP2, and the increased presence at chromocenters could arise from the binding of the wt form.

In the course of these experiments, we noticed that transfection of Neuro-2A cells with GFP- Δ MeCP2

systematically resulted in a decrease in the volume of the nuclei when compared with the cells transfected with GFP-MeCP2, similar to what has been observed in neurons of the Mecp2^{tm1.1Jae} mouse (29). A quantification of this phenomenon carried out by comparison of the nuclear size in wt and $Mecp2^{tm1.1Jae}$ mouse cortical neurons showed a 21% decrease in the latter (Figure 4D and E). Qualitative observations of reduced neuronal nuclear size in the hippocampus were made in the original publication describing $Mecp2^{im1.IJae}$ line, but these were not quantified (29). Interestingly, the decreased nuclear size we observed is comparable with the 28% reduction in neuronal nuclear size in the brains of MeCP2-null *Mecp2^{tm1.1Bird}* mice (56). An MeCP2-related size dependence was also observed in embryonic stem cells by the same authors, and it has attributed to an increase in histone H1 on MeCP2 depletion (24,56). Similarly, neurons from the Mecp2^{tm1.1Bird} line were shown to have smaller chromocenters and an increase in the number of nucleoli that exhibit a smaller size when compared with the wt neurons (57). However, as we have described earlier, the brain cells still retain a significant amount (~35%) of Δ MeCP2 in $Mecp2^{tm1.1Jae}$ (Figure 1B-3B), while retaining the exact same amount of histone H1 (Figure 4F). This suggests that the decrease in nuclear volume and nucleolar organization on MeCP2 depletion is not the direct result of a compensatory H1 increase but rather of an MBD-dependent change in MeCP2-mediated chromatin organization that is normally associated to an increase in size of the nucleus.



Figure 4. (A–C) Δ MeCP2 distribution is altered in Neuro-2A cells and does not concentrate in heterochromatin. (A) Live Neuro-2A cells expressing GFP-tagged wt or Δ MeCP2 from transfected plasmid constructs. Nuclei were counterstained with Hoechst 33 342. wt GFP-MeCP2 is restricted to the nucleus and co-localizes with heterochromatin puncta. Mutant GFP- Δ MeCP2 is diffuse throughout the cell but reaches its highest concentration in the nucleus. Dark patches in the nucleus co-localize with bright Hoechst puncta, indicating reduced levels in heterochromatin. Red arrows indicate end points of the line segments plotted in (B). Scale bar = 20 µm. (B) Line plots of greyscale intensities for GFP-MeCP2 (left axis) and Hoechst 33342 (right axis). Although GFP- Δ MeCP2 peak intensities coincide with Hoechst staining and are brightest at heterochromatin puncta, this correlation is not observed with GFP- Δ MeCP2. Greyscale values were squared to highlight subtle differences in staining patterns. (C) The ratio of the variability of GFP- Δ MeCP2 relative to Hoechst staining. The inverse of that observed in the wt. The coefficient of variation was obtained for both GFP and Hoechst greyscale intensity over the entire nucleus. In the wt, a ratio value >1 is observed, indicating a greater variance in the distribution of GFP-MeCP2 relative to Hoechst staining. The inverse is observed in the mutant (Mann–Whitney U-test, P < 0.0001). (D and E) Nuclear size is reduced in cortical neurons of Δ MeCP2 mutant mice. (D) Wide field fluorescence images of Hoechst 33342-stained nuclei. Mutant males (85.00 ± 1.04 µm² in wt versus 67.84 ± 1.71 µm² in mutant, $t_4 = 9.284$, P < 0.01). (F) Western blot analysis of histone H1 and histone H3 with an IRDye fluorescent antibody detection. CE: chicken erythrocyte histones used as a histone marker. B_{wt}: nuclear brain from wt mice; B_{\Delta}: nuclear liver from Wt mice; L_{\Delta}: nuclear liver from Mecp2^{Im1.1Jae} mutant. The numbers underneath the image represent the percentile of MeCP2 with r

ΔMeCP2 exhibits decreased nucleosome-binding affinity

Although all the results described so far indicate that Δ MeCP2 is able to localize to the nucleus, its binding to chromatin remained to be established. To this end, nuclei from wt and $Mecp2^{tm1.1Jae}$ mutant mice were isolated, digested with micrococcal nuclease and fractionated as described earlier (22). As shown in Figure 5A despite the reduced amount of Δ MeCP2 present in the $Mecp2^{im1.1Jae}$ mouse and the lack of a functional MBD, mutant MeCP2 exhibits a similar partitioning to the wt on nuclease digestion in both the wt and mutant mice. The fractions exhibited a similar biochemical composition (Figure 5B and C), although a larger quantity of di-nucleosomes was consistently observed in nuclei from *Mecp2*^{tm1.1Jae} mouse brain as compared with the wt (see Figure 5C SE- Δ). Fraction SI, which is highly enriched in Pol2 (58), contained the largest amount of MeCP2 in both instances (Figure 5A).

