15q11.2–13.3 chromatin analysis reveals epigenetic regulation of *CHRNA7* with deficiencies in Rett and autism brain

Dag H. Yasui^{1,*}, Haley A. Scoles¹, Shin-ichi Horike², Makiko Meguro-Horike^{2,†}, Keith W. Dunaway¹, Diane I. Schroeder¹ and Janine M. LaSalle¹

¹Department of Medical Microbiology and Immunology, Genome Center, University of California Davis School of Medicine, One Shields Avenue, Davis, CA 95616, USA and ²Frontier Science Organization, Kanazawa University, 13-1 Takaramachi, Kanazawa 920-0934, Japan

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Copy number variations (CNVs) within human 15q11.2-13.3 show reduced penetrance and variable expressivity in a range of neurologic disorders. Therefore, characterizing 15q11.2-13.3 chromatin structure is important for understanding the regulation of this locus during normal neuronal development. Deletion of the Prader–Willi imprinting center (PWS-IC) within 15q11.2–13.3 disrupts long-range imprinted gene expression resulting in Prader-Willi syndrome. Previous results establish that MeCP2 binds to the PWS-IC and is required for optimal expression of distal GABRB3 and UBE3A. To examine the hypothesis that MeCP2 facilitates 15q11.2–13.3 transcription by linking the PWS-IC with distant elements, chromosome capture conformation on chip (4C) analysis was performed in human SH-SY5Y neuroblastoma cells. SH-SY5Y neurons had 2.84-fold fewer 15q11.2-13.3 PWS-IC chromatin interactions than undifferentiated SH-SY5Y neuroblasts, revealing developmental chromatin de-condensation of the locus. Out of 68 PWS-IC interactions with15g11.2-13.3 identified by 4C analysis and 62 15q11.2-13.3 MeCP2-binding sites identified by previous ChIP-chip studies, only five sites showed overlap. Remarkably, two of these overlapping PWS-IC- and MeCP2-bound sites mapped to sites flanking CHRNA7 (cholinergic receptor nicotinic alpha 7) encoding the cholinergic receptor, nicotinic, alpha 7. PWS-IC interaction with CHRNA7 in neurons was independently confirmed by fluorescent in situ hybridization analysis. Subsequent quantitative transcriptional analyses of frontal cortex from Rett syndrome and autism patients revealed significantly reduced CHRNA7 expression compared with controls. Together, these results suggest that transcription of CHRNA7 is modulated by chromatin interactions with the PWS-IC. Thus, loss of long-range chromatin interactions within 15q11.2-13.3 may contribute to multiple human neurodevelopmental disorders.

INTRODUCTION

Imprinting centers (ICs) are loci that control imprinted gene expression in *cis* over several megabases (Mb) of genomic DNA. The Prader–Willi IC (PWS-IC) within 15q11.2–13.3 is genetically defined as the smallest region of overlap of paternal deletions in Prader–Willi patient DNA samples (1). Prader–Willi syndrome (PWS, OMIM 176270) is characterized by hypotonia, obesity, hypogonadism and mild mental retardation (2). The PWS-IC maps to a 4.3 kilobase (kb) region

in the promoter, first exon and part of the first intron of *SNRPN* (1) and forms a bipartite imprinting control center with the Angelman syndrome IC (AS-IC) (3).

15q11.2–13.3 provides an ideal locus for studying the epigenetic effect of chromatin structure on neuronal gene expression for several reasons. First, the PWS-IC acts in *cis* to regulate paternal expression of a 2 Mb gene cluster within 15q11.2–13.3 encompassing *MKRN3*, *MAGEL2*, *NDN*, *SNRPN*, the small nucleolar RNA transcripts (snoRNAs) *HBII-436*, *HBII-13*, *HBII-437*, *HBII-438*, *HBII-85*, *HBII-52*

*To whom correspondence should be addressed. Tel: +1 5307547906; Fax: +1 5307528692; Email: dhyasui@ucdavis.edu *M.M-H. is a Japanese Society for the Promotion of Science Research Fellow.

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and UBE3A-AS (reviewed in 4). Second, 15q11.2-13.3 contains numerous genes with critical neurologic function such as CHRNA7 (cholinergic receptor nicotinic alpha 7), UBE3A and GABRB3 (5–7). Third, deletions and duplications of 15q11.2-13.3 are observed in 1-3% of autism patients (OMIM 209850), revealing the importance of this locus for normal neurologic development (8). Fourth, a neuron-specific 460 kb paternal transcription unit extends through SNRPN and the snoRNA clusters to UBE3A-AS (9). Post-natal neurons also show paternal allele-specific chromatin decondensation of the SNRPN-UBE3A-AS locus that corresponds to increased snoRNA transcription (10). Fifth, the PWS-IC has been shown in previous studies to contain the strongest MeCP2binding site in the 15q11.2-13.3 locus (11,12). Finally, the PWS-IC has been shown to be necessary for normal post-natal development in mice (13).

Proximal 15q is enriched for blocks of segmental duplications that extend to 15q13.3 and predispose the locus for chromosomal rearrangements. 15q13.3 contains the gene CHRNA7, encoding the cholinergic receptor nicotinic alpha 7 subunit (5,14) and its partial duplication CHRFAM7A, a gene that appears to be transcribed but has unknown function (15). The protein product of CHRNA7, CHRNA7, homooligomerizes to form a ligand-gated ion channel, thereby allowing passage of calcium ions when bound by acetylcholine or other ligands such as nicotine (5). CHRNA7/ CHRNA7 appears to be critical for normal neurodevelopment as copy number variation (CNV) of 15q13.3 is associated with autism (16,17), mental retardation (17-19) as well as schizophrenia and epilepsy (20-22). Furthermore, mice with targeted genetic deletion of Chrna7 exons 8-10 have learning and memory deficits (23-25). For these reasons, genetic alterations in 15q13.3 may contribute to multiple human neurodevelopmental disorders.

Epigenetic mechanisms have recently been shown to underlie complex neurologic processes (26). MECP2 loss of function mutations result in Rett syndrome (27) (OMIM 312750), an X-linked dominant neurodevelopmental disorder characterized by regression in late infancy similar to autism (28,29). MeCP2 is a key interpreter of epigenetic marks as a member of a family of proteins that contain a methyl CpG-binding domain (MBD) (reviewed in 30). The MBD of MeCP2 allows the protein to bind to as little as a single methylated CpG di-nucleotide in a DNA sequence (31) and recruit co-factors such as Sin3a and histone de-acetylase HDAC activity (32,33). Therefore, MeCP2 was originally hypothesized to be a transcriptional silencer of methylated gene promoters in cis (32,33). However, recent evidence suggests that MeCP2 has other important epigenetic function including a role in long-range chromatin organization (34). Studies show that MeCP2 can form a molecular bridge between chromatin structures in vitro (35-37). In addition, MeCP2 ChIP-chip studies in SH-SY5Y neurons revealed predominately intergenic MeCP2-binding patterns particularly in 15q13, a region encompassing CHRFAM7A and CHRNA7, suggesting a role in chromatin loop organization and gene expression of specific loci (11). Furthermore, loss of MeCP2 leads to dysregulation of the 15q11.2-13.3-encoded genes GABRB3 and UBE3A (27,38). Thus, MeCP2 with multiple epigenetic

functions including organization of chromatin structure affects gene expression at 15q11.2-13.3 in developing neurons.

