CARM1 is a protein arginine methyltransferase (PRMT) that acts as a coactivator in a number of transcriptional programs. CARM1 orchestrates this coactivator activity in part by depositing the H3R17me2a histone mark in the vicinity of gene promoters that it regulates. However, the gross levels of H3R17me2a in CARM1 KO mice did not significantly decrease, indicating that other PRMT(s) may compensate for this loss. We thus performed a screen of type I PRMTs, which revealed that PRMT6 can also deposit the H3R17me2a mark in vitro. CARM1 knockout mice are perinatally lethal and display a reduced fetal size, whereas PRMT6 null mice are viable, which permits the generation of double knockouts. Embryos that are null for both CARM1 and PRMT6 are noticeably smaller than CARM1 null embryos, providing in vivo evidence of redundancy. Mouse embryonic fibroblasts (MEFs) from the double knockout embryos display an absence of the H3R17me2a mark during mitosis and increased signs of DNA damage. Moreover, using the combination of CARM1 and PRMT6 inhibitors suppresses the cell proliferation of WT MEFs, suggesting a synergistic effect between CARM1 and PRMT6 inhibitions. These studies provide direct evidence that PRMT6 also deposits the H3R17me2a mark and acts redundantly with CARM1.

Growing evidence shows that protein arginine methylation plays a significant role in a number of cellular processes, including gene regulation, mRNA splicing, translation, DNA damage response, and signal transduction (1). In mammalian cells, three types of methylarginine are catalyzed by nine protein arginine methyltransferases (PRMTs): Type I methyltransferases generating asymmetric dimethylarginine (ADMA), Type II methyltransferases generating symmetric dimethylarginine (SDMA), and Type III methyltransferases generating monomethylarginine (MMA) (2). CARM1 (also called PRMT4) is recognized as a transcriptional activator and belongs to the Type I methyltransferase class. CARM1 deposits methylation marks on both histone tails (3) and a number of nonhistone proteins (4–6). CARM1 executes its coactivator function by modifying a diverse substrate repertoire that often shares a splicing and/or transcriptional theme (7–10) and includes the H3R17 histone code mark as one of the major regulatory nodes (8, 11–13). CARM1 can methylate histone H3 on both Arg-17 and Arg-26 residues but has a preference for Arg-17 over Arg-26 (14), at least in vitro. It is the only mammalian PRMT known to methylate H3R17 (15); however, interestingly, ablation of CARM1 in mouse embryonic fibroblasts (MEFs) does not result in the total loss of the H3R17me2a mark (9), indicating that another PRMT(s) may methylate this site when CARM1 is absent.

PRMT6 is also a Type I arginine methyltransferase that catalyzes the H3R2me2a mark, which in turn blocks enzymes from depositing the adjacent H3K4me3 and impedes the binding of the effectors of this transcriptional activation mark (16–18). Thus, in this setting, PRMT6 functions as a transcriptional corepressor. Most of the PRMTs methylate glycine- and arginine-rich (GAR) motifs within their substrates, but CARM1 displays unique substrate specificity and cannot methylate GAR motifs (19, 20). Similar to other PRMTs, PRMT6 typically methylates GAR motifs in substrates like HMG1A and fibrillarin (21, 22), but it also methylates non-GAR motifs in the HIV-1 Tat protein and on PRMT6 itself (an automethylation site) (23, 24). PRMT6 methylates multiple sites on histones apart from H3R2, including H2A/H4R3 (17), H2AR29 (25), and H3R42 (26), all of which are non-GAR motifs. Interestingly, both CARM1 and PRMT6 can methylate histone H3 at Arg-42 (26), suggesting that PRMT6 and CARM1 can share substrates in some scenarios. Moreover, an in vitro methylation assay performed on nearly 200 putative CARM1 substrates revealed that PRMT6 could also methylate some of these substrates, but PRMT1 and PRMT5 could not (27). Supporting this possible redundant role for PRMT6 and CARM1 is the evidence that both of these PRMTs function as coactivators in the context of nuclear receptor–mediated transcriptional activation (28).