On sucrose gradient fractionation (Figure 5D), both MeCP2 and Δ MeCP2 remained bound to mono-nucleosomes of the SI fraction and in both instances a similar amount of free (non-nucleosome-bound) MeCP2 (sedimenting at the upper part of the gradient) is present in this fraction as described previously (22). MeCP2 is also bound to the mono-nucleosome and di-nucleosomes of the SE fraction, although no free sedimenting fraction was observed for Δ MeCP2 (Figure 5E).

A time course of micrococcal nuclease digestion of cell nuclei (Figure 6A) showed that chromatin from *Mecp2^{tm1.1Jae}* mutant brain tissue digested much faster than chromatin from wt brains, but at a similar rate as wt liver nuclei. Interestingly, a di-nucleosome-'resistant' particle is observed at high-digestion times (Figure 6A, SE, 24 and 32 min), as it was also observed in Figure 5C, SE. Although the differences in the micrococcal nuclease digestion rates could be attributed to the differences in the amount of MeCP2



Figure 5. Brain chromatin distribution of wtMeCP2 and Δ MeCP2 in wt and $Mecp2^{im1.IJae}$ mutant mice (Δ). (A) Western blots of nuclear MeCP2 partitioning in the SI, SE and P chromatin fractions on digestion with micrococcal nuclease (20). (B) SDS–PAGE analysis of the SI and SE chromatin fractions. Note the absence of histone H1 (square) in SI. (C) Native PAGE analysis of the DNA composition of the SI and SE chromatin fractions. Cf: pBR322-CfoI–digested DNA marker (the numbers on the left hand side of the gel correspond to the size of the electrophoretic bands in base pairs); D: di-nucleosomes; M: mono-nucleosomes. The arrow head points to the increased presence of di-nucleosomes in $Mecp2^{im1.IJae}$ mutant mice. (D) Sucrose gradient fractionation of the SI chromatin fraction. The graph (upper panel) depicts the absorbance (260 nm) profile of the numbered fractiona collected along the gradient, shown in the lower panel by western blot analysis using MeCP2 antibody. (E) Sucrose gradient fraction [as described in (D)].

present in the different cell types (being the lowest in the wt liver), the amount of histone H1 was almost identical in all of them (Figure 4F). It is possible that the slower rate of digestion observed in wt brain cell nuclei is the result of a higher MeCP2 content in these cells, which is associated with a preference for binding the linker regions connecting adjacent nucleosomes (17,20). The preference of wt MeCP2 for linker DNA in proximity to the nucleosome, similar to histone H1, is likely the result of the highest levels of DNA methylation within these chromatin regions (59) and the affinity of MeCP2 for four-way junction DNA (60), a structure that resembles the DNA organization at its site of entry and exit from the nucleosome. Accordingly, histone H1, which also binds at this nucleosome site, exhibits a similar preference for four-way junctions (61).

Interestingly, the presence of a structurally and functionally impaired MBD in Δ MeCP2 results in a highly transient weak chromatin binding as indicated by the FRAP experiments (Figure 6B). As shown, at almost any time, most of GFP- Δ MeCP2 is free in solution, and the half time residence (t⁵⁰) of the protein was not measurable (below detection limits). GFP- Δ MeCP2 had a t⁵⁰ of 1.3 s in comparison with 21 s for the wt GFP-MeCP2

(Figure 6B, blue diamonds), in agreement with similar results that used GFP-MeCP2 constructs lacking the whole MBD (62) [compare Figure 6A of Kumar et al. (62) with Figure 6B]. Thus, although we have shown that Δ MeCP2 is able to bind to chromatin in the cells of the $Mecp2^{tm1.1Jae}$ mouse model, the binding is weak and dynamic. This finding suggests that in vivo the contribution of the MBD of MeCP2 is critical for proper chromatin binding, and it largely overrides its electrostatic interaction potential. The in vivo relevance of the FRAP data is validated by the similar nuclear/cytoplasmic localization (Figure 4A and B) and decreased nuclear size (Figure 4A) of cells transfected with similar expression vectors of GFP-tagged MeCP2 when compared with neurons from the $Mecp2^{tm1.1Jae}$ mouse (Figure 3A and 4D). Moreover, a single-point mutation in the region of MBD truncated in the Mecp2^{tm1.1Jae} mouse model exhibits a FRAP that is almost identical to MeCP2 lacking MBD (62) or to shown in Figure 6B.