In this study, the role of the PWS-IC and MeCP2 as longrange epigenetic regulators of 15q11.2-13.3 chromatin organization was assayed in human neuronal cells by chromatin conformation capture on microarray (4C) and fluorescent in situ hybridization (FISH) analyses. Unexpectedly, a specific PWS-IC interaction with the 15q13.3 genes CHRNA7 and CHRFAM7A overlapped with MeCP2-binding sites. 4C analysis also demonstrated that 15q11.2-13.3 undergoes dynamic chromatin decondensation with neuronal maturation. Quantitative transcriptional analyses showed significantly reduced CHRNA7 transcripts in human frontal cortex samples from Rett syndrome or autism patients with reduced levels of MeCP2. Together, these results suggest that recruitment of CHRNA7 to the PWS-IC by MeCP2 during neuronal development modulates its expression. Therefore, by establishing a link between MeCP2, chromatin organization and CHRNA7 transcript levels, these results are relevant to understand the epigenetic complexity of neurodevelopmental disorders with genetic association with the 15q11.2-13.3 locus and to develop pharmacological therapies common to multiple neurodevelopmental disorders.

RESULTS

To assay 15q11.2-13.3 chromatin structure during neuronal differentiation by identifying chromatin interactions with the PWS-IC, 4C analysis (39,40) was performed. For these experiments, a model human cell culture system for neuronal maturation was used. Treatment of cycling SH-SY5Y neuroblasts with phorbol 12-myristate 13-acetate (PMA) induces cellcycle arrest, a 2-fold increase in MeCP2 expression (41), neurite outgrowth and synapse formation with surface NeuN expression, a marker of mature neurons (42). 4C analysis identifies genomic loci that physically interact with a defined locus of interest or 'anchor locus' and thereby form chromatin loops (40). Specific interactions of the PWS-IC internal 'anchor' site with surrounding 15q11.2-13.3 loci were assayed by hybridizing 'captured' loci to a custom microarray containing 13 Mb of 15q11.2-13.3 and 13 Mb of non-15q11.2-13.3 control loci tiled at high density. As a positive 4C control for chromatin interaction, a known binding of the H19 locus with Igf2 (43,44) was assayed and observed in parallel experiments (Supplementary Material, Fig. S1) providing proof of principle for 4C results.

Identification of PWS-IC interactions with flanking 15q11.2–13.3 sites by 4C analysis was used to characterize overall chromatin organization. First, PWS-IC interactions with 15q11.2–13.3 were determined by bioinformatic analysis of log 2 microarray signal intensities from three replicate 4C libraries using the Tamalpais peak calling algorithm (http://genomics.ucdavis.edu/farnham) (45). PWS-IC interactions are represented as bars (Fig. 1A, rows 2 and 4) shown above the log 2 signal peaks (Fig. 1A, rows 3 and 5) that were uploaded as custom tracks to the UCSC Genome Browser (http://genome.ucsc.edu). Thus, every identified interaction of a 15q11.2–13.3 site with the PWS-IC represents a

chromatin loop. Therefore, chromatin loop organization as assayed by 4C is a high-resolution representation of overall chromatin structure (40).

The PWS-IC forms chromatin loops with heterogeneous elements of 15q11.2–13.3

4C analysis of the entire 15q11.2–13.3 locus during neuronal maturation revealed several insights into chromatin organization of this locus. Closer examination of PWS-IC interaction patterns revealed a dynamic chromatin landscape as chromatin organization was found to be heterogeneous throughout 15q11.2–13.3 both in terms of number and location of PWS-IC interactions forming chromatin loops. To facilitate this analysis, the entire 13 Mb 15q11.2–13.3 region was partitioned based on Giemsa banding pattern as an unbiased means of dividing the locus into three, approximately equivalently sized, subregions (Fig. 1A, boxes A–C, data in Table 1).

In general, initial 15q11.2-13.3 chromatin organization was heterogeneous. Surprisingly, the central 15q12-15q13.1 subregion adjacent to the PWS-IC (Fig. 1A, box B) displayed a more open chromatin conformation in neuroblasts as shown by the significant probability (P = 0.02555) that the number of PWS-IC interactions observed were less frequent than the number expected by random chance (Table 1, row 9) and than either of the more distal subregions (Fig. 1A, box A or C). In contrast, PWS-IC interactions occurred significantly more frequently than random in subregions 15q11.2 and 15q11.2-13.3 (P = 0.00004 and P =0.00001, respectively) as shown in Table 1, row 10. As a control, PWS-IC trans-interactions with a different autosomal locus 6p22.3 occurred with significantly lower frequency than random in both neuroblasts (P = 0.00298) and neurons (P = 0.00629) as expected (Table 1, rows 6 and 9). Surprisingly, there was no obvious correlation between intensity of 15q11.2-13.3 Giemsa bands and density of PWS-IC interactions.

Not only did the initial level of chromatin condensation vary considerably between 15q11.2-13.3 subregions in neuroblasts, but the level of decondensation upon neuronal differentiation also varied by subregion. In the 15q11.2-13.3 locus, 4C analysis of PWS-IC interaction sites revealed marked chromatin decondensation 48 h after PMA treatment in SH-SY5Y neurons (Fig. 1A). This is shown by an overall 2.84-fold reduction (Table 1, row 11) in PWS-IC interactions with flanking 15q11.2-13.3 loci in neurons (Fig. 1A, rows 4 and 5) versus PWS-IC interactions observed in differentiated neuroblasts (Fig. 1A, rows 2 and 3). Interestingly, subregion 15q13.2-15q13.3, which displayed the most compact chromatin in neuroblasts (Fig. 1A, box C), not only showed the lowest level of induced chromatin decondensation at 2.44-fold (Table 1, row 11) during the neuroblast-to-neuron transition but also displayed significantly higher levels of PWS-IC binding in neurons (P = 0.00266, Table 1, row 7) than expected by random chance. Therefore, PWS-IC interaction was particularly abundant in the 15q13.2-13.3 subregion (Fig. 1A, box C), containing the CHRFAM7A and CHRNA7 loci, thereby marking this subregion for further study.

The PWS-IC forms chromatin loops containing key neurodevelopmental genes

4C analyses also revealed a number of novel PWS-IC interactions with genes necessary for normal neurologic function. For example, the *HBII-85* small snoRNA cluster, the loss of which contributes to the Prader–Willi phenotype (9,46), is flanked by PWS-IC interactions. These results suggest incorporation of the snoRNA loci into a distinct 100 kb chromatin loop in undifferentiated SH-SY5Y neuroblasts and an open structure in maturing neurons (Fig. 1B), similar to what has previously been demonstrated by FISH (10).