In this study, we performed in vitro methylation assays using a panel of recombinant type I PRMTs, showing that PRMT6 can efficiently methylate the histone H3R17 site. The CARM1 knockout mice died at birth and showed a reduced fetal size (9). In contrast, the PRMT6 knockout mice were viable (29), allowing us to generate CARM1 knockout embryos on a PRMT6 knockout background. We found that embryos that are null for...
both CARM1 and PRMT6 are smaller than littermates that are only null for CARM1. Importantly, MEFs generated from these double knockout embryos displayed a reduction in the H3R17me2a mark. We also detected elevated levels of DNA damage in these double knockout cells. Using two recently developed small molecular compounds that specifically inhibit CARM1 (30) and PRMT6 (31), we show that there is a synergistic effect between CARM1 and PRMT6 inhibition on cell proliferation. Thus, these data unmask partially redundant functions for CARM1 and PRMT6.

**Results**

**PRMT6 generates the H3R17me2a mark in vitro**

Previously, we showed that global levels of H3R17me2a are not impacted by the loss of CARM1 (9), suggesting that other PRMT(s) may also be able to methylate this site. To test this hypothesis, we purified core histones from the aforementioned CARM1 knockout MEFs and CARM1 enzyme-dead knock-in MEFs (32) (Fig. S1A). Using two different H3R17me2a-specific antibodies, we observed no decrease in H3R17me2a levels in either the CARM1 knockout or the enzyme-dead knock-in MEF lines (Fig. 1A). We confirmed the specificity of these antibodies by using a panel of arginine-methylated peptides (Fig. S1B). The Millipore antibody for H3R17me2a was used in the following studies because there is an additional nonspecific band (Fig. 1A) using the Abcam antibody. To search for the PRMTs that generate the H3R17me2a mark, we performed an *in vitro* methylation assay with a few recombinant type I PRMTs: PRMT1, PRMT3, PRMT6, and CARM1. All four PRMTs can methylate histone H3, to varying degrees, to generate the H3R2me2a mark; however, only CARM1 and PRMT6 can generate the H3R17me2a mark (Fig. 1B). Like CARM1, PRMT6 can generate the H3R26me2a mark, albeit weakly (Fig. 1B). In addition, recombinant PRMT6 can methylate a histone H3 peptide (amino acids 10–27) that harbors the H3R17 motif; however, when the H3R17 site is masked with a dimethyl mark or mutated to alanine, it can no longer be methylated by PRMT6 (Fig. 1C and Fig. S1C). Finally, we performed a time course *in vitro* methylation assay and observed a steady increase in the methylation of both H3R17 and H3R2 sites by PRMT6 (Fig. 1D). These findings suggest that both CARM1 and PRMT6 deposit an ADMA mark on H3R17, at least *in vitro*.

