A Role for Methyl-CpG Binding Domain Protein 2 in the Modulation of the Estrogen Response of \( pS2/TFF1 \) Gene

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Abstract

Background: In human Estrogen Receptor \( \alpha \) (ER\( \alpha \))-positive breast cancers, 5’ end dense methylation of the estrogen-regulated \( pS2/TFF1 \) gene correlates with its transcriptional inhibition. However, in some ER\( \alpha \)-rich biopsies, \( pS2 \) expression is observed despite the methylation of its TATA-box region. Herein, we investigated the methylation-dependent mechanism of \( pS2 \) regulation.

Methodology/Principal Findings: We observed interplay between Methyl-CpG Binding Domain protein 2 (MBD2) transcriptional repressor and ER\( \alpha \) transactivator: (i) the \( pS2 \) gene is poised for transcription upon demethylation limited to the enhancer region containing the estrogen responsive element (ERE); (ii) MBD2-binding sites overlapped with the methylation status of the \( pS2 \) 5’ end; (iii) MBD2 depletion elevated \( pS2 \) expression and ectopic expression of ER\( \alpha \) partially overcame the inhibitory effect of MBD2 when the ERE is unmethylated. Furthermore, serial chromatin immunoprecipitation assays indicated that MBD2 and ER\( \alpha \) could simultaneously occupy the same \( pS2 \) DNA molecule; (iv) concomitant ectopic ERE expression and MBD2 depletion resulted in synergistic transcriptional stimulation, while the \( pS2 \) promoter remains methylated.

Conclusions/Significance: MBD2 and ER\( \alpha \) drive opposite effects on \( pS2 \) expression, which are associated with specific steady state levels of histone H3 acetylation and methylation marks. Thus, epigenetic silencing of \( pS2 \) could be dependent on balance of the relative intracellular concentrations of ER\( \alpha \) and MBD2.

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Introduction

Global loss of DNA methylation and localized CpG island hypermethylation is a common characteristic of cancer cells [1–3], leading respectively to aberrant ectopic gene activation or inversely to gene silencing. The \( pS2 \) gene (also called \( TFF1 \)) has been identified by differential screening of a cDNA library from the human breast cancer cell line MCF7 [4]. In this cell line, its transcription is directly controlled by estrogens [5] and an estrogen responsive element (ERE) has been identified at nt positions 405 to −393, from transcription start site [5]. In breast tumors, expression of the \( pS2 \) gene is correlated with the presence of estrogen receptors (ER), and it had been suggested that \( pS2 \) expression increases cell proliferation and tumor cell survival [6,7]. Analysis of breast cancer biopsies or microdissected cells from formalin-fixed breast tissues has shown that \( pS2 \) is hypomethylated in sub-classes of breast cancers [8,9].

We have previously shown [8] that the hypomethylation of the CCGG site close to the \( pS2 \) ERE correlates with its expression in human breast cancer biopsies. Southern blots performed with methylation sensitive enzymes and bisulfite sequencing have indicated that the breast tumors analyzed exhibited different DNA methylation patterns at the 5’ end of \( pS2 \) [8]. Biopsies can display either methylated, unmethylated and partially methylated 5’ end \( pS2 \) sequences at CpGs analyzed (nt positions −84 to +16) [8]. These observations prompted us to investigate the methylation-linked mechanisms of \( pS2 \) gene repression and the potential involvement of DNA methylation in its response to estrogen stimulation.

In mammals, mechanisms implicated in the generation of a repressive state of chromatin associated with methylated DNA sequences have been investigated for over 20 years [10–13]. Pioneering studies led to the discovery of the Methyl-CpG binding domain (MBD) proteins family [14], which mediate DNA methylation-dependent gene silencing. The five \( \textit{ bona fide } \) MBD proteins, MeCP2, MBD1, MBD2, MBD3, and MBD4, share a canonical MBD. Biochemical and genetic analyses of these proteins have provided evidence of a direct link between DNA methylation and repressive chromatin architecture. MeCP2, MBD1 and MBD2 proteins bind to methylated DNA and recruit different histone deacetylase (HDAC)- and histone methyltransferase (HMT)-containing complexes that control chromatin compaction and gene silencing [15–17]. Mammalian MBD3, which lacks a functional MBD, does not recognize methylated DNA but is part of the histone deacetylase and chromatin remodeling Mi2/NuRD complex [18–20]. The last member of
This protein family, MBD4, is a thymine glycosylase primarily involved in DNA repair [21].

The involvement of MBD proteins in gene imprinting [22], X inactivation [23], and transcriptional silencing of genes possessing hypermethylated CpG islands in cancer cells [2,3] is now well documented. However, in contrast to the situation observed for DNMT-deficient mice, which either fail to develop or else die shortly after birth [24], the loss of MBD proteins, with the exception of MBD3, does not result in dramatic phenotypes [17], suggesting that MBD proteins deficiency causes subtle gene-expression changes.

Although the involvement of MBD proteins in gene silencing is well established, new facets regarding the links between DNA methylation, MBD proteins and gene transcription are emerging. For instance, it has been reported that DNA methylation in the body of genes can alter chromatin structure and reduce transcriptional elongation [25]. In parallel with the above findings, we have shown that the association of MBD2 with a methylated CpG island located downstream of the promoter region reduces the transcription of NBR2 gene [26].