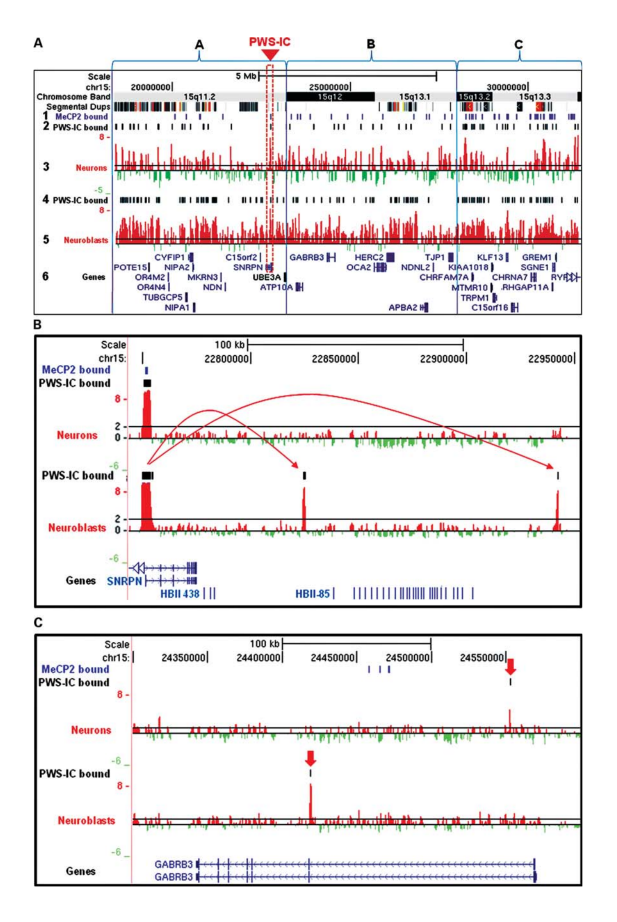
The autism candidate gene *GABRB3* (47) contains a PWS-IC interaction site in the first intron in neuroblasts that divides the gene into two loop domains (Fig. 1C, bottom histogram), but in differentiated neurons this is replaced by an interaction proximal to the transcriptional start site (Fig. 1C, top histogram). This chromatin structure suggests that *GABRB3* chromatin adapts an altered configuration with neuronal maturation. Although single-representative experiments are shown in Figure 1A–C, chromatin structure patterns as assayed by 4C analysis were consistent between three replicate experiments for SH-SY5Y neuroblasts and PMA-treated SH-SY5Y neurons (Supplementary Material, Fig. S2).

MeCP2-binding sites are more frequent in the 15q13.2 and 15q13.3 subregions

To address the hypothesis that MeCP2 recruits discrete 15q11.2-13.3 sub-loci to the PWS-IC, MeCP2-binding sites mapped in a previous study (11) were compared with PWS-IC interaction sites in 15q11.2-13.3. In Figure 1A, MeCP2-binding sites as determined from ChIP-chip data are depicted as the top row of bars (Fig. 1A, row 1) in the UCSC Genome Browser as a custom track. On a 15q11.2-13.3 locus wide scale, 38 MeCP2-binding sites clustered in 15q13.2–15q13.3 (Fig. 1A, row 1, box C), where there were also discrete clusters of sites that were bound to the PWS-IC in both neurons (Fig. 1A, row 2) and neuroblasts (Fig. 1A, row 4). Therefore, MeCP2-binding sites are 6.07 times more frequent in the 15q13.2-15q13.3 subregion than in the 15g11.2 subregion and 3.58 times more frequent than in the 15q12-13.1 subregion (Table 1, row 14). These results suggest that, of all the 15q subregions, MeCP2 was most likely to affect chromatin organization of the 15q13.2-15q13.3 subregion and this was therefore the most likely region to contain genes affected by MeCP2 binding.

MeCP2-binding sites in CHRNA7 interact with the PWS-IC in maturing neurons

Previous studies had identified 62 high-confidence MeCP2binding sites in the 13 Mb comprising the non-repetitive fraction of 15q11.2–13.3, particularly around 15q13.2 and 15q13.3, the majority of which were intergenic (11). These results and *in vitro* studies on chromatin templates (36,37) suggest that MeCP2 might be involved in higher order chromatin organization of 15q11.2–13.3. To test this hypothesis, bioinformatic analysis of the relationship between MeCP2binding sites and sites of PWS-IC chromatin loop interactions



Subregion	15q11.2	15q12-13.1	15q13.2-13.3	Total 15q11.2-13.3	Control 6p22.3
PWS-IC-binding sites in neurons	25	16	27	68	5
PWS-IC-binding sites in neuroblasts	85	42	66	193	19
Total base pairs (TBP)	4 900 000	4 800 000	3 400 000	13 100 000	2 915 400
Neuronal PWS-IC-binding frequency (sites/10 ⁶ TBP, row 2/row 4)	5.10	3.33	7.94	5.19	1.71
P = probability that the number of PWS-IC interactions is at least as <i>low</i> by chance	0.77288	0.09760	0.99873	0.94464	0.00629
P = probability that the number of PWS-IC interactions is at least as <i>high</i> by chance	0.30126	0.93962	0.00266	0.07900	0.99800
Neuroblast PWS-IC-binding frequency (sites/10 ⁶ TBP, row 3/row 4)	17.35	8.75	19.41	14.73	6.52
P = probability that the number of PWS-IC interactions is at least as <i>low</i> by chance	0.99997	0.02555	0.99999	0.99999	0.00298
P = probability that the number of PWS-IC interactions is at least as <i>high</i> by chance	0.00004	0.98258	0.00001	0.0000007	0.99848
Decondensation ratio (row 8/row 5)	3.40	2.63	2.44	2.84	3.81
MeCP2-binding sites in neurons	9	15	38	62	10
MeCP2-binding frequency (sites/10 ⁶ TBP, row 12/row 4)	1.84	3.12	11.18	4.73	3.43
MeCP2 sites/10 ⁶ TBP in 15q13.2–13.3 divided by the total number of MeCP2 sites	6.07	3.58	1.0	2.36	3.26