**PRMT6 overexpression results in elevated levels of the H3R17me2a mark in HeLa cells**

To determine whether PRMT6 can methylate the H3R17 site in mammalian cells, PRMT6 and CARM1 were overexpressed in HeLa cells, and the levels of H3R17me2a were tested by Western blotting. Increased levels of H3R2me2a by PRMT6 overexpression were observed as reported previously (18). In addition, the levels of H3R17me2a were also elevated when PRMT6 was ectopically expressed (Fig. 2A). In contrast, the overexpression of CARM1 in HeLa cells did not cause an increase in the H3R17me2a mark, likely because the ectopically expressed CARM1 is predominantly cytoplasmic (Fig. S2A) and does not engage its nuclear substrates. We also note that the GFP–CARM1 protein level is lower than the endogenous CARM1 levels (Fig. 2A), whereas ectopic PRMT6 expression is much higher than its endogenous counterpart. When CARM1 is forced to express in the nucleus (using a GFP expression vector with a nuclear localization signal), a substantial increase in the levels of H3R17me2a was observed as expected (Fig. S2, B and C). Next, we analyzed a HeLa cell line that stably expresses a TAP-tagged form of PRMT6. The protein G domains in the TAP tag allow tight binding by IgG; thus, the ectopic expression of PRMT6 can be visualized by any IgG-type antibody (33). Again, increased H3R17me2a levels were observed in the PRMT6-overexpressing lines (Fig. 2B). We also generated a cell line that stably expresses an ER–PRMT6 fusion protein, which was also linked to a FLAG tag (34). Upon treatment with tamoxifen, the fusion protein was stabilized and translocated into the nucleus. In this experiment, we mixed ER–PRMT6 cells with the same number of WT cells, to have PRMT6-overexpressing cells and control cells adjacent to each other in the same field of view. This mixture culture was treated with tamoxifen and then prepared for immunofluorescence with anti-FLAG and anti-H3R17me2a antibodies (Fig. 2C). FLAG-tagged ER–PRMT6–positive cells showed significantly higher H3R17me2a signals. Even though the H3R17me2a antibody can recognize additional CARM1 substrates, the predominant signal generated by this antibody in ER–PRMT6–positive cells is due to an increase of the H3R17me2a histone mark (Fig. S2D). Together, these findings indicate that PRMT6 can methylate the H3R17 site both *in vitro* and in cells.

**Double knockout mice reveal cross-talk between CARM1 and PRMT6**

Because both CARM1 and PRMT6 have the ability to deposit the H3R17me2a mark, it would be important to gauge the degree of redundancy between these two PRMTs in mice. Previously, we made CARM1 knockout mice and showed that CARM1 knockout embryos are similar in size to their WT littermates at E12.5 but are significantly smaller than their littermates at E18.5 (9). Whereas CARM1 knockout mice died at birth, PRMT6 knockout mice survived to adulthood and did not show any signs of stunted growth (29). Importantly, no PRMT double knockouts have yet been performed. First, we intercrossed PRMT6–/– CARM1+/– double heterozygous mice to generate the single and double knockout embryos in the same litter. At E13.5, both the CARM1 knockout and heterozygous embryos were the same size, and the CARM1/PRMT6 double knockout embryos were distinctly smaller than the CARM1 knockout embryos (Fig. 3A). At E18.5, CARM1 knockout embryos were smaller than CARM1 WT embryos, as we previously reported (9), and the CARM1/PRMT6 double knockout embryos were even smaller (Fig. 3B). Second, we crossed CARM1+/– mice onto a PRMT6 null background, which is fertile, and then intercrossed these mice. Again, CARM1/PRMT6 double knockout embryos were smaller than single knockout embryos at E13.5 (Fig. S3A). Genotyping was performed by PCR (Fig. S3B), and Western blot analysis confirmed the loss of PRMT6 and CARM1 in both embryo extracts and the corresponding cell lines derived from those embryos.
Next, we used two independent sets of primary MEF lines derived from WT, CARM1^{-/-}, PRMT6^{-/-}, and CARM1/PRMT6 double knockout embryos and tested the level of MMA, ADMA, and SDMA, using GAR motif antibodies, as well as the H3R17me2a antibody (which recognizes a cadre of CARM1 substrates) (9) (Fig. 3C). Interestingly, we observed that the depletion of CARM1 and PRMT6 together led to the disappearance of a strong ADMA signal of approximately 42 kDa (Fig. 3C), which could represent an as yet uncharacterized shared substrate for these two PRMTs. In addition, the extracts from CARM1/PRMT6 double knockout embryos display an increase in both MMA and SDMA signals, suggesting that PRMT5 may methylate these substrates when they are no longer blocked with ADMA marks. The phenomenon of substrate scavenging by PRMTs has previously been reported in both PRMT1 knockout cells and CARM1 knockout cells (27, 35).

