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To cite this version:
Robert Feil, Yui Imaizumi. Emerging chromatin structural roles of the methyl-CpG binding protein MeCP2. Epigenomics, Future Medicine, 2021, 13 (6), pp.405-409. 10.2217/epi-2021-0053. hal-03434520

HAL Id: hal-03434520
https://hal-cnrs.archives-ouvertes.fr/hal-03434520
Submitted on 18 Nov 2021

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Emerging chromatin structural roles of the methyl-CpG binding protein MeCP2

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“Irrespective of the precise underlying mechanisms, the novel studies clearly establish a role for MeCP2 in the architecture and compaction of heterochromatin, and reveal likely cooperation with HP1 proteins and major-satellite RNAs.”

First draft submitted: 3 February 2021; Accepted for publication: 14 February 2021; Published online: 9 March 2021

Keywords: DNA methylation • heterochromatin • MBD proteins • MeCP2 • protein condensates

DNA methylation is an essential epigenetic modification, present in mammals predominantly at cytosine residues that are followed by guanines. In somatic cells, this so-called CpG methylation is found at high levels all along the genome. Different readers have evolved that bind to the methylated DNA and influence chromatin and gene expression. Methyl-CpG binding domain (MBD) proteins constitute the largest family of such proteins, and share a conserved MBD [1]. Extensive research has shown that their binding patterns evolved to facilitate appropriate gene expression levels in the context of highly methylated genomes [2].

Exciting recent studies on the best-studied MBD protein, MeCP2, evoke a new role besides transcriptional repression. They show that MeCP2 contributes to the 3D structuration and compaction of heterochromatin as well [3–6]. Below, we present these newly emerging insights and discuss their possible relevance for neurological disorders in humans that are caused by genetic mutations in the MeCP2 gene.

The MeCP2 gene is positioned on the X chromosome and is mutated in almost all patients with Rett syndrome (RTT, OMIM 312750), a progressive and devastating autism-spectrum disorder that affects about one in 10–15,000 female births [7,8]. Another genetic neurological syndrome is caused by MeCP2 duplication, and is found mostly in boys [8]. Constitutive and brain-specific knockout (KO) studies reproduced the clinical phenotype of RTT in mice, and highlighted the essential role of Mecp2 in brain functions [7,9]. Although MeCP2-deficient neurons are functionally abnormal, they do not die. Remarkably, it was shown that the defective brain functions can be largely reversed by reactivating the Mecp2 gene in the KO mice [10,11]. Thus, although RTT affects neuronal functions, it is not a developmental syndrome.

Mecp2 produces two similar isoforms that both comprise the MBD but differ slightly in the N-terminus. In the brain, this very highly expressed protein binds not only methyl-CpG but also the methyl-CAC tri-nucleotide motif, which is globally rare in the genome, but more abundant in this than other tissues [12,13]. Extensive research has shown that the protein’s primary role is to recruit different proteins involved in transcriptional repression. Among these are the HDAC-containing NCoR1/2 and SIN3A complexes [7]. The recruitment of these repressive complexes by MeCP2 requires the NCoR1/2 interaction domain (NID) (Figure 1) [7]. Particularly in neurons – where its expression is highest – MeCP2 influences the transcription of many genes in a subtle manner, leading to moderate gains and losses of expression. At genes that show preferential MeCP2 recruitment, its depletion enhances gene expression, indicating that MeCP2’s primary role is to dampen gene expression [7]. It had been suggested that MeCP2-binding to genes could also control RNA splicing but this is not supported by recent studies, which also do not provide evidence for a prominent role of methylation in alternative splicing [14]. Besides the MDB and NID, other parts of MeCP2 are also highly conserved and thus could also be functionally important.
Pericentromeric heterochromatin comprises satellite repeat DNA and is particularly rich in methyl-CpG dinucleotides. Immunofluorescence staining on fixed mouse cells shows a marked focal accumulation of MeCP2 in heterochromatic foci. What could be the role of MeCP2 at these gene-poor heterochromatic domains? To address this question, Linhoff et al. developed ChromAtin, a quantitative approach that combines IF staining and low-temperature DNA FISH with a method that defines nuclear compartments using array tomography imaging [4]. They carefully compared normal and MeCP2-deficient neurons. Remarkably, the lack of MeCP2 increased DAPI staining at chromocenters, indicating higher compaction levels within pericentromeric heterochromatin. MeCP2 depletion also correlated with increased IF staining of histone H4 lysine-20 tri-methylation (H4K20me3) [4], a modification that was known to control chromatin compaction [15].

Following the discovery that MeCP2 influences chromatin compaction, several groups set out to perform further studies. Applying high-resolution confocal and electron microscopy, Ito-Ishida et al. explored heterochromatic foci of neurons in Mecp2 KO versus WT mice [3]. In hemizygous Mecp2 +/− females, about half of the neurons do not express the WT Mecp2 allele, due to X-inactivation (Mecp2 maps to the X-chromosome). In these mice, the MeCP2-deficient neurons showed a markedly stronger DAPI staining of constitutive heterochromatin than the MeCP2-expressing neurons. MeCP2-depleted heterochromatin foci were more spherical in shape as well, providing another indication that MeCP2 ‘loosens’ heterochromatin. Concordantly, when the authors overexpressed MeCP2 in neurons, heterochromatin foci showed reduced DAPI staining and were no longer spherical in shape.