Table 1. (Quantitative	analysis	of 15q1	1.2-13.3	chromatin	condensation
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The table summarizes chromatin condensation and MeCP2-binding density in subregions of 15q11.2-13.3 chosen based on PWS-IC interaction density in neuroblasts and a control 6p22.3 locus. 15q11.2-13.3 subregions 15q11.2-15q13.3 plus 15q13.2-15q13.3 as shown in Figure 1 were assayed for PWS-IC interactions in neuroblasts (rows 2 and 3). Next the total number of PWS-IC sites was divided by the total number of pWS-IC interactions at least as low or as high as observed was calculated in rows 6, 7, 9 and 10. To determine levels of chromatin decondensation with neuronal differentiation for each region, the frequency of PWS-IC sites per base in neuroblasts was divided by the frequency of PWS-IC sites was divided by total base pairs (row 4) to calculate MeCP2 sites per TBP (row 13). MeCP2 sites/TBP for 15q13.2-15q13.3 were divided by MeCP2 sites/TBP for the other subregions as a means of comparison (row 14).

in 15q11.2-13.3 was performed. Analysis of the total 15q11.2-13.3 MeCP2-binding sites and total PWS-IC interaction sites revealed that distribution of MeCP2-binding sites is random relative to PWS-IC interaction sites overall (data not shown). However, 5 out of 68 or 7.3% of discrete PWS-IC chromatin interactions were also bound by MeCP2 (Supplementary Material, Table S2) identifying them as overlapping binding sites. Furthermore, the proportion of MeCP2 sites that are also sites of interaction with the PWS-IC is much higher than expected by random chance according to Fisher's exact test (Supplementary Material, Table S2), suggesting that these specific PWS-IC and MeCP2 interaction sites have an important biologic function in neurons. As expected, one of the PWS-IC and MeCP2 overlapping binding sites corresponds to the 4C anchor sequence in the PWS-IC itself, where MeCP2 had previously been shown to bind (11,12). However, the remaining four of the PWS-IC and MeCP2 overlapping binding sites in neurons were located in 15q13.2 and 15q13.3 at CHRNA7/CHRFAM7A (Fig. 1A, box C). Two of the PWS-IC- and MeCP2-binding sites flanked the CHRNA7 gene (Fig. 2A), whereas two sites were found in an intron and upstream of the partial

CHRNA7 duplication gene CHRFAM7A (Fig. 2B). Thus, both CHRNA7 and CHRFAM7A interact with the PWS-IC located >7 Mb distal via MeCP2-binding sites, consistent with a long-range chromatin loop structure in neurons.

To further examine the connection between *CHRNA7*/PWS-iC interactions and MeCP2, 4C analysis was performed in neurons following knockdown of MeCP2 (Fig. 2C). For these studies, SHSY-5Y neurons were treated with siRNA targeting the 3' UTR, and MeCP2 knockdown was confirmed by western blot (Supplementary Material, Fig. S8). Knockdown of MeCP2 protein levels in neurons correlated with disrupted PWS-IC interactions with *CHRNA7* as shown by reduction in peak height compared with neurons with normal levels of MeCP2 (Fig. 2C). These results suggest that MeCP2 connects *CHRNA7* to the PWS-IC through long-range chromatin interactions.

CHRNA7 interactions with the PWS-IC are confirmed by FISH

In order to confirm the interaction of the PWS-IC with CHRNA7 detected by 4C analyses, an independent assay using FISH analysis was performed. For this assay,

Figure 1. Dynamic 15q11.2-13.3 chromatin decondensation during neuronal maturation. (A) 4C analysis of 15q11.2-13.3 chromatin structure in developing neurons shows developmentally regulated chromatin decondensation in maturing neurons. 15q11.2-13.3 loci that contact the PWS-IC (red box) are shown as a series of log 2 signal peaks from representative hybridization of 4C libraries isolated from developing SH-SY5Y neurons (row 3) and untreated SH-SY5Y neuroblasts (row 5). 4C peaks corresponding to sites of PWS-IC interaction were determined using a bioinformatic analysis of three independent replicate experiments for both PMA-treated neurons and untreated neuroblasts and are shown as a series of black bars above the corresponding peaks (rows 2 and 4). MeCP2-binding sites determined from previous studies are shown as a series of purple bars in row 1. 15q11.2-13.3 was subdivided into regions A–C based on Giemsa banding patterns in metaphase chromosomes (top row). Log 2 signal ratios are shown as red and green histograms in the UCSC Genome Browser with known genes and transcripts shown in blue font (row 6). A black line at a log 2 signal ratio of 1 is included to enable comparison of peaks in row 3 and 5. Segmental duplications are shown as a series of colored bars below chromosome bands. (B). A high-resolution example of PWS-IC interactions flanking the *HBII-85* snoRNA cluster in neuroblasts (bottom histogram) that are lost during neuronal differentiation (top histogram). Arrows show PWS-IC-binding interactions. (C). PWS-IC interactions shown as black lines above the log 2 signal peaks with the autism candidate gene *GABRB3* are also altered by neuronal differentiation. PWS-IC interactions are indicated by red arrows.

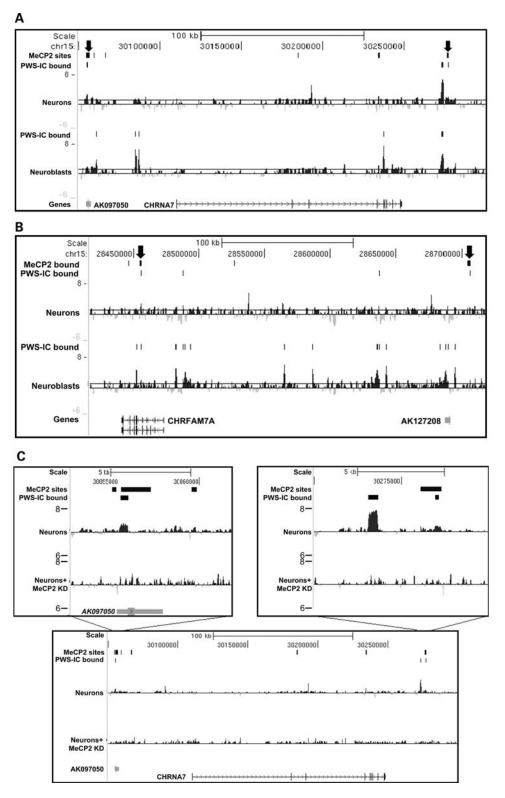


Figure 2. PWS-IC- and MeCP2-bound sites map to *CHRNA7* and *CHRFAM7A*. 4C analysis reveals PWS-IC interaction sites that are also bound by MeCP2 in 15q13.3. (**A**) A 250 kb scale view of *CHRNA7* chromatin structure reveals two sites (arrows) that are bound both by the PWS-IC (PWS-IC peaks) and MeCP2 (top bars) as determined by previous ChIP-chip analysis. 4C interactions shown as log 2 signal ratio peaks of differentiated SH-SY5Y 4C libraries identify one bivalent site 50 kb upstream of *CHRNA7* near the *AK097050* transcript in neurons undergoing differentiation (upper histogram) and another site ~30 kb down-stream. (**B**) Overlap between MeCP2-bound and PWS-IC-bound sites was also observed within an intron of *CHRFAM7A* (left arrow) and upstream of the transcript (right arrow) in maturing neurons. (**C**) MeCP2 knockdown by siRNA in SH-SY5Y cells reduces interaction of *CHRNA7* with the PWS-IC. Insets show PWS-IC interactions in neurons with normal levels of MeCP2 (top histogram) as peaks compared with neurons with reduced MeCP2 levels (bottom histogram) with diminished PWS-IC interaction signals.