Figure 1. PRMT6 methylates histone H3 at the Arg-17 site in vitro. A, acid-extracted core histones from CARM1 WT, KO, and enzyme-dead KI MEFs were immunoblotted using the indicated methyl-specific antibodies. Ponceau staining served as an additional loading control. B, in vitro methylation reactions were performed using recombinant histone H3 (2 μg) with the indicated GST-tagged PRMTs (0.5 μg) in the presence of [3H]SAM. Reactions were separated on SDS-PAGE, transferred to PVDF membranes for fluorograph, and immunoblotted using the indicated arginine methyl-specific antibodies to histone H3. C, unmethylated histone H3 peptide and Arg-17-methylated peptides were incubated with GST-PRMT6 and [3H]SAM. Methylation signal by PRMT6 was detected by fluorography. D, in vitro methylation reactions were performed using recombinant histone H3 with the indicated recombinant GST-fused PRMT in the presence of [3H]SAM and incubated at the indicated time. Reactions were separated on an SDS-polyacrylamide gel, transferred to PVDF membranes for fluorograph, and immunoblotted using the indicated methyl-specific antibodies. *, nonspecific band detected by the Abcam H3R17me2a antibody. #, nonspecific band detected by the H3R26me2a antibody.
These data, particularly the reduced size of the CARM1/PRMT6 double knockout embryos, provide genetic evidence for a degree of redundancy between PRMT6 and CARM1. It should be noted that a very small subset of CARM1 substrates are likely methylated by PRMT6, because the H3R17me2a antibody, which cross-reacts with a large number of CARM1 substrates, does not lose any immunoreactivity on whole-cell lysates from PRMT6 knockout lines. We next focused on the histone H3R17me2a mark itself, using acid-purified core histones from these mutant MEF lines.

Loss of CARM1 and PRMT6 leads to reduced H3R17me2a levels in mitosis

Previous studies performed by Sakabe and Hart indicate that the H3R17me2a mark is increased during mitosis (36). To validate these published findings, we performed an expanded cell cycle analysis of the H3R17me2a mark, along with known cell cycle markers like cyclin B1 levels and histone H3S10 phosphorylation states (37). We synchronized HeLa cells with thymidine and nocodazole and tested the levels of H3R17me2a and H4R3me2a at the indicated time points after the cells were released from cell cycle arrest. The intensity of the H3R17me2a mark mirrors the H3Ser10ph signal, clearly supporting the regulation of this mark during the cell cycle (Fig. 4A). The H4R3me2a mark was not altered during this time course (Fig. 4A). Next, we thus focused on studying the impact of CARM1 and PRMT6 loss on the H3R17me2a mark during mitosis, when the level of this mark peaks. WT, CARM1−/−, PRMT6−/−, and CARM1/PRMT6 double knockout MEFs were arrested in mitosis, and the levels of H3R17me2a were tested by Western blotting. During mitosis, H3R17me2a levels were elevated in WT, CARM1 knockout, and PRMT6 knockout MEFs. However, elevated levels of H3R17me2a were not seen in the double knockout MEFs (Fig. 4B). Asynchronous CARM1/PRMT6 double knockout MEFs only showed a slightly lower level of H3R17me2a than other asynchronous MEF cells, suggesting that other methyltransferases may help maintain the basal level of H3R17me2a in the double knockout MEFs or that this antibody weakly recognizes the unmethylated histone. H3Ser10ph is widely known to be a biomarker during mitosis, and elevated levels of it are found in the mitotic cells of all four genotypes tested (Fig. 4B). These findings indicate that both CARM1 and PRMT6 are required for the deposition of the H3R17me2a mark during mitosis.

CARM1 and PRMT6 share some sites of methylation

CARM1 and PRMT6 double knockout MEFs experience increased double-strand DNA damage

CARM1 is reported to methylate p300, which promotes the recruitment of BRCA1 to the promoter of p21 and is required...
for the induction of p21 and GADD45 expression (38). Also, p21 levels are up-regulated in PRMT6-deficient MEFs (29), whereas GADD45a expression is suppressed by PRMT6 in the early development of zebrafish (39). PRMT6 methylates DNA polymerase β to promote efficient base excision repair (40). CARM1 deposits marks that are read by the TDRD3 effector protein, which in turn recruits TOP3B to resolve R-loops and suppress DNA damage (41). These findings implicate both CARM1 and PRMT6 in the DNA damage response. Thus, we were interested to examine the impact of CARM1/PRMT6 loss on the DNA damage response. PRMT6 knockout MEFs showed increased staining for γ-H2AX, as compared with WT or CARM1 knockout MEFs, whereas the double knockout MEFs showed the strongest signal of this biomarker for DNA damage (Fig. 5A). Western blotting analysis further confirmed that there is an increase in γ-H2AX in double knockout MEFs (Fig. 5B). In summary, we showed that there is significantly increased DNA damage response in the CARM1/PRMT6 double knockout MEFs compared with the single knockout lines.