DISCUSSION

Although the interactions of MeCP2 with different chromatin templates have been extensively characterized



Figure 6. Nuclear size, histone H1 composition and chromatin accessibility of wt and $Mecp2^{im1.1Jae}$ mutant mice (Δ). (A) Agarose gel electrophoresis of the DNA fragments obtained by micrococcal nuclease time course digestions of mouse brain and liver at the times indicated on top of the gel. M1 and M2 are the GeneRuler 50 bp and GeneRuler 1 kb DNA ladder markers (Fermentas), respectively. The arrow points to the higher intensity of the band corresponding to the di-nucleosome in the SE fraction of the $Mecp2^{im1.1Jae}$ mutant mice nuclei at high times of nuclease digestion. B_{wt}, B_{Δ} and L_{wt} same as in Figure 4F. D: di-nucleosome; M: mono-nucleosome; T: tri-nucleosome. The numbers under the lanes shown in SE indicate the ratios D: M as determined by densitometry. (B) FRAP of wtMeCP2-GFP and Δ MeCP2-GFP.

in vitro (2), its binding to chromatin in the native in vivo setting is less well understood. Wild-type MeCP2 E1 has a pI of 9.9, and removal of the protein sequence encoded by exon 3 [which removes part of the MBD and part of the N-terminal region (Figure 1A)] renders a protein $(\Delta MeCP2)$ with a 10.1 pI, which is close to that of histone H1 (i.e. human H1.2, pI = 10.9). Hence, the interaction of MeCP2 in vitro with the nucleosome or nucleosome array substrate in the absence of competitor DNA can be largely electrostatically driven (20). The extent to which this mimics physiological conditions and the prevalence of the ionic interactions over those of the MBD domain within the cell is not well known. The in vitro studies of MeCP2 and reconstituted nucleosome complexes have provided evidence for the existence of several specific sites of interaction of MeCP2 with the nucleosome (18,63) in addition to its MBD. Little is known about the role these sites play in the interaction of MeCP2 with the native chromatin template in vivo.

To address some of these issues in the *in vivo* setting, we have taken advantage of the $Mecp2^{tm1.1Jae}$ RTT mouse model, as shown in Figure 1 and 2, which expresses Δ MeCP2 in all tissues examined, albeit at a lower amount than MeCP2 in wt mice (~50% between cytoplasm and nuclei). The data obtained with this model nicely complement previous experiments directed to understand the role of the MBD in MeCP2–chromatin binding *in vitro*, based on transfection of plasmids encoding wt and truncated MeCP2 constructs (3,55,62). The cause of reduced levels of Δ MeCP2 [Figure 1 B and (29)] in the $Mecp2^{tm1.1Jae}$ mouse model is not obvious, however. Although the data shown in Figure 1C could suggest a decrease in Mecp2 expression or mRNA instability, the statistical significance for the observed difference is

not strong. A lower level of expression would be in agreement with the lower presence of Δ MeCP2 RNA detected by northern blot analysis (29). However, the high variability of the data (Figure 1C) could also be in agreement with a previous study that found no difference in mRNA abundance using the same technique in another characterization of the same mouse model (30). A potential alternative and/or complementary explanation could be a higher cellular turnover of \triangle MeCP2 that binds transiently to the chromatin template (Figure 6B) and partitions itself between the cytoplasmic and nuclear compartments differently than native MeCP2 in wt mouse (Figure 3) and in transfected cells (Figure 4). In this regard, it is interesting to note that removal of exon 3 in \triangle MeCP2 results in the loss of one of the PEST sequences (4) of the protein but does not affect the NLS (Figure 1A). PEST sequences are hydrophilic, at least 12 amino acids in length, and typically signal the protein containing them for rapid proteolytic degradation by the 26S ubiquitin proteasome system after ubiquitination at lysine residues within the PEST sequence (64,65). Notably, phosphorylation within PEST motifs precedes ubiquitination of proteins (66); one of the best characterized sites of MeCP2 phosphorylation (S80 in MeCP2 E2; S97 in MeCP2 E1) (67), as well as one of the recently identified ubiquitination sites (K82/K99) (58), both fall within one of these regions. Therefore, it is reasonable to assume that the decrease in expression and the unusual cytoplasmic/nuclear partitioning ratio are just a reflection of an enhanced or altered protein turnover.

We observed a more diffuse distribution of Δ MeCP2 throughout the nucleus of the $Mecp2^{tm1.1Jae}$ mouse model (Figure 3 and 4), which coincides with a reduction in the nuclear size, both in the comparison of transfected N2A cells or in the native brain, without a change in the

histone H1 content (Figure 4F). Such heterogeneous distribution of Δ MeCP2, when compared with the wt, is in good agreement with the results of Nan *et al.* (3), who used an MTase^S cell line that contains only ~5% of the levels of methyl cytosine to show that the location of MeCP2 depends on DNA methylation.