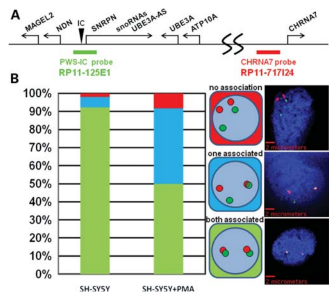


Figure 3. PWS-IC interaction with *CHRNA7* is validated by FISH. Two-color FISH analysis confirms the interaction of the PWS-IC with *CHRNA7*. (A) BAC clones complementary to the PWS-IC (RP11-125E1) and *CHRNA7* loci (RP11-717124 as shown on the 15q11.2–13.3 map were differentially detected as green and red fluorescent signals, respectively). (B) PWS-IC and *CHRNA7* signals were considered associated if there was a spatial overlap or contact between the red and green signals. One hundred nuclei from undifferentiated and maturing (SH – SY5Y + PMA) neurons were scored for signal overlap in each allele and the possible combinations (middle diagram) and were graphed as a percentage of total nuclei (left bar graph). Representative images of each PWS-IC/*CHRNA7* allelic combination are shown in the right panel with scale bars.

differentially labeled genomic probes to the PWS-IC (green) and *CHRNA7* genomic loci (red) (Fig. 3A) were hybridized to SH-SY5Y cells treated with the same neuronal differentiation protocol established for 4C analyses (10). Consistent with 4C results, >90% of neuroblast nuclei showed PWS-IC signals overlapping with *CHRNA7* signals on both 15q11.2–13.3 alleles (Fig. 3B, bar graph), indicating close physical proximity of the two loci (Fig. 1A, rows 4 and 5). In contrast, FISH analysis of SH-SY5Y neurons revealed that 50% of nuclei had a clear separation of PWS-IC (green) and *CHRNA7* (red) signals on one (40%) or both (10%) of 15q11.2–13.3 alleles compared with neuroblasts (Fig. 3B, bar graph).

As controls for FISH analysis of PWS-IC interactions, the 15q11.2–13.3 internal locus *TJP1* and an external *PML* locus in 15q24.1 revealed that only in 8.7% in SH-SY5Y neuroblasts and 4.1% neuronal nuclei was there any association of the PWS-IC with *TJP1* on one or both alleles, whereas only 18.8% of one or both *PML* alleles in neuroblasts or 23.5% of neurons showed association with the PWS-IC (Supplementary Material, Figs S3 and S5). This suggests that *TJP1*, which is 2.3 Mb closer, and *PML*, which is over 50 Mb distant from the PWS-IC than *CHRNA7* rarely interact with the PWS-IC unlike *CHRNA7* and that neuronal maturation did not lead to increased association. Together, these results independently confirm that the PWS-IC is in close proximity to *CHRNA7* on both alleles in neuroblasts and that neuronal maturation

correlates with dissociation of the PWS-IC from *CHRNA7* on one and sometimes both alleles. These results also show that the PWS-IC interaction with *CHRNA7* is not a general property of the distal chromosome 15 *PML* locus. Most importantly, these results are consistent with the 4C analyses that reveal specific interactions between the PWS-IC and *CHRNA7* primarily in neuroblasts and 15q11.2–13.3 chromatin decondensation with differentiation to neurons.

CHRNA7 transcript levels in human cortex decrease with age

As CHRFAM7A is reported to have very low levels of expression in the brain (48) and its biologic function is obscure, quantitative reverse transcriptase-polymerase chain reaction (QRT-PCR) analyses focused on CHRNA7 transcript levels. Although CHRNA7 has been previously assayed in brains of adult schizophrenics (48,49), little is known about its transcript levels during early human developmental stages. For CHRNA7 analysis, RNAs were extracted post-mortem, from Brodmann area 9 obtained from a wide age range of normally developing human subject brains, 7 brains from classic Rett patients and 11 patients diagnosed with autism. QRT-PCR analysis of CHRNA7 transcript levels normalized to GAPDH levels from typically developing cortices revealed high levels of expression in subjects younger than 1 year of age and a reduction in transcript with increasing post-natal age before leveling off around 20 years of age (Fig. 4). Strikingly, CHRNA7 transcript levels were roughly 10-fold higher in the youngest (<1 year of age) than the oldest typically developing (control) brain samples (57 years of age). Therefore, a one-way analysis of co-variance (ANCOVA) was performed, which revealed a significant (P < 0.0001) effect of age on CHRNA7 expression. Interestingly, CHRNA7 transcripts appeared to be reduced in Rett and autism cortices (Fig. 4, filled symbols).

CHRNA7 transcript levels are significantly reduced in Rett and autism cortices

To test the hypothesis that MeCP2 modulates expression of CHRNA7 in vivo, QRT-PCR analysis was extended to RNAs isolated from frontal cortices of classic Rett syndrome patients and autism samples, where MeCP2 has been shown to be significantly less abundant (50). Since our findings in Figure 4 suggest that there is an apparent reduction of CHRNA7 in Rett and autism cortical samples, a one-way ANCOVA using age as a covariate was performed. This age covariate analysis determined that CHRNA7 transcript levels were significantly reduced (P < 0.05) compared with control samples for both Rett (RTT, P = 0.022) and autism (AUT, P = 0.012) (Fig. 5) even when cortical samples of all ages were included. Also, when cortical RNA samples were segregated into two age groups based on our findings in Figure 4. brain cortices of subjects 20 years of age and younger displayed a significant reduction in CHRNA7 transcripts compared with controls (Supplementary Material, Fig. S7). The results shown in Figure 5 and Supplementary Material, Figure S7, are consistent with results in Mecp2-deficient mice, where *Chrna7* transcripts are significantly decreased

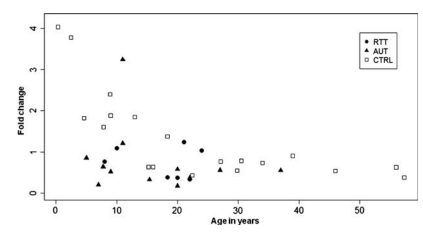


Figure 4. *CHRNA7* transcript levels decline significantly with age in the human cortex. QRT–PCR analyses of *CHRNA7* expression in normally developing control (CTRL, N = 20), Rett (RTT, N = 7) and autism (AUT, N = 12) were performed on RNAs isolated from frozen, post-mortem Brodmann area 9 cortices using *CHRNA7*- and *GAPDH*-specific primers. Fold change of *CHRNA7* expression relative to *GAPDH* was calculated using the comparative Ct method for each sample. *CHRNA7* expression declines rapidly from 4.0-fold relative to *GAPDH* in 4-month-old cortex to ~0.4-fold in 57-year-old cortex. Analysis of *CHRNA7* expression levels was done using a one-way ANCOVA with age as a covariate showing that age has a significant relationship to *CHRNA7* expression (P < 0.0001).