**Synergistic effect between CARM1 and PRMT6 small molecule inhibitions**

Recent advances in small molecular compounds allow us to specifically inhibit the enzyme activities of certain PRMTs (42). TP-064 is a selective and noncompetitive inhibitor to CARM1 (30), and EPZ020411 is a potent and selective inhibitor to PRMT6 (31). By using these compounds, we can avoid the variation between different cell lines. We thus hypothesize that the loss of activity of both CARM1 and PRMT6 will result in reduced cell proliferation, as suggested from the genetic experiments (Fig. 3A and B) and Fig. S3A). To test this hypothesis, we combined increasing concentrations of CARM1 (TP-064) and PRMT6 (EPZ020411) inhibitors to treat WT MEFs, and we tested the relative proliferation of the cells after 6 days (Fig. 6A). As individual agents, 3–10 μM EPZ020411 mildly inhibited the cell growth of WT MEFs, whereas 3–10 μM of TP-064 slowed the proliferation rates. When combined, lower concentrations of TP-064 and EPZ020411 (0.3–3 μM) inhibited the proliferation rates of WT MEF cells (Fig. 6A), suggesting that
there is a synergistic effect between the two compounds. After 4 days of the combined use of a 3 μM concentration of the CARM1 inhibitor and 3 μM PRMT6 inhibitor, a clear inhibition of cell growth was observed with combination treatment over single inhibitor treatment (Fig. S4). Next, we analyzed the combined drug effect values shown in Fig. 6A by using the combination index theorem of Chou and Talalay (43), which is based on the mass-action law principle. A combinatorial index (CI) of <1 indicates synergism, whereas a CI of >1 demonstrates antagonism. Effect-oriented mapping of the combination data demonstrated synergism between the CARM1 inhibitor and the PRMT6 inhibitor (Fig. 6B). These results show that the loss of enzyme activity of both CARM1 and PRMT6 has dramatic effects on cell proliferation rates and cell viability of WT MEFs.

**CARM1 and PRMT6 enzyme activities are required for normal DNA damage response**

Compared with the MEFs deficient in CARM1 or PRMT6 only, MEFs with both CARM1 and PRMT6 knocked out displayed increased γ-H2AX levels (Fig. 5). To make sure this effect on DNA damage response was not due to the idiosyncra-

**Discussion**

In this study, we discovered that PRMT6, like CARM1, has the ability to methylate histone H3 at Arg-17 both in vitro and in vivo (Figs. 1 and 2). To date, PRMT6 has been shown to deposit methylation marks on histone H3R2 (16), H3R17, H3R42 (26), H2A/H4R3 (17), and H2AR29 (25). Among these sites, H3R2me2a and H2AR29me2a are associated with repressed gene expression (16, 17, 25), whereas H3R17me2a, H4R3me2a, and H3R42me2a are generally associated with transcriptional activation (26, 44). Histone H3R2 was thought to be the major histone target site of PRMT6 in cells, and PRMT6 was widely considered a transcriptional repressor (29, 44–46); however, in a few other cases, PRMT6 was reported to act as a transcriptional coactivator (34, 47). Moreover, recent RNA-Seq data from PRMT6 knockout cells showed almost equal amounts of up-regulated and down-regulated genes (48). Interestingly, ~25% of deregulated genes have promoters that are associated with H3R2me2a peaks, and 70% of these H3R2me2a-marked genes are down-regulated in the absence of PRMT6 (48). These data suggest that PRMT6 functions as both a transcriptional repressor and activator, depending on the context. It is unclear how these opposite functions of PRMT6 are regulated, but it could be due to its presence in different transcriptional complexes, which may direct the enzyme activity of this PRMT to different sites on histone tails. Our data revealed that PRMT6 can methylate H3R17 and that it can impinge on the CARM1 pathway.