The density of methylation seems to be an important parameter in the MBD proteins-dependent repression. Several years ago, using in vitro methylated plasmids, it had been shown that the density of methylated CpG and promoter strength modulate transcriptional repression mediated by McCP1 complexes containing MBD2 [27]. Furthermore, analysis of the transcriptional activity of patch-methylated plasmids microinjected into Xenopus oocytes has suggested a competition between transactivators and MBD proteins for the establishment of an open conformation [28]. All together, these data suggest that transcriptional repression mediated by DNA methylation is a consequence of a cross-talk between methylated CpG density, MBD proteins and transactivators.

To further explore the relative roles of methylated CpG patterns and the competition between transactivators and MBD proteins to influence or modulate gene transcription, we here investigate the expression of the estrogen-regulated pS2/TFF1 (TFF1, Tissue factor Factor 1) gene [29] in cell lines exhibiting different DNA methylation patterns at its 5' end, unmethylated, regionally methylated, and fully methylated. In these cell lines, it was possible to manipulate artificially the Estrogen Receptor-α (ERα), the natural transactivator of pS2, and MBD protein levels, and therefore, use them to determine the contribution of these proteins to pS2 expression.

**Results**

**Correlation between methylation patterns and pS2 transcriptional repression**

Expression of pS2 is driven by a complex promoter containing a promoter/enhancer region responsive to estrogens, EGF, a phorbol ester tumor promoter, c-Ha-ras oncoprotein, and c-jun protein [33]. Specifically, the 5' end of pS2 possesses an estrogen-responsive element (ERE), conferring potential estrogen inducibility. In the ERα-rich MCF7 cell line, it has been shown [34] that proteins present on the pS2 promoter in the absence of estradiol (E2) include basal transcriptional factors, active polymerase II, certain HATs and HMTs. This basal transcriptional activity implies steroid-independent expression of pS2. Moreover, the 5' end of pS2 (nt positions –464 to +314) is included in a CpG-poor region, G + C = 0.54%, CpG observed / CpG expected = 0.35. A correlation between the methylation status of the pS2 promoter region and its expression has been observed in human tissues and the breast cancer cells lines [8,35]. Experimental evidences are also in favour of a role of DNA methylation in the repression of pS2 transcription. In the ERα-rich MCF7 cells, pS2 is unmethylated and transcriptionally active while in the ERα-negative MDA MB231 cell line the 5' end of pS2 (nt positions –665 to +17) is fully methylated and the pS2 gene is silenced, [35]. We have extended this analysis to the down stream region of pS2 promoter, since it had been suggested that DNA methylation of the regions adjacent to a promoter region may affect transcription [26,36,37].

The methylation status of the 5’ end of pS2, from nt positions –464 to +294, was investigated in MCF7 cells expressing pS2 at a high level and MDA MB231 a pS2 negative cell line (Figure 1A). Bisulphite conversion of DNA and sequencing of cloned PCR fragments, indicated that this region is unmethylated in MCF7 and fully methylated in MDA MB231 cells (Figure 1B). A screening failed to detect a human breast cancer cell lines exhibiting intermediate methylation patterns similar to that observed in breast cancer biopsies by Southern blot experiments.

![Figure 1. pS2 gene expression and DNA methylation patterns in MCF7, HeLa and MDA MB231 cells. (A) The expression of endogenous pS2 gene in MCF7, HeLa and MDA MB231 cell lines. pS2 mRNA levels were monitored by relative RT-PCR. Briefly, pS2 transcripts were simultaneously amplified with β-actin transcripts as a loading control and expression standard. (B) Methylation patterns at CpG sites of pS2 5’ flanking sequence from nt positions –464 to +314 in MCF7, HeLa, and MDA MB231 cell lines. A schematic representation of the human pS2 gene is shown. The transcription start site is indicated by a black arrow. Black box, AP1 site; dark-grey box, Estrogen-Responsive Element (ERE); light-grey box, TATA-box; hatched boxes, pS2 exons. The studied region (from nt positions –464 to +314) is presented on an expanded scale. This region contains 20 CpG sites, represented by white circles. The bisulphite-sequencing status of this 5’ pS2 region in MCF7, HeLa and MDA MB231 cells (number of analyzed clones, n = 10) is represented. Each line corresponds to a single DNA template molecule. Black and open circles represent methylated and unmethylated CpGs, respectively.

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Nevertheless, some of these patterns, associated with low level of pS2 transcripts (Figure 1A), are very similar to that observed in HeLa cells [8]. Thus, these cells were chosen for further analysis. HeLa cells exhibit an intermediate DNA methylation pattern, the CpGs spanning the −464 to −84 region, which includes the ERE are unmethylated, while the TATA-box region is methylated (Figure 1B). These data confirm the inverse correlation between pS2 expression and the density of the methylation of its 5’end, suggesting that the methylation patterns around the transcription start site impact in the activity of pS2 promoter in these cells.

**Specific binding of MBD2 to the methylated pS2 promoter**

Among the proteins involved in the methylation-dependent repression of transcription, MBD proteins seem to play a major role. Therefore, we assessed the presence of MBD proteins on pS2 promoter by chromatin immunoprecipitation (ChIP) assays using antibodies directed against MBD1, MBD2 and MeCP2.