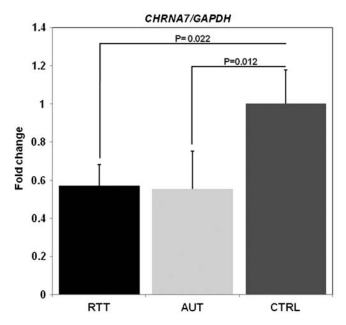


Figure 5. *CHRNA7* transcripts are significantly reduced in Rett and autism corticies. QRT–PCR analyses identify significant reductions in *CHRNA7* expression in Rett and autism frontal cortices. RNAs were isolated from Brodmann area 9 of patients with classic Rett syndrome (RTT, N = 7) and autism patients (AUT, N = 11). RNAs were amplified and quantified using *CHRNA7*- and *GAPDH*-specific primers, and *CHRNA7/GAPDH* fold changes were calculated using the comparative Ct method and graphed, and normalized to the typically developing controls (CTRL, N = 20). Significant reductions (P < 0.05) in *CHRNA7* transcripts are observed in both RTT and AUT samples compared with CTRL while controlling for the effect of age. Fold changes for RTT and AUT samples are normalized to CRTL samples and error bars correspond to standard error of the mean. *P*-values are derived from ANCOVA with age as a covariate.

(P < 0.05) at 1 week of post-natal development in total RNAs isolated from the whole brain (Supplementary Material, Figure S4). Age was found to have a significant effect on *Chrna7* expression (P < 0.001) by ANCOVA (data not

shown). Together, these novel results suggest that MeCP2 positively regulates *CHRNA7* expression in human cortex during neuronal maturation by long-range interaction with the PWS-IC.

DISCUSSION

This study used a novel 4C analysis in order to gain insight into the role that chromatin structure plays in the regulation of 15q11.2-13.3 genes. These results provide many insights into the organization and regulation of this locus. Primarily, these results indicate that PWS-IC interactions within 15q11.2-13.3 are dynamic during neuronal maturation, suggesting dramatic chromatin remodeling at this chromosomal subregion. These results complement studies of the 15q11.2-13.3 syntenic region 7q5 in the mouse brain that show a developmentally regulated decondensation of *Snrpn-Ube3A* preceding an exponential rise in SnoRNA accumulation and maturation (10). Together, these *in vitro* and *in vivo* results suggest that chromatin decondensation and remodeling regulates 15q11.2-13.3 gene transcription during neuronal maturation.

The PWS-IC is a well-characterized long-range control element of 15q11.2-13.3 (51). 4C chromatin structure results confirmed by FISH analyses showed that the PWS-IC makes specific contact with sites flanking CHRFAM7A and CHRNA7 in maturing neurons. FISH analysis of 15q11.2-13.3 conformation in developing SH-SY5Y neurons also revealed PWS-IC and CHRNA7 signal disassociation, consistent with decondensation of the intervening chromatin at this locus during neuronal maturation on one allele. Again, this result is consistent with previous in vivo results by Leung et al. (10), where the 15q11.2-13.3/7q5 paternal allele visualized by FISH analysis becomes strikingly decondensed during neuronal development . 4C chromatin structure analysis also reveals a PWS-IC interaction site upstream of GABRB3, a neuro-developmentally essential gene known to be dysregulated by loss of MeCP2 (27,38). Therefore, chromatin structure analysis directed our hypothesis that *CHRNA7* makes a functional molecular connection with the PWS-IC. Although ICs can regulate imprinted gene expression over several megabases, it is surprising that the PWS-IC may epigenetically regulate *CHRNA7*, which is over 7 Mb away.

In addition, 4C results indicate that clusters of PWS-IC interaction sites in the 15q13.2–15q13.3 subregion overlap with regions containing segmental duplications in both neurons and neuroblasts. In particular, PWS-IC interactions cluster in two 300 kb regions of segmental duplication including portions of the *CHRFAM7A* and *CHRNA7* genes. It has recently been shown that low copy repeat elements predispose the 15q13.3 locus to duplications and deletions that are often found in patients with neurodevelopmental disorders (reviewed in 52). Therefore, it is tempting to speculate that PWS-IC interactions form the bases of chromatin loops bearing the low copy number repeats, making them susceptible to deletion or duplication by homologous recombination.

4C analysis in SH-SY5Y neurons with and without MeCP2 knockdown suggests that MeCP2 has a role in linking the 15q13.3 region, located over 7 Mb distant, with the PWS-IC. Support for the role of MeCP2 in chromatin loop formation comes from Horike et al., who described a role for MeCP2 in chromatin loop formation and transcriptional regulation of the Dlx5/Dlx6 locus in the developing striatum (34,53). Also, in vitro chromatin assays show that MeCP2 can bridge and condense chromatin arrays (35) through multiple DNAbinding domains to form complex looped structures (37). Interestingly, a neuronal enhancer of Chrna7 expression has been recently identified in the Chrna7 intron 4 (54) near the human PWS-IC/MeCP2 interaction site observed in this study, further suggesting a potential site of action for MeCP2 binding to Chrna7. Recent results from Skene et al. (55) also indicate a function for MeCP2 in neuronal chromatin structure and repression of genome-wide endogenous retrovirus transcription. Together, the results shown here are consistent with independent studies examining the role of MeCP2 as a neuronal chromatin structural organizer at high abundance in maturing neurons.

The 15q11.2-13.3 chromatin organization studies revealed a potential and clinically relevant MeCP2 target gene CHRNA7. CHRNA7 levels were found to be significantly reduced in Rett and autism brain frontal cortices. In these studies, CHRNA7 expression was first found to be developmentally regulated with postnatal age in humans. This is shown by a 10-fold reduction in CHRNA7 from a high at 4 months of age to a leveling off of transcript levels around 20 years of age in normal cortices. Intriguingly, CHRNA7 levels in the frontal cortex were found to decline rapidly around 1-2 years of age, corresponding with the age of onset of autistic symptoms (56) and age of diagnosis of Rett symptoms (57). These initial results led to the discovery of reduced CHRNA7 levels in the human cortex from autism and Rett brains, where MeCP2 is reduced or mutated as described previously (50). For these studies, Brodmann area 9 was selected, as it is part of the dorsolateral prefrontal cortex, where previous studies have linked changes in this area to loss of function in autism (58). Analysis of this cortical region is also justified by transgenic mouse studies where genetically engineered loss of MeCP2 expression

in neurons is sufficient to cause symptoms similar to Rett syndrome (59,60). These results, along with the previous finding that *MECP2* levels are also significantly reduced in these samples (50), suggest that MeCP2 positively modulates expression of *CHRNA7* and that MeCP2 deficiency results in *CHRNA7* deficiency in childhood and adolescence. Most importantly, these results are the first *in vivo* evidence that MeCP2 may positively regulate *CHRNA7* through long-range chromatin interaction and suggest that reduced expression may lead to abnormal brain function observed in autism and Rett syndrome.