Most PRMTs, including PRMT6, prefer a GAR motif in their substrates, whereas CARM1 has a preference for a proline residue near the methylated arginine (4, 19). In a few cases, PRMT6 was shown to methylate non-GAR motifs in HIV Tat proteins and in PRMT6 itself (23, 24). Another study showed that both CARM1 and PRMT6 can methylate histone H3 at Arg-42, stimulating the activation of p53-dependent transcription in vitro (26). In this study, we showed that both CARM1 and PRMT6 can methylate H3R2, H3R17, and H3R26 in vitro (Fig. 1, B and D), with varying degrees of efficiency. In cells, we
validated that PRMT6 has the ability to methylate H3R17 (Fig. 2), and we showed the loss of an unknown methylated protein in CARM1 and PRMT6 double knockout MEFs (Fig. 3). Our data further support the hypothesis that CARM1 and PRMT6 share additional common substrates (Fig. 3C), apart from H3R17 and H3R42.

The best example of PRMTs collaborating during certain cellular processes is PRMT1 and PRMT5. Together, they regulate hypoxia and ischemia-induced apoptosis (49), the formation of stress granules (50), cell proliferation (51), and RNA splicing (52). Besides the collaboration between PRMT1 and PRMT5, PRMT1 and CARM1 work together to synergistically regulate p53-dependent and p53-independent genes (53–56). Also, knocking down PRMT1 or CARM1 leads to an increased RNA-binding ability of hnRNPUL1 (57). Even though PRMT6 and PRMT1 in vitro share similar substrates, such as H4R3 (58), Npl3 (21), and TOP3B (59), not much is known about their collaboration in vivo. In the case of CARM1 and PRMT6, they are known to work together to positively regulate hormone-dependent transcription (28, 60). Knockdown of both CARM1 and PRMT6 significantly inhibits the estrogen-stimulated proliferation of breast cancer cells, but it does not inhibit cell proliferation in the absence of estrogen (28). In a recent study, MCF-7 cells were treated with a small molecular inhibitor of both CARM1 and PRMT6, and the estrogen-dependent transcription of GREB1 significantly increased, whereas the loss of CARM1 alone only partially mimicked this effect (61). Finally, in early mouse embryonic development, CARM1 methylation of the H3R26 site seems to be critical for driving cell fate decisions (62, 63), but it remains unclear whether PRMT6, which can weakly methylate the H3R26 site (Fig. 1, B and D), functions redundantly in this setting. In summary, previous reports showed that the combinatorial effect of CARM1 and PRMT6 regulates cell proliferation in a hormone-dependent manner (28). Here, we provide a mechanistic explanation for the reported synergy between CARM1 and PRMT6, which relies on the ability of these two PRMTs to methylate the same motifs on histone tails and perhaps also on nonhistone substrates.

**Materials and methods**

**Antibodies**

Two H3R17me2a antibodies used in this study were from Millipore (#07-214) and Abcam (#ab8284), mostly from Millipore (if not indicated). Other antibodies were anti-H3R2me1 (Abcam, #ab15584), anti-H3R2me2a (Millipore, #07-585), anti-H3R26me2a (Millipore, #07-215), anti-H4R3me2a (Active Motif, #39705), anti-histone H3 (Abcam, #ab18521), anti-FLAG (Sigma, #F1804), anti-β-actin (Sigma, #A1978), anti-CARM1 (Bethyl, #A300-421A), anti-PRMT6 (Bethyl, #A300-929A), anti-H3Ser10ph (Active Motif, #61623), anti-cyclin B1 (BD Biosciences, #554177), anti-H3K9me3 (Active Motif,
CARM1 and PRMT6 share some sites of methylation