Representative experiments from at least three independent assays for each antibody are shown in Figure 2A. As a control, the fractions immunoprecipitated with a non-MBD protein-specific antibody (anti-mouse IgG) were also analyzed. In order to determine pS2 DNA fragment enrichment in MBD immunoprecipitated fractions, a dose-dependent and quantitative (Q-PCR) amplifications (Figure 2A, B) using an equal quantity of DNA (0.5 ng) per PCR assay, were performed with each fraction obtained from the ChIP procedure. We focused this analysis on the pS2 methylated promoter region, shared by HeLa and MDA MB231 cells (nt positions −11 to +292).

In the pS2-methylated cells, MDA MB231 and HeLa, when antibodies against MBD2 were used, the amount of pS2 promoter per ng of total DNA in immunoprecipitated fraction (Figure 2A,
“bound”) was greater than in the input, or in the non-retained fractions (Figure 2A, “input”, “unbound”, and “IgG”), indicating that this methylated region is immunoprecipitated by anti-MBD2 antibodies. In contrast, ChIP assays with anti-MeCP2 or anti-MBD1 antibodies led to a depletion of this DNA segment in the bound fractions (Figure 2A). Western blot analysis, using antibodies directed against MeCP2 and MBD1, produced a signal of the expected sizes [14, ~85 kDa and ~70 kDa, respectively (data not shown), indicating that both proteins are expressed in MDA MB231 and HeLa cells. As an additional control of MBD2-ChIP assays, we also amplified the −11 to +292 pS2 region from MCF7-chromatin immunoprecipitated by anti-MBD2 antibodies (Figure 2A). As expected, no enrichment was observed in the bound fraction, since this region was unmethylated in the MCF7 cell line (Figure 2A). Taken together, these data strongly suggest that MBD2 binds selectively and specifically the methylated region of this promoter.

High resolution MBD2 binding profiles analysis of pS2 promoter indicates that MBD2 specifically binds the methylated pS2 promoter region and does not spread to the unmethylated ERE in HeLa cells

In HeLa cells, the bimodal methylation status of the pS2 5’ end suggests that only some regions would actually be bound by MBD2. The unmethylated region containing the ERE (nt positions −405 to −393) is very close to the methylated TATA-box region (beginning at nt -9). ChIP experiments are not appropriate to discriminate between these two regions, as they are below the limit of resolution of the assay. Standard sonication of crosslinked chromatin leads to 300–500 bp DNA fragments and attempts to reduce its length (100–200 bp) resulted in a loss of efficiency in immunoprecipitation. Therefore, to precisely map the MBD2 binding sites we used a high-resolution method based on a ChIP-on-chip approach (Chatagnon et al., manuscript in preparation). DNAs obtained from HeLa cell chromatin immunoprecipitated by anti-MBD2 antibodies were hybridized on an Affymetrix Human Promoter 1.0 R Array (ChIP-on-chip).

ChIP-on-chip experiments indicated that MBD2 associated specifically the region containing the methylated pS2 TATA-box, where a strong positive signal (red bars) is observed, while the region containing the unmethylated ERE is devoid of MBD2. Thus, the positive signals for MBD2 binding parallels the methylation status of the pS2 5’ end and indicates that MBD2 does not spread outside the methylated region on pS2 promoter (Figure 3A). As a control, results obtained for a previously identified MBD2 free promoter [26], BRCA1, are also shown on Figure 3B. Consistent with previous findings, no MBD2 positive signal was observed in the region spanning the nucleotides −1000 to +1000 of BRCA1 (Figure 3B).

MBD2 acts as a methylation-dependant transcriptional repressor of pS2 transcription

The correlation between levels of pS2 expression and the presence of MBD2 in the TATA-box region argues in favour of a repressive effect of MBD2 on pS2 transcription. To examine this further, we depleted cells from MBD2 by transient transfection of siRNA targeted MBD2 transcripts. Quantitative competitive RT-PCR assays indicated a significant reduction of the MBD2 mRNA

Figure 3. ChIP-on-chip analysis of MBD2-binding sites on pS2 5’ end regionally methylated in HeLa cells. (A) Array peaks on pS2 5’ end of MBD2 log2 signal ratio (MBD2 / Input) values are shown below the Affymetrix’ Integrated Genome Browser (IGB) window. Red bars, MBD2 binding sites; yellow bars, MBD2 free sites. Genes are transcribed from right to left. pS2 methylation status from nt positions −464 to +314 is shown by a diagram. “pS2 ERE fragment” and “pS2 promoter fragment” analyzed by PCR following ChIP are represented by a white box. (B) BRCA1 5’ end viewed as a MBD2 free control.

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level (by 87–93%) (Supplemental Figure S1A). Moreover, in HeLa cells, western blot analysis also revealed a dramatically lower abundance of MBD2 proteins in the MBD2 siRNA-treated cells compared with control cells (Supplemental Figure S1B). Furthermore, the expression of MeCP2 and MBD1 was not different in MBD2 knockdown HeLa cells than in wild-type or mock-treated cells (Supplemental Figure S1B).