Independent support for the role of MeCP2 as a positive modulator of Chrna7 comes from RNA expression profiling that revealed significant up regulation of Chrna7 expression in 6 week post-natal mouse hypothalamus engineered to overexpress MeCP2 (61). As CHRNA7/Chrna7 expression is highest in the earliest stage of postnatal development examined, it may also be dysregulated in embryonic stages of development, where reduced abundance may have a persistent effect on later neuronal development and function. In support of this hypothesis, CHRNA7 has reported to have high expression in nuclei that receive sensory input such as the hippocampus and cortex in the fetal brain (62). In contrast, MeCP2 has very low levels of expression in mouse embryonic brain outside of the thalamus and early post-natal cortex (63). Therefore, it is tempting to speculate that a rising concentration of MeCP2 protein in cortical neuronal nuclei during post-natal life dampens the expression of CHRNA7/Chrna7 during late infancy through adolescence. However, the results presented here indicate that loss or reduction in MeCP2 protein is concomitant with reduced CHRNA7/Chrna7 expression in developing humans and mice. Therefore, MeCP2 may act to keep CHRNA7 at an optimal level during postnatal development by both positive and negative modulation. Although the evidence of direct MeCP2-binding sites around CHRNA7 suggests that MeCP2 is a direct transcriptional modulator of CHRNA7 similar to the case of Bdnf (64), it is possible that the effect of MeCP2 deficiency on CHRNA7 is indirect. Nonetheless, these novel results suggest that reduced CHRNA7/Chrna7 expression may contribute to neurologic abnormalities observed in Rett and autism patients as well as in Mecp2-deficient mice.

Epigenetic mechanisms contribute to multiple neurologic disorders (26,65). Specifically, previous results have clearly demonstrated that loss or altered expression of the epigenetic factor MeCP2 leads to severe defects in neurologic development characteristic of Rett syndrome (57). Although the mechanisms underlying the Rett phenotype are not completely understood, the results presented here suggest that MeCP2 acting through the PWS-IC may contribute to the precise epigenetic 15q11.2-13.3 chromatin remodeling process that regulates expression of CHRNA7 and other key genes such as GABRB3 and UBE3A during neuronal maturation. Therefore, aberrant 15q11.2-13.3 chromatin remodeling caused by defects in MeCP2 mutation or deficiency could lead to neurodevelopmental gene dysregulation and thereby contribute significantly to the overlapping Rett and autism phenotypes. A recent phase 2 clinical trial of a CHRNA7 partial agonist in schizophrenic patients significantly reduced negative symptoms, and suggests a potential therapeutic strategy for Rett and autism patients (66).

MATERIALS AND METHODS

4C analysis of chromatin structure

4C analysis was performed according to Simonis *et al.* (40). Briefly, SH-SY5Y neurons were treated with formaldehyde to crosslink chromatin, followed by quenching of the reaction with 1 M glycine and cell lysis by resuspension in hypotonic buffer and dounce homogenization. Cell nuclei were resuspended in restriction buffer and digested overnight with BglII (NEB, Ipswich, MA, USA) at 37°C followed by ligation using T4 ligase (Promega, Madison, WI, USA). DNA was purified from nuclei using proteinase K and RNAseA digestion, followed by phenol/chloroform extraction. Purified DNA was digested using DpnII (NEB), re-ligated with T4 ligase (NEB), re-precipitated and then re-digested with Mnl1 (NEB) overnight at 37°C. 4C libraries were created by linear amplification of DNA fragments with Expand Long Template PCR System (Roche Applied Sciences, Indianapolis, IN, USA). Amplified 4C library fragments were evaluated by gel electrophoresis and column-purified using a PCR clean-up kit (Qiagen, Valencia, CA, USA).

A custom DNA microarray tiling 26.3 Mb of NCBI34/hg17 human genome sequence including 13.3 Mb of 15q11.2-13.3, 2.6 Mb of 11p15.5, 3.8 Mb of 7q21.3, 2.9 Mb of 19p13.2, 0.3 Mb of 20q11.21, 1.5 Mb of 2p25.1, 0.4 Mb of 1p36.12, 2.9 Mb of 6p22.3 and 0.2 Mb of 11p14.1 with repetitive sequences removed was designed and produced by Roche-NimbleGen, Madison, WI, USA (11).

Labeling and hybridization of DNA for 4C analysis were performed according to the Nimblegen Array User's Guide version 3.1 (Nimblegen). One microgram of each DNA sample was denatured in the presence of 5'-Cy3- or Cy5-labeled random nonamers (TriLink Biotechnologies) and incubated with 100 units of (exo-) Klenow fragment (NEB) and dNTP mix (6 mM) each in TE buffer (10 mM Tris/1 mM EDTA, pH 7.4) (Invitrogen) for 2 h at 37°C. Reactions were terminated by addition of 0.5 M EDTA (pH 8.0), precipitated with isopropanol and resuspended in water. Then, 13 µg of the Cy5-labeled 4C sample and 13 µg of the Cy3-labeled genomic input sample were mixed together, dried down and resuspended in 40 µl of Nimblegen Hybridization Buffer (Nimblegen) plus 1.5 µg of human COT1 DNA. Hybridization was carried out in a MAUI Hybridization System (BioMicro Systems, Salt Lake City, UT, USA) for 18 h at 42°C.

The arrays were washed using Nimblegen Wash Buffer System, dried by centrifugation and scanned at 5 µm resolution using the GenePix 4000B scanner (Axon Instruments, Union City, CA, USA). Fluorescence intensity raw data were obtained from scanned images of the arrays, using NIM-BLESCAN 2.0 extraction software (Nimblegen). For each spot on the array, log 2 ratios of the Cy5-labeled test sample versus the Cy3-labeled reference sample were calculated. Then, the bi-weight mean of this log 2 ratio was subtracted from each point by the software. To control for labeling effects, Cy3/5 dye labels were also swapped between the 4C samples and total input DNA prior to hybridization. 4C peaks in three biologic replicate experiments were determined using Tamalpais peak-finder algorithm (45) and categorized into levels L1, L2

MeCP2 knockdown by siRNA

MeCP2 levels were knocked down in SH-SY5Y neurons, TAAAGACA sequence in MECP2 designed and produced by iGENE Therapeutics (Tsukuba, Japan). SH-SY5Y cells were subjected to siRNA treatment and tested for MeCP2 knockdown by western blot (Supplementary Material, Fig. S8). For western blot analysis, 25 µg of SH-SY5Y protein was resolved by SDS-PAGE, transferred to PVDF nylon membranes and probed for MeCP2 using a custom carboxyl terminal antibody raised in chicken and GAPDH using a mouse monoclonal antibody (Advanced Immunochemical, Inc., Long Beach, CA, USA).