Knockout mice and genotyping

CARM1\(^{+/−}\) mice (9) were bred with PRMT6\(^{−/−}\) mice (29) to generate CARM1\(^{+/−}\)-PRMT6\(^{+/−}\) double heterozygous mice. CARM1\(^{+/−}\)-PRMT6\(^{+/−}\) mice were further intercrossed to produce WT, CARM1\(^{−/−}\), PRMT6\(^{−/−}\), and CARM1\(^{−/−}\)-PRMT6\(^{−/−}\) embryos. For timed pregnancies, CARM1\(^{+/−}\)-PRMT6\(^{+/−}\) mice were mated overnight. Females were inspected for vaginal plugs the following morning, and the following noon was taken as day 0.5 of gestation (E0.5). Genomic DNA was isolated from liver biopsies and analyzed by PCR using the following primers: 5′-AGT CCA TGC TGA GCT CGG T-3′ and 5′-TCC ATG CAG CTC ATA TCC A-3′ for PRMT6 WT allele; 5′-AGG GTC ACT GGA AGA-3′ and 5′-ACT CTC AGA ATG TAG-3′ for PRMT6 knockout allele; 5′-CCC ACT TCT GTT ACC TCC TTT G-3′ and 5′-TAA CTA AAA GAA AAT GGA ATG G-3′ for CARM1 genotyping (both WT allele and KO allele). All mouse experiments were reviewed and approved by an institutional animal care and use committee at MD Anderson Cancer Center.

Plasmids and cell culture

GST-tagged PRMTs were expressed from constructs based on the vector pGEX-6P-1 (Amersham Biosciences). Constructs expressing GST-PRMT1, GST-PRMT3, GST-CARM1, and GST-PRMT6 have been described previously (21). NTAP-PRMT6 was generated by cloning PRMT6 cDNA into pCeMM NTAP(GS) vector (EUROSCARF). CARM1 and PRMT6 cDNAs were inserted into pGEP-C1 vector to generate constructs expressing GFP-tagged CARM1 and PRMT6 in mammalian cells. GFP-NLS-CARM1 was generated by inserting mouse CARM1 coding sequence into the BglII site

**Figure 6.** Synergistic effect of CARM1 and PRMT6 inhibitors on the cell proliferation of WT MEFs. A, WT MEFs were treated with a combination of the indicated concentration of CARM1 inhibitor (CARM1i) and PRMT6 inhibitor (PRMT6i) for a total of 6 days. At the end of the culture, cell counting was performed with the CellTiterGlo kit, and the viabilities of different drug treatment groups were normalized by the group with no CARM1 inhibitor or PRMT6 inhibitor (which was set as 100%). B, normalized cell viabilities lower than 100% in A were allowed to be used to calculate the CI values, with CompuSyn software (CompuSyn, Inc., Paramus, NJ, USA). Synergistic interactions are implied by values of <1, whereas values of >1 indicate antagonistic interactions. C and D, WT MEFs were treated with 3 \(\mu M\) CARM1 inhibitor alone, 3 \(\mu M\) PRMT6 inhibitor alone, or the inhibitors combined for 6 days. The treated cells were stained with anti-\(γ\)-H2AX or DAPI to visualize the DNA damage foci in the nucleus (C) or lysed for Western blotting to detect the levels of \(γ\)-H2AX and H3R17me2a (D).
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in pAcGFP1-Nuc (Takara). CARM1 WT, KO, and KI MEFs have been described previously (32). HeLa and HEK293 cell lines were obtained from ATCC. HeLa cells stably expressing NTAP or NTAP-PRMT6 were generated by transfecting NTAP or NTAP-PRMT6 construct into WT HeLa cells, followed by selection of single clones using G418. Stable ER*-FLAG-PRMT6 or NTAP-PRMT6 were generated by transfecting lines were obtained from ATCC. HeLa cells stably expressing arginine-rich peptides were incubated with GST-tagged PRMTs and [3H]SAM at 30°C for 1 h (or the indicated time course). Recombinant histones or peptides were incubated with GST-tagged PRMTs and [3H]SAM at 30°C for 1 h (or the indicated time course). SDS loading buffer was added to the reactions, and the samples were boiled for 5 min. The samples were then applied to electrophoresis on an SDS-polyacrylamide gel and transferred to a PVDF membrane. Then the membrane was dried, and the radioactive signals were recorded with X-ray films.