In the pS2-fully methylated cells MDA MB231, MBD2 depletion (about 90%), did not induce pS2 expression (Supplemental Figure S1C). However, in MBD2 siRNA-treated HeLa cells, pS2 expression is stimulated approximately 3-fold (Figure 4), while the methylation level of pS2 TATA box region and ERα expression remain unaffected (data not shown). In MBD2 siRNA-treated HeLa cells, transient expression of an Mbd2 cDNA, refractory to siRNA-mediated decay [26], shifted down the pS2 mRNA level (Figure 4). Additional dose of Mbd2 in HeLa cells, containing normal levels of MBD2 proteins, did not affect pS2 expression (Figure 4). Thus, the amount of MBD2 protein is not a limiting factor in the transcriptional repression of pS2 in these cells. In MCF7 cells, quantitative RT-PCR showed that the level of pS2 expression is unaffected by MBD2 depletion, indicating that MBD2 siRNA did not elevate pS2 expression by an off-target effect since pS2 is not bound by MBD2 in these cells (Supplemental Figure S1C).

ERα only bound pS2 ERE when unmethylated in cell lines

To determine whether ERα can be recruited on the pS2 ERE sequence when this region or the adjacent region is methylated, we artificially manipulated the level of ERα in MDA MB231 and HeLa cells, which are deficient in this protein.

ChIP assays, using anti-ERα antibodies, were performed from HeLa and MDA MB231 cells transiently transfected with the vector HEG0 encoding ERα. ERα-rich MCF7 cells were used as a positive control and untransfected HeLa and MDA MB231 cells as negative controls. As expected, ChIP assays performed from MCF7 chromatin indicated that the amount of pS2 DNA per ng of total DNA in immunoprecipitated fraction (Figure 5, “bound”) is higher (about 8-fold, Q-PCR assays) than in input, or non-retained fractions (Figure 5, “input”, “unbound”, and “IgG”) while no enrichment in pS2 sequence was observed in immunoprecipitated fraction obtained from untransfected HeLa and MDA MB231 cells (Figure 5). All together, these data indicate that anti-ERα antibodies specifically precipitated chromatin bound by ERα.

Figure 4. MBD2 specifically and directly represses pS2 transcription. Real time RT-PCR analysis of pS2 transcripts in HeLa and MBD2-depleted HeLa cells (HeLa cells pretreated for 72 h with MBD2 siRNA) transfected with an MBD2 vector expressing a transcript resistant to RNAi (pRev-MBD2 vector) or with an empty basic vector pGL3. Transcriptional expression of pS2 was analyzed 24 h after transfection. The fold change of pS2 expression was calculated from the relative pS2 mRNA in pRev-MBD2-transfected cells compared to pGL3-transfected cells. Values are presented as the mean ± standard deviation of at least three independent transfection experiments. A significant difference between the two cell groups is represented by an asterisk * (P<0.05).

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Figure 5. ERα only associates hypomethylated ERE region of pS2. Representative experiments of ERα ChIP assays in ERα-rich MCF7 cells, in ERα-negative HeLa and MDA MB231 cells, and in HeLa and MDA MB231 expressing the vector HEG0 encoding ERα (HeLa::ERα, and MDA MB231::ERα). ChIP assays were performed as described in Figure 2. The position of the “pS2 ERE fragment” analyzed by PCR are represented on the pS2 5’ end schema.

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In HeLa cells, transient expression of the vector HEG0 coding for ERα, leads to enrichment in pS2 sequence in the immunoprecipitated fraction (Figure 5). This enrichment (about 10 fold, Q-PCR assays) was comparable to that observed in MCF7 cells, indicating that ERα is efficiently recruited on the pS2 ERE site in HeLa cells. In contrast, in MDA MB231 cells, despite ectopic expression of ERα, pS2 sequence was not selectively immunoprecipitated by the anti-ERα antibodies (Figure 5). Thus, ERα does not bind the methylated pS2 ERE in MDA MB231 cells, suggesting that full DNA methylation induces chromatin changes that prevent ERα binding.

**Ectopic expression of ERα enhances pS2 gene expression only when its 5’ end region is partially methylated, in ERα negative cells**

The ERE of the pS2 promoter can act as a strong enhancer in the presence of E2 and ERα [29,33]. To analyze the potential antagonistic activity of MBD2 and ERα on pS2, ectopic expression of ERα was induced in MDA MB231 and HeLa cells.

Transient expression of ERα elevated pS2 expression by a 4-fold (Figure 6), in HeLa cells. This stimulation by ERα of pS2 transcription did not affect the methylation status of HeLa pS2 promoter (data not shown). ERα stimulation of pS2 expression was fully reversed by antiestrogen (4-hydroxytamoxifen, OHT) treatments (Figure 6), while the basal level of pS2 expression was conserved. These results indicate the existence of estrogen-dependent transactivation of pS2 in HeLa cells. As expected, pS2 expression was not induced in MDA MB231 cells (data not shown), since ERα did not bind pS2-ERE sequence in this cell line (Figure 5). These results suggest that demethylation of ERE region allows the estrogen response of pS2 in HeLa cells.

**MBD2 is not displaced from the pS2-methylated promoter region by ERα transactivation, in HeLa cells**

HeLa cells expressing the vector encoding ERα were used to identify the proteins bound to the 5’ end of pS2. Dose-dependent and quantitative PCR amplifications of each fraction obtained from the ChIP procedure were performed. These assays showed that MBD2 was still present on the methylated region of pS2 promoter (Figure 7A), and the enrichment in the bound fraction was not modified by the presence of ERα on the ERE (Figure 7A). These results provide evidence that ERα can overcome, at least partially, the inhibitory effect of MBD2 binding to pS2 promoter and imply that both proteins can occupy the 5’ end of the pS2 gene.

