Upregulation of Protein Phosphatase 2Ac by Hepatitis C Virus Modulates NS3 Helicase Activity through Inhibition of Protein Arginine Methyltransferase 1

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Hepatitis C virus (HCV) is a major cause of chronic liver disease, cirrhosis, and hepatocellular carcinoma worldwide. HCV has a positive-strand RNA genome of about 9.4 kb in size, which serves as a template for replication and for translation of a polyprotein of about 3,000 amino acids. The polyprotein is cleaved co- and posttranslationally by cellular and viral proteases into at least 10 different mature proteins. One of these proteins, nonstructural protein 3 (NS3), has serine protease and NTPase/RNA helicase activity. Arginine 467 in the helicase domain of NS3 (arginine 1493 in the polyprotein) can be methylated by protein arginine methyltransferase 1 (PRMT1). Here we report that the methylation of NS3 inhibits the enzymatic activity of the helicase. Furthermore, we found that PRMT1 activity itself is regulated by protein phosphatase 2A (PP2A). PP2A inhibits PRMT1 enzymatic activity and therefore increases the helicase activity of NS3. This is important, because we found an increased expression of PP2A in cell lines with inducible HCV protein expression, in transgenic mice expressing HCV proteins in hepatocytes, and in liver biopsy samples from patients with chronic hepatitis C. Interestingly, up-regulation of PP2A not only modulates the enzymatic activity of an important viral protein, NS3 helicase, but also interferes with the cellular defense against viruses by inhibiting interferon-induced signaling through signal transducer and activator of transcription 1 (STAT1). We conclude that up-regulation of PP2A might be crucial for the efficient replication of HCV and propose PP2A as a potential target for anti-HCV treatment strategies.

Pathway, i.e., DNA binding. The complex formation between STAT1 and PIAS1 is regulated by an important posttranslational modification of STAT1, arginine methylation (30). Methylation of STAT1 is catalyzed by protein arginine methyltransferase 1 (PRMT1) and protects STAT1 from binding and inactivation by PIAS1 (30).

We have previously reported that HCV inhibits IFN-α-induced signaling at the level of STAT DNA binding (5, 14). Further analysis of the mechanism responsible for HCV interference with STAT signaling led to two a priori independent observations made both with extracts from liver cells of HCV transgenic mice and with liver biopsy samples from patients with CHC: (i) STAT1 was hypomethylated and bound by PIAS1, and (i) protein phosphatase 2Ac (PP2Ac) expression was increased (10). PP2A is a heterotrimeric protein phosphatase consisting of a 36-kilodalton catalytic C subunit (PP2Ac), a 65-kilodalton structural A subunit, and a variable regulatory B subunit. PP2A is expressed in all cell types, is primarily a serine/threonine phosphatase, and is involved in a wide range of cellular processes, including cell cycle regulation, cell morphology, development, signal transduction, translation, apoptosis, and stress response (15, 27). An involvement of PP2A in the regulation of PRMT1 has not been reported before. However, we found that expression of a constitutively active form of PP2Ac in human hepatoma cells (Huh7) resulted in hypomethylation of STAT1 and inhibition of IFN-α-induced signaling (10), demonstrating that PP2Ac is upstream in a signaling pathway.© 2005, American Society for Microbiology. All Rights Reserved.


pathway that regulates STAT1 methylation. We therefore further investigated the role of PP2A in the regulation of STAT1 methylation. In the present paper, we report that PP2Ac interacts directly with PRMT1 and that it inhibits the enzymatic activity of PRMT1.

Interestingly, PRMT1 is also responsible for arginine methylation of NS3, a nonstructural HCV protein with protease and helicase activity (31). The NS3 helicase-NTPase domain consists of the 442 C-terminal amino acids of NS3 and belongs to the DEAD (Asp-Glu-Ala-Asp) box RNA helicase family (19, 21, 24). The domain has probably multiple functions, including RNA-stimulated NTPase activity, RNA binding, and unwinding of RNA regions with extensive secondary structure. Mutational analysis of NS3 helicase revealed that arginines 1490 and 1493 (position in the polyprotein) are essential for enzymatic activity (16). Interestingly, arginine 1493 (arginine 467 of the helicase domain) has been shown to be posttranslationally modified by methylation through PRMT1 (31). However, the functional consequence of this modification is not known. Here we show that arginine methylation inhibits the enzymatic activity of NS3. Furthermore, we show that PRMT1 itself is negatively regulated by PP2A. Therefore, by increasing the expression level of PP2Ac, HCV can indirectly regulate the helicase activity of NS3.

Taken together, our results support an important role of PP2A in the regulation of the viral life cycle of HCV. By inducing PP2Ac overexpression, HCV achieves both an inhibition of IFN-α signaling and an increase in NS3 helicase activity.

MATERIALS AND METHODS

Reagents, antibodies, and plasmids. Human IFN-α was obtained from Hoffmann-LaRoche (Basel, Switzerland). Purified PP2Ac and okadaic acid (OA) were purchased from Upstate (LucernaChem, Luzern, Switzerland) and Sigma (St. Louis, MO). Purified PP2Ac and okadaic acid, D-thiogalactopyranoside, and 3H-AdoMet (specific activity, 15 Ci/mmol) were purchased from Amersham Biosciences (Amersham Pharmacia Biotech Europe GmbH, Uppsala, Sweden). Glutathione S-transferase (GST) columns (MicroSpin GST purification module), IPTG (isopropyl-β-D-thiogalactopyranoside), and 3H-AdoMet (specific activity, 15 Ci/mmol) were obtained from Amersham Biosciences (Amersham Pharmacia Biotech Europe GmbH, Uppsala, Sweden). For protein expression, transfected bacteria cells were grown overnight in LB medium supplemented with ampicillin or kanamycin until they reached an optical density of 0.6 to 0.7. GST-PRMT1 expression was induced with 1 mM IPTG for 3 h at 30°C. The reaction was stopped by heating to 95°C for 5 min. SYBR-PCR was performed based on SYBR green fluorescence (SYBR green PCR master mix, Applied Biosystems, Foster City, CA). To prevent genomic DNA amplification, the primers for RPL19 and PP2Ac were designed across exon-intron junctions. The primers for RPL19 were 5′ GATGCCCCGAAAAACACCTTGG 3′ and 5′ TGGCTGTCACCCCTGCCTT 3′. The primers for PP2Ac were 5′ CCA CACGAACTACACATGG 3′ and 5′ CAGGACATCTGCGCTACAA 3′.

RNA isolation, reverse transcription, and SYBR-PCR. Total RNA was isolated from the cells using a Perfect RNA Eukaryotic Mini kit (Eppendorf, Vaudaux-Eppendorf, Basel, Switzerland) according to the manufacturer’s instructions. RNA was reverse transcribed by Moloney murine leukemia virus reverse transcriptase (Promega, Promega Biosciences Inc., Wallisellen, Switzerland) in the