PWS-IC regional binding analysis

In order to determine whether the probability that the number of 4C peaks identified in a subregion is greater or less than expected by chance, the following binomial probability formula was used:

$$P_{(k \text{ out of } N)} = \frac{N!}{k!(N-k)!} (p^k)(q^{N-k}),$$

where N denotes the total number of peaks; k, the peak (number ranging from 0 to N); p, percentage of area subsection takes in array; q = 1 - p.

Thus, this equation calculates the chance that a number of peaks (k) would be identified in a subregion with probability (P) with the total number of peaks found (N). To determine the chance that at least the number of peaks would be found by chance, all values of P were summed from k equaling the peaks found to k equaling N. To determine the chance that no more than the number of peaks would be found by chance, all values of P were summed from k equaling 0 to kequaling the number of peaks found. Therefore, all summed *P*-values < 0.05 were called as significant because the 4C peaks were not random with >95% certainty.

Data analysis of 4C peak relationship with MeCP2-binding sites

To determine whether MeCP2 peaks (at the L3 level) were significantly closer or further from 4C peaks (at the L3 level) than expected by chance, each of the 68 4C peaks were assigned a nearest MeCP2 peak neighbor. The 68 distances were then categorized as either near or far. To determine how often these nearest-neighbor distances would be expected by chance, 100 random peak data sets were generated over the same genomic interval, each with 62 peaks as in the MeCP2 peak data set. Nearest neighbors and distances to the 4C peaks were computed for each data set and the results were combined for use in the two-sided Fisher's exact test shown in Supplementary Table S1.

FISH analysis

DNA FISH analysis was performed as described in Leung et al. (10). Briefly, FISH was performed in SH-SY5Y neuroblasts and SH-SY5Y neurons differentiated by 48 h treatment with PMA (11,12). Bacterial artificial chromosome (BAC) DNA corresponding to the PWS-IC and 5' region of CHRNA7 (Fig. 3) was isolated from clones obtained through BacPacResources (Oakland Children's Hospital, Oakland, CA, USA). BAC DNA was labeled by nick translation to create probes, hybridized to cells fixed in Histochoice (Amresco, Solon, OH, USA) and then dehydrated in ethanol. FISH probes were denatured with the fixed cells at 80°C for 3 min and then hybridized overnight at 37°C on cover-slipped slides. Cells were washed three times in 50% formamide/50% $2 \times$ SSC for 5 min, $2 \times$ SSC for 5 min and $2 \times$ SSC/0.1% IGEPAL for 5 min, all at 46°C prior to drying and staining with Rhodamine and Oregon Green-labeled secondary antibodies, washing three times with $1 \times PBS/0.5\%$ Tween 20 and mounting in Vectashield (Vector Laboratories, Burlingame, CA, USA) containing 5 µg of DAPI.

Microscopy

FISH slides were analyzed on a Zeiss Axioplan 2 fluorescence microscope (Carl Zeiss, Inc., NY, USA) equipped with a digital camera. The microscope and peripherals were controlled by a computer-running iVision (Scanalytics, Vienna, VA, USA) software. Images were taken from the top of the specimen, and then repeated at $0.5-1 \mu m$ sections through the depth of the tissue. Each image stack was digitally de-convolved to remove out-of-focus light using HazeBuster software (Vaytek, Fairfield, IA, USA). Following haze removal, image stacks were merged and stacked to create a two-dimensional image representing all of fluorescence within the section. Images were taken using a $100 \times oil$ objective with a $1 \times zoom$. FISH spots were counted as associated if the signals were touching or overlapping. All other red and green signal distances were counted as not associated.

CHRNA7/Chrna7 RT-PCR analysis

Seven classic Rett syndrome, 11 autism and 20 typically developing (controls), post-mortem brain cortices were obtained through the Autism Speaks-supported Autism Tissue Program (ATP) from the Harvard Brain Tissue Resource Center (HBTRC) and the University of Maryland Brain Tissue Resource Center or from the University of Miami Brain and Tissue Bank for Neurodevelopmental Disorders (Supplementary Material, Table S3). For mice, whole brains were isolated from *Mecp2*-deficient B6.129P2(C)-*Mecp2*^{tm1.1Bird}/J mice.

RNAs were isolated using TRIzol[®] Reagent (Invitrogen) from Brodmann area 9 of the frontal cortices. To remove DNA contamination, the total RNA was treated with DNaseI (New England Biolabs, Ipswich, MA, USA) according to the manufacturer's instructions. Single-stranded complementary DNA (cDNA) was synthesized using Quantitect Reverse Transcription Kit (Qiagen). For each reaction, a tube without reverse transcriptase was used as a control for genomic DNA contamination. Primers were designed to cross an intron or span intron/exon boundaries to further limit the effect of possible genomic DNA contamination using (http://frodo.wi.mit.edu/primer3/) Online Primer3 PCR primer design software or Biosearch Technologies Real Time Design software (http://www.biosearchtech.com/rea ltimedesign). PCR amplification of cDNA was performed using 200 nM primer concentration and Express Sybr[®] Green ER Universal Master mix (Invitrogen). Cycling conditions were 20 s at 95°C followed by 40 cycles of 3 s at 95°C and 30 s at 60°C. PCR amplification was performed using the Mastercycler[®] Ep realplex (Eppendorf, Hamburg, Germany) and crossing points were analyzed using Realplex software (Eppendorf). For each reaction, a well without reverse transcriptase and a well without control cDNA were amplified to evaluate genomic DNA contamination, non-specific product formation or other contamination. All samples were normalized to GAPDH expression using the comparative CT method (Applied Biosystems, Foster City, CA, USA) to measure fold change relative to calibrator, following normalization to GAPDH. Melting curve analysis was also performed to determine homogenous product formation and to detect non-specific products.

Student's *t*-test was performed to verify that there were no differences in the ages of disease groups prior to running the one-way analysis of covariance (ANCOVA). Statistical analyses were performed using Mann–Whitney U and one-way ANCOVA tests. All *P*-values were considered significant if the associated value was <0.05.

Primers employed for QRT-PCR

Human *CHRNA7*—forward: GACGTGGATGAGAAGAAC CAA; reverse: TGGGAAACGAACAGTCTTCA. Human *GAPDH*—forward: TGAACCATGAGAAGTATGACAAC; reverse: GTCCTTCCACGATACCAAAG.

Mouse *Chrna7*—forward: TGGGCATTGCCAGTATCT; reverse: AAAGGGAACCAGCGTACATC. Mouse *Gapdh* forward: GCGGAGATGATGACCCTT; reverse GCTGAG TATGTCGTGGAGT.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

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Conflict of Interest statement. None declared.

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