To address this matter, we performed serial ChIP assays. A first round of immunoprecipitation was carried out with an anti-ERα antibody from HeLa cells transiently transfected with the vector HEG0 encoding ERα. Then, immunoprecipitated cross-linked DNA-protein complexes were isolated and subjected to reimmunoprecipitation using antibodies directed against MBD2. PCR amplification of pS2 promoter region from the fraction reimmunoprecipitated with anti-MBD2 antibodies gave a positive signal.
Synergic activity of MBD2 depletion and ectopic ERα expression on pS2 transcription, in HeLa cells

The opposite effects of MBD2 and ERα proteins on pS2 expression suggest an antagonistic action between these two transcriptional regulators, in HeLa cells. To investigate this possibility, the concentrations of MBD2 and ERα proteins were artificially manipulated in these cells. After MBD2 depletion mediated by RNA interference, ectopic ERα expression resulted in a dramatic (approximately 31-fold) enhancement of pS2 mRNA concentration, approaching the level to that observed in MCF7 (Figure 8A, B). Thus, pS2 responses to ERα activation (4-fold increase) and MBD2 depletion (3-fold decrease) are not additive and suggest a cross-talk between these two transcriptional regulators. Concomitant exposure to OHT knocked down pS2 expression to the level observed in HeLa cells transfected by MBD2 siRNA alone, (about 3-fold) when compared with control HeLa cells (Figure 8A, B). It should be noted that pS2 transcription cannot be induced by concomitant ectopic expression of ERα and MBD2 depletion when its 5’ end is fully methylated, as observed in MDA MB231 cells (data not shown). In cells exhibiting a bimodal methylation profile as HeLa cells, a synergic activation was observed. Thus, the binding of MBD2 to the methylated TATA-box of pS2 reduces but does not abolish pS2 response to ERα, suggesting that as a result of regional demethylation, pS2 is poised for transcription.

Post-translational modifications of histone H3 are associated with pS2 expression induction

To get further insight on the mechanism involved in the opposite effect of MBD2 and ERα on pS2 transcription, we investigated histone modifications in HeLa cells. It is well known that both proteins, MBD2 and ERα, regulate the transcription by the recruitment of chromatin remodeling complexes [15,38]. Importantly, ERα has been shown to interact with several coactivators with histone acetyltransferase activity (CBP, p300, p/CAF and the members of p160 family) [38], or histone demethylase (LSD1) [39]. Conversely, MBD2 recruits corepressors with histone deacetylase activity (Mi2/NurD) [17]. Histone H3 acetylation (H3Ac) and histone H3 lysine 9 (H3K9) trimethylation chromatin marks have been the subject of intense investigation during the past few years and appear to be associated with active and silent promoters, respectively.

In our study, ChIP assays indicated that ERα pS2 stimulation was associated with increase in histone H3 acetylation (~2.5 fold) and enhanced the demethylation of H3K9 (~500 fold) at pS2 promoter, when compared with wild type HeLa cells (Figure 9). Moreover, in HeLa cells, the synergistic activity of MBD2 depletion and ectopic ERα expression on pS2 transcription led to a stronger induction of histone H3 acetylation (~28 fold) at pS2 promoter, while H3K9 methylation was still lowered (~10 fold), (Figure 9). From these findings we conclude that the transcriptional MBD2 repressor and ERα transactivator co-participate to the regulation of pS2 expression by mediating a balance between repressive histone H3 lysine 9 trimethylation and active histone H3 acetylation marks at pS2 promoter. Thus, the repressive effect of MBD2 on the transcription of pS2 mediated by ERα is linked to histone modifications.

Discussion

In cancer tissues and cell lines, transcriptional silencing associated with aberrant methylation of promoter regions is now regarded as an almost universal epigenetic marker of malignant transformation [1–3]. Since the first experiments showing that the MBD2 binds in vivo to the methylated regulatory regions of p16 and p14 and could thereby contribute to gene silencing in colon carcinoma cell lines [40], a body of evidence has accumulated concerning associations between MBD proteins and hypermethylated promoter regions [41–43]. In non-pathological situations, MBD proteins are also directly involved in the repression of
The present study also indicates that ERα only associates the unmethylated pS2 ERE independently of the methylation status of the TATA-box region, since the same level of ERα was observed at the ERE region in MCF7 cells and in HeLa cells transiently transfected with the HEG0 vector encoding this receptor. In contrast, strong expression of ERα upon transient transfection with HEG0 did not lead to ERE binding in MDA MB231 cells, suggesting that full methylation of the 5′ end of pS2 prevents its binding. Although a direct effect cannot be totally excluded, we have previously shown, using electrophoretic mobility assays, that methylated oligonucleotides containing the pS2 ERE are efficiently recognized by ERα [44]. In vivo experiments also suggest an indirect effect of DNA methylation. In the HE5 cell line derived from MDA MB231 that expresses functional ERα, but not pS2, DNase I hypersensitive sites are not modified by ERα expression, while in the ERα and pS2 positive MCF7 cells display hormone-dependent hypersensitive DNase sites at the pS2 ERE region [45]. Thus, the methylation of TATA-box region does not seem to lead to diffusible alteration of chromatin structure, in HeLa cells.