Histone purifications

Core histones were purified from cell pellets using a Histone Purification Mini kit (Active Motif) according to the manufacturer’s instructions.

In vitro methylation

GST-tagged PRMT1, PRMT3, CARM1, and PRMT6 were purified and used as an enzyme source. Recombinant histones or peptides were incubated with GST-tagged PRMTs and [3H]SAM at 30°C for 1 h (or the indicated time course). SDS loading buffer was added to the reactions, and the samples were boiled for 5 min. The samples were then applied to electrophoresis on an SDS-polyacrylamide gel and transferred to a PVDF membrane. Then the membrane was dried, and the radioactive signals were recorded with X-ray films.

Immunostaining

Cells were seeded on coverslips. At the end of cell culture, the indicated cells were rinsed with PBS and subsequently fixed and permeabilized in buffer containing 2% paraformaldehyde and 0.5% Triton X-100 in PBS at 4°C for 30 min. Then the cells were blocked in 20% fetal bovine serum and incubated with the indicated primary antibody for 1 h at room temperature. Cells were then washed and incubated with Alexa Fluor 647–conjugated secondary antibody (Invitrogen) for 30 min. Finally, they were washed again and stained with DAPI for 5 min at room temperature.

Western blotting

Cells were harvested and lysed, and the cell lysates were applied to Western blot analysis. In brief, cells were harvested and washed three times with cold PBS, and then radioimmune precipitation assay buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1% Nonidet P-40, 0.1% SDS, 1% sodium deoxycholate, 5 mM EDTA, supplemented with proteinase inhibitor mixture) was added to obtain the cell lysates. Cell debris was pelleted and discarded, whereas the supernatant was kept. Protein samples were added with SDS loading buffer and boiled for 5 min, followed by SDS-PAGE. Then the proteins were transferred to a PVDF membrane. The membrane was blocked with 5% fat-free milk for 1 h at room temperature and then incubated with primary antibodies at 4°C overnight. The blot was then washed three times with PBST and incubated with secondary antibodies for 1 h at room temperature. After washing three times with PBST, the membrane was incubated with ECL reagent, and the signals were detected on X-ray film.

Synchronization

HeLa cells were cultured to 40% confluence and treated with 2 mM thymidine for 24 h. Then thymidine was removed by washing with PBS. Fresh medium was then added, and the cells were incubated for 3 h. Afterward, nocodazole was added at a concentration of 100 ng/ml for 12 h to arrest cells at mitosis.

PRMT inhibitor treatment and cell counting

TP-064 (Cayman Chemical, #20256) and EPZ020411 (Cayman Chemical, #19160) were dissolved in DMSO. Mouse embryonic fibroblasts were treated with these drugs for a total of 6 days, with the medium and drugs changed every other day. At the end of the culture, CellTiter-Glo luminescent kit (Promega) was used to measure the cell viability. Briefly, cells were trypsinized and diluted with fresh Dulbecco’s modified Eagle’s medium to the culture volume. Then 100 μl of prewarmed reagent was directly added to 100 μl of the harvested cell culture. The plates were shaken on a horizontal shaker at room temperature for 10 min and applied to a luminescence plate reader. Cell viabilities were normalized to the untreated group.

Data availability

The data sets supporting the conclusions of this article are included within the article and the supporting information.

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Abbreviations—The abbreviations used are: PRMT, protein arginine methyltransferase; CARM, coactivator-associated arginine methyltransferase; MMA, mono methylarginine; SDMA, symmetric dimethylarginine; ADMA, asymmetric dimethylarginine; GAR, glycine- and arginine-rich; γ-H2AX, phosphorylated histone H2AX; DAPI,
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