Another remarkable recent study, by Li et al., reported that MeCP2 has physico-chemical properties compatible with the induction of liquid-liquid phase separation (LLPS) condensates in vitro, and that this propensity of
the protein to form droplets is enhanced by the presence of methylated DNA [6]. By studying truncated forms of the protein, they discovered that the formation of condensates required a large intrinsically disorganized region (IDR) covering the C-terminal portion of the protein. Generating amino-acid alterations, the authors showed that basic residues within this IDR-2 domain (Figure 1) are essential to the process. Whether the C-terminus exerts phase separation condensates in living cells as well remains unclear, with the confounding issue that IDR-2 deletion removes the NID domain as well. Interestingly, however, an RTT-associated nonsense mutation that removed the entire IDR-2 gave strongly reduced accumulation of MeCP2 within heterochromatic foci. Concomitantly, reduced incorporation of a fluorophore-tagged HP1α was observed within heterochromatin in these mutant cells. In vitro, concordantly, phase-separated MeCP2 droplets incorporated tagged-HP1α when this was added to the tube, showing that these two proteins can form joint condensates [6]. These observations are interesting given that also HP1α displays phase separation properties – at least in vitro – and given that this heterochromatin component restricts the compaction levels of heterochromatin in somatic cells [16,17].

Further studies are required to unravel the molecular details, for instance of how MeCP2 and the HP1 isoform HP1α are linked within heterochromatin. The data so far evoke a model in which local methyl-CpG (and methyl-CAC) concentration triggers the initial MeCP2 enrichment at heterochromatin, which requires the MBD domain (Figure 1). Further protein condensation is achieved through the LLPS-like properties of the IDR-2. The phase separation of MeCP2, HP1α and yet other heterochromatin components may be facilitated by the focally high concentration of nucleosomes as well [18]. In neurons, MeCP2 condensation within heterochromatin correlates with focal HP1α enrichment, reduced H4K20me3 and with lower compaction levels (Figure 1). Whether MeCP2 condensation also influences HP1β levels would now be interesting to explore, given that this HP1 isoform is functionally associated with H4K20me3 and with increased chromatin compaction levels [16].

What could be the role of the intriguing MeCP2-mediated heterochromatin configuration? Without the MBD-induced recruitment of MeCP2 to its methylated DNA, a further accumulation – possibly through LLPS – would not take place at heterochromatin [6]. Molecular details are yet insufficient, however, to understand the dampening effects of MeCP2 condensates on repressive histone methylation and compaction levels.

A missing piece of the puzzle could be RNA. In recent studies, different nuclear RNAs were reported to influence chromatin structure and the formation of protein condensates in distinct nuclear compartments [19], including at pericentromeric heterochromatin [20]. Within this context, Fioriniello et al. recently explored the importance of RNA for heterochromatic MeCP2 condensates in neural cells [5]. After a brief RNase-A treatment, which completely ablated the major satellite RNA, strongly-reduced levels of MeCP2 were observed at heterochromatin foci. A similar, but smaller, effect was seen in experiments in which the authors used RNA-ablating gapmers directed against satellite repeat RNA [5]. In Mecp2 KO cells, conversely, major-satellite RNAs (transcribed in the forward orientation) were strongly reduced at heterochromatin. Dissecting this further in missense mutations, the authors found that the MBD and the NID-comprising transcriptional repression domain are both required for the targeting of satellite-RNA to heterochromatin. Combined with additional experiments, the authors propose that MeCP2 interacts with major-satellite RNAs and contributes to their retention at heterochromatin. The major-satellite repeat RNAs, in turn, contribute to focal MeCP2 condensation, as they do for other heterochromatin components [20,21], and this could involve dose-dependent threshold effects.

The recent studies generated intriguing insights into MeCP2 and heterochromatin, but also evoke new questions. Notably, it remains unclear how important the structural effects of MeCP2 on heterochromatin are in relation to RTT. Several of the mutations that had chromatin-structural effects are causally involved in RTT [6], but this might be for other reasons not related to heterochromatin, including possible effects on MBD or NID functions or on the protein’s stability. In Mecp2 KO mice, the different RTT-like neurological defects can be prevented or reversed by expressing a truncated mini-protein that comprises just the MBD and NID domains [22]. This suggests that the neuronal functions mediated by MeCP2 are possibly not linked to the protein’s structural effects on heterochromatin, in case the latter are indeed mediated by regions other than the MBD and NID [6].

Irrespective of the precise underlying mechanisms, the novel studies clearly establish a role for MeCP2 in the architecture and compaction of heterochromatin, and reveal likely cooperation with HP1 proteins and major-satellite RNAs. Future research will undoubtedly provide other pieces of this rather complicated puzzle.

Acknowledgments
We apologize to colleagues whose work was not cited due to space limitations. We thank D. Ll`eres for expert advice and proof-reading.
Financial & competing interests disclosure
Research in the R Feil laboratory is supported by the Agence Nationale de la Recherche (ANR), through grant ANR-18-CE12-0022-02 (‘IMP-REGULOME’) and Labex EpiGenMed (ANR-10-LABX-12-01), and the Institut National de Cancer (INCa, grant PLBIO18-094). Y Imaizumi acknowledges Fellowship funding from TOYOBO Biotechnology Foundation, Japan (2019-01). The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript.

No writing assistance was utilized in the production of this manuscript.

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