As expected from the analysis of methylation and ERα binding sites profiles, ectopic expression of ERα in MDA MB231 cells did not stimulate pS2 expression. In HeLa cells, ERα induction of pS2 expression was observed suggesting that the demethylation of ERE region seems to be a prerequisite for estrogen-dependent pS2 stimulation. Nevertheless, the level of pS2 transcript remained relatively low when compared with that observed in E2 treated MCF7 cells. The present study indicates that high levels of ERα can at least partially overcome the transcriptional repression mediated by MBD2 without affecting the methylation status of the pS2 promoter. Thus, the binding of MBD2 to the methylated TATA-box of pS2 reduces but does not abolish pS2 response to ERα.

The mapping of MBD2 binding sites at pS2 locus indicated that MBD2 proteins lay downstream the initiation start site. The analysis of the 5′ region of the endothelin receptor B gene in human cell lines shows that extensive methylation closely downstream of the initiation site does not abolish gene expression [46]. However, the impact of intragenic methylation has been studied from transgenes methylated exclusively in a region downstream of the promoter, into a specific genomic site. This methylation pattern induces a close chromatin conformation and decrease transcription levels, suggesting that this epigenetic mark may reduces the efficiency RNA polymerase II elongation [25]. Thus, we cannot exclude that DNA methylation and MBD2 binding downstream pS2 promoter region may impact elongation rate. However, we observed post-translational modifications of histone H3 (acetylation and demethylation) at the promoter region, when pS2 transcription is induced by MBD2 siRNA and/or ERα expression, suggesting that MBD2 depletion influences chromatin conformation at pS2 promoter.

Recently, it has been published that, upon E2 induction, synchronized human cell lines exhibit a cyclical methylation/demethylation of the pS2 ERE region correlated with cyclical binding of transcriptional repressors/activators [47,48], when the 5′ end of pS2 is unmethylated [33,48]. Although the maintenance of DNA methylation patterns begin to be well described, it will also be important to consider mechanisms that enable the removal of these marks to fully comprehend the dynamic behavior of DNA methylation, as suggested by recent reports [47,48]. Nevertheless, the mechanisms and enzymatic activities that are responsible for DNA demethylation in mammals although potentially linked to DNA repair are controversial [47–53].

Transcriptional repression mediated by MBD proteins can be reversed by various mechanisms. In mouse and rat, Mecp2 is

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**Figure 9. Histone H3 marks on pS2 promoter in presence or absence of MBD2 and/or ERα.** HeLa cells, wild type (HeLa), expressing ectopically ERα (HeLa:ERα), and depleted in MBD2 and expressing ectopically ERα (MBD2 KD HeLa:ERα), were subjected to ChIP analysis using anti-histone H3 acetylation (H3Ac) or an anti-histone H3 lysine 9 trimethylation (H3K9me3) antibodies. The pS2 promoter was amplified by real-time PCR from an equal amount (0.5 ng) of total DNA immunoprecipitated by the different antibodies. Relative amounts of H3Ac or H3K9me3 marks were measured by comparing fractions immunoprecipitated by the anti-H3Ac or anti-H3K9me3 antibodies to fractions immunoprecipitated by the anti-histone H3 pan antibody. Each bar represents the mean ± standard deviation.

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directly involved in a depolarization-controlled repression of the brain-derived neurotrophic factor (Bdnf) gene in neurons [54,55]. This gene, associated with the corepressor Sin3a, binds to a CpG-poor methylated region of Bdnf promoters. Upon the initiation of Ca\textsuperscript{2+} signaling, Mecp2 becomes phosphorylated and is liberated from the promoter as Bdnf is activated. Mbd2 is also able to repress CpG-poor methylated promoter. In Mbd2 null mice, ectopic expression of interleukin-4 (Il4) disrupts T-helper cell differentiation, suggesting that Mbd2 is a transcriptional repressor of Il4 in naive T cells. Indeed, in naive wild type T cells, overexpression of the transcription factor GATA-3, normally required for Il4 expression, displaces Mbd2 from the promoter and activates Il4 transcription [56].

The interplay between ER\textsubscript{x} and MBD2 in pS2 transcription indicates that the partial reversion of transcriptional repression mediated by MBD2 does not necessarily involve the displacement of this repressor from its CpG-poor promoter region. In the model studied, the amounts of pS2 gene immunoprecipitated by anti-MBD2 antibodies was not affected by the binding of ER\textsubscript{x}. In addition, serial ChIP assays showed that pS2 promoter region can be simultaneously bound by ER\textsubscript{x} and MBD2 proteins. Furthermore, the synergetic effects of ectopic ER\textsubscript{x} expression and depletion of MBD2 on pS2 transcription also suggest that both proteins do not compete for pS2 promoter occupancy.

Data, obtained from experiments performed on cell lines, indicate that methylation-dependent repression of pS2 expression is mediated by MBD2. Nevertheless, ER\textsubscript{x} binding can counteract the inhibitory effect of DNA methylation without displacing MBD2 proteins from the TATA-box region. Taken together these data might explain pS2 expression in some ER\textsubscript{x}-rich breast cancers despite the methylation of its TATA-box.

Materials and Methods

Cell culture

MDA MB231, MCF7, and HeLa cells were obtained from the American Type Culture Collection (ATCC, Rockville, MD). Cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) containing 1g/l glucose (Eagle, Sigma, L’Isle d’Abeau, France) and supplemented with 10% of heat inactivated fetal bovine serum (Lonza, Vervier, Belgium) and grown at 37\textdegree{}C in a humidified 5% CO\textsubscript{2} atmosphere.