It is possible also that the lower levels of $\Delta MeCP2$ exhibited by the $Mecp2^{tm1.1Jae}$ mouse model are indirectly contributing to this model's phenotype as opposed to $\Delta MeCP2$ itself. In other words, a decrease in the nuclear volume could result in a concentration increase of all the nuclear proteins relative to wt, which could lead to all kinds of biological effects currently unpredictable from our data.

Histone H1 and MeCP2 have both been shown to bind cooperatively to DNA and to methylated DNA, respectively (18,68). The binding constant of MeCP2 to methylated DNA is $\sim 40 \times 10^{-9}$ M (18,69). This is much lower than that corresponding to the binding of the SPKK motif characteristic of the C-terminal motif of some H1 histones $(4 \times 10^{-9} \text{ M})$ (70), or that determined for the binding of H1 to four-way junction DNA constructs $(16 \times 10^{-9} \text{ M})$ (71). The binding affinity of MeCP2 for methylated DNA is also lower than that of histone H1 to the nucleosome $(10 \times 10^{-9} \text{ M})$ (72). Consequently, the mechanisms involved in the competition of MeCP2 with histone H1 for binding to the methylated region of the genome are ambiguous (21). Interestingly, the amount of histone H1 remains the same in the brains of Mecp2^{tm1.1Jae} mice (Figure 4F), suggesting that the reduced levels of Δ MeCP2 relative to wt do not result in a compensatory increase by histone H1, as compared with previous data from embryonic stem cells lacking MeCP2 (56). Furthermore, an increase in H1 should have resulted in a more compact chromatin organization that was more resilient to nuclease accessibility, in contrast to what we observe (Figure 6A). Moreover, this also indicates that the unusually large nuclear volume size of wt neurons where MeCP2 is present in large amounts (Figure 4D) is not the result of a histone H1 deficiency but of the inherent presence of native MeCP2.

Ultimately, any interpretation of the interactions of MeCP2 with chromatin needs to take into consideration the difference in the binding affinity of MBD proteins, including MeCP2, for specific methylated DNA compared with generic non-methylated DNA binding. The difference exhibits a decrease from the nanomolar to the micromolar range (73) (~10-fold for MeCP2), which is likely exacerbated by the presence of chromatin and is far beyond the binding affinity of linker histones. Therefore, in addition to the lacking the ability to bind to methylated DNA, any binding of Δ MeCP2 to chromatin will be secondary to H1 binding and will have to rely on other chromatin-binding sites of the molecule.

The *in vitro* characterization of the interaction of MeCP2 with nucleosomal templates has shown that, in addition to the MBD, MeCP2 contains a methylation-in-dependent chromatin-binding site at its C-terminal end [residues 196–355 (312–501 in wt)]. The weakest interaction of MeCP2 in the absence of a functional MBD observed by us (Figure 6B) and others (62) indicates

that within the cell, MeCP2 binds primarily to methylated DNA through its MBD, and it is in turn further stabilized by the interaction of its C-terminal end with the nonmethylated DNA. This occurs in the opposite order to an earlier proposal based on their *in vitro* analysis (18). The presence of binary chromatin-binding sites within the same terminal region of MeCP2, such as the 195–271 (311–387 in wt) and 272–354 (388–470 in wt) (18) or the 63–102 (179–218 in wt), DNA-binding domain of the intervening domain (ID) (63) may be responsible for the 'resistant' nucleosome dimers observed on nuclease digestion of brain chromatin (Figure 5E and 6A). Such structures may correspond to the existence of MeCP2 molecules sandwiched between two adjacent nucleosomes through interactions with these regions (63).

The MBD of MeCP2 thus seems to confer high specificity and affinity to chromatin binding within the cell. It is important to point out here that the removal of exon 3 of MeCP2 in the Mecp2^{tm1.1Jae} model in addition to an important part of MBD also removes 114 amino acids adjacent to this domain in the N-terminal region (Figure 1A). It has been shown that this region structurally synergises with the MBD to increase its binding to DNA and to chromatin (63). The coupling between these two regions most likely allows for both recognition of the four-way junction-like DNA organization at the entry and exit site of the DNA in the nucleosome, as well as for DNA methylation sites, which preferentially occur in linker regions of chromatin (59). It is possible that in cell types such as neurons where MeCP2 is highly abundant (20,22) other methylation-independent chromatin interactions such as those observed here for Δ MeCP2 also take place. These are likely to involve the C-terminal end of MeCP2, and their more labile interaction with chromatin could account for the apparently unbound MeCP2 that appears on the SI and SE fractions that are quickly released on micrococcal nuclease digestion of wt brain nuclei (4). Overall, the results presented here clearly demonstrate that in the in vivo setting, the DNA methylation-independent interactions of MeCP2 with chromatin are transient, weak and secondary in nature.

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