Due to the poor efficiency of repeated siRNA treatments associated with vector transfections in estradiol depleted medium by charcoal extraction, all the experiments were performed in standard DMEM medium supplemented with fetal bovine serum. To investigate the estrogen dependence of pS2 expression, cells were exposed to estradiol or to 5 \mu M 4-hydroxytamoxifen (OHT).

Sodium bisulphite modification

Sodium bisulphite reactions were carried out as previously described [30]. Two regions (nt positions -46 to +167, and nt positions +37 to +314 from the pS2 transcription start site) within the pS2 promoter gene were analyzed. PCR amplifications were accomplished in 100 \mu l using the HotStart Tag DNA polymerase Kit (Qiagen, Courtaboeuf, France) and 0.25 \mu M of the primers (Supplemental Table S1), after 15 min at 95\textdegree{}C for Tag polymerase activation and 35 cycles (30 s denaturation at 94\textdegree{}C, 1 min annealing at 53\textdegree{}C, and 1.5 min extension at 72\textdegree{}C) for the promoter fragment. PCR products were analyzed on a 2% agarose gel containing 1 \mu g/ml ethidium bromide and were quantified by densitometry. Real-time PCR was carried out using LightCycler
\textsuperscript{®} Fast Star DNA Master PLUS SYBR Green I System (Roche Molecular Biochemicals, Maylan, France) and 0.4 \mu M of the primers (Supplemental Table S1). Cycling parameters were 95\textdegree{}C for 10 min followed 45 cycles at 95\textdegree{}C for 10 s, 64\textdegree{}C for 5 s and 72\textdegree{}C for 10s.

ChIP-on-chip

For ChIP-on-Chip analysis, the specific protein-DNA complexes were obtained from independent immunoprecipitations using two different polyclonal anti-MBD2 antibodies (kindly provided by Dr. P. Wade and Dr. E. Ballestar). The ChIP DNAs from the input and bound fractions were labelled and hybridized on microarrays by ProfileXpert service according to Affymetrix\textsuperscript{\textregistered} protocols. Brieﬂy, the ChIP DNA was amplified by random PCR. Enrichment of MBD2-bound sites during the amplification procedure was assayed, by PCR amplification of NBR2, and pS2 promoters, on each ChIP samples before and after amplification. The amplified DNAs were then labelled using the GeneChip\textsuperscript{®} WT Double - Stranded DNA Terminal Labelling Kit and hybridized to the human tiling arrays (Human Promoter 1.0R Arrays), which were then washed and scanned. Raw data from the scans were analyzed using Affymetrix\textsuperscript{®} Tiling Analysis Software (TAS) and the results were viewed in Affymetrix\textsuperscript{®} Integrated Genome Browser (IGB) Software.

Serial ChIP assays (ChIP re-ChIP)

The ERE and the TATA box regions of pS2 are placed about 450 bp apart. In order to immunoprecipitate, from the same DNA fragments, the proteins bound to both regions, nucleoprotein

Chromatin Immunoprecipitation (ChIP) assays

Nucleoprotein complexes were sonicated to reduce the length of DNA fragments to 300–600 bp, and ChIP assays were carried out as described previously [26]. Fifteen \mu l of two different polyclonal anti-MBD2 antibodies (kindly provided by Dr. P. Wade and Dr. E. Ballestar) or 20 \mu l of polyclonal anti-MBD1 (Abcam, Cambridge, UK), anti-MeCP2 (Upstate Biotechnology, Lake Placid, NY), antibodies, or 2.5 \mu g of polyclonal anti-ER\textsubscript{x} antibody (Santa Cruz Biotechnology, Inc, Santa Cruz, CA) or anti-mouse IgG (Dakocytomation, Trappes, France), were used for immunoprecipitation. DNA samples obtained from the input, unbound and bound fractions were quantified by densitometry using the VersaFluor\textsuperscript{TM} fluorometer (Biorad, Ivry, France) and RiboGreen reagent (Molecular Probes, Interchim, Montluçon, France). The amounts of DNA in IgG control fractions were at the limit of the fluorometric detection methods. Thus, PCR quantification of DNA fragments in IgG control fractions was not accurate as in other fractions, since we have used very large parts of the “IgG fractions” when compared with the other fractions.

PCR assays were performed to assess the binding of the proteins to the pS2 5’ flanking sequence. Two regions were analyzed: “pS2 ERE fragment” (from nt positions -461 to -204 from the pS2 transcription start site) and “pS2 promoter fragment” (from nt positions -11 to +292). We amplified, by dose-dependent and quantitative PCR (Q-PCR), equal amounts of total DNA (0.5 ng) from the input, unbound and bound fractions. HotStar Taq polymerase kit (Qiagen) and 0.4 \mu M of the primers (Supplemental Table S1) were used in classical PCR. After 15 min at 95\textdegree{}C for Taq polymerase activation and 37 cycles (30 s denaturation at 94\textdegree{}C, 1 min annealing at 53\textdegree{}C, and 1.5 min extension at 72\textdegree{}C) for the pS2 ERE fragment or 36 cycles (30 s denaturation at 94\textdegree{}C, 1 min annealing at 58\textdegree{}C, and 1.5 min extension at 72\textdegree{}C) for the pS2 promoter fragment, PCR products were analyzed on a 2% agarose gel containing 1 \mu g/ml ethidium bromide and were quantified by densitometry. Real-time PCR was carried out using LightCycler\textsuperscript{®} Fast Star DNA Master PLUS SYBR Green I System (Roche Molecular Biochemicals, Maylan, France) and 0.4 \mu M of the primers (Supplemental Table S1). Cycling parameters were 95\textdegree{}C for 10 min followed 45 cycles at 95\textdegree{}C for 10 s, 64\textdegree{}C for 5 s and 72\textdegree{}C for 10s.
complexes were sheared to reduce the length of DNA fragments to 500–1000 bp. In serial ChIP experiments, following primary immunoprecipitation, the cross-linked complexes were eluted from the immunoprecipitated fraction by incubation with elution buffer (1% SDS, 50 mM NaHCO3) at room temperature for 30 min, and then with 1:10 in ChIP dilution buffer (0.01% SDS; 1.1% Triton X-100, 1.2 mM EDTA, 16.7 mM Tris-HCl, pH 8.0, 167 mM NaCl) followed by reimmunoprecipitation with a second set of antibodies.

Transient MBD2 siRNA knockdown and ERα vector transfections

siRNA duplexes for MBD2 [sense: 5'-GGAGGAAGGUACCG-GAAATT-3'; antisense: 5'-UUUCGGAUCAUCUCCUCCTT-3'] and non-specific siRNA control were obtained from Eurogentec (Eurogentec, Seraing, Belgium). HEG0, an expression vector coding for human ERα [31], and the empty vector pSG5, were provided by Prof. P. Chambon. MBD2 siRNA and ERα vector was transfected with Lipofectamine 2000 [Invitrogen, Carlsbad, CA] according to the manufacturer’s instructions. Briefly, cells were seeded at 2 × 105 cells per well in six-well plates, and grown to 50–60% confluence on the day of transfection. All transfections were done in Opti-MEM medium (Invitrogen) with 625 nM of MBD2 siRNA and 1 μg of ERα expression plasmid. Lipofectamine 2000 complexes were incubated for 4–5 hours. The medium was then removed and replaced with fresh medium. Cells were grown and harvested at various times after the transfection.

Reverse-transcription-PCR analysis

Total RNA was extracted from the cell lines using the RNeasy Mini kit (Qiagen). After extraction, the integrity of total RNA was examined on a 1.2% agarose gel containing 1 μg/ml ethidium bromide and quantified by densitometry using a Fluor’s fluorimeter and Quantity One software (Biorad, Ivy, France) by comparison with serial dilutions of a standard RNA (Roche, Molecular Biochemicals, Maylan, France).

pS2 mRNA was detected by relative RT-PCR using primers described in Supplemental Table S1. Briefly, 0.1 μg of total RNA were amplified simultaneously for β-actin and pS2 using the One Step RT-PCR kit (Qiagen). After 30 min incubation at 50°C, RT was inactivated by heating at 95°C for 15 min, PCR amplification was then performed under the following conditions: 30 cycles, 30 s denaturation at 94°C, 1 min annealing at 55°C and 1.5 min extension at 72°C. PCR products were analyzed on a 2% agarose gel and quantified. The ratio between pS2 and β-actin signals was then determined.

Real-time RT-PCR were also carried out to quantify pS2 mRNA using LightCycler® RNA Master SYBR Green I One-Step RT-PCR mix on a LightCycler® 2.0 system according to the manufacturer’s instructions (Roche). β-actin mRNA was used as a reference. The primers sequences used for reverse-transcription-PCR are available in Supplemental Table S1.

MBD2 mRNA was quantified by competitive quantitative RT-PCR as previously described [32].

Supporting Information

Figure S1 MBD2 siRNA treatments, supplementary data. (A) Expression in MBD2 siRNA transfected cells. Bar chart representing the efficiency of MBD2 siRNA in HeLa, MCF7 and MDA MB231. In mock treated cells, the initial amount of MBD2 molecules / μg of total RNA was: 7.4 × 106 ± 2.8 × 106, in HeLa cells; 5.8 × 106 ± 1.2 × 106, in MDA MB231 cells and 3.2 ± 0.6 × 106, in MCF7 cells. The efficiency of MBD2 siRNA was calculated from the MBD2 mRNA in treated cells compared with mock-treated cells. Each bar represents the mean ± standard deviation of, at least, three independent analyses. (B) MBD1, MBD2 and MeCP2 protein quantifications in HeLa cells expressing transient MBD2 siRNA. HeLa cells were pretreated for 72 h with MBD2 siRNA and again for 24 h. Mock-treated cells were transfected with a non-specific siRNA. Immunoblot analysis of MBD2, MBD1 and MeCP2 proteins in mock-treated and in MBD2 siRNA-treated HeLa cells. MBD2, MBD1 and MeCP2 proteins were probed using rabbit polyclonal antibodies. The same membrane was then stripped and probed using a mouse β-tubulin antibody as a loading control. (C) Bar chart showing the fold change of pS2 expression in MCF7 and MDA MB231 cells depleted in MBD2. pS2 transcripts were quantified by real-time RT-PCR. The fold change was calculated from the amount of pS2 mRNA in treated cells compared with mock-treated cells. Each bar represents the mean ± standard deviation of, at least, three independent analyses.

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Table S1 List of primers.

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Author Contributions

Conceived and designed the experiments: AC EB RD. Performed the experiments: AC. Analyzed the data: AC EB ME. Contributed reagents/materials/analysis tools: AC EB ME. Wrote the paper: AC EB ME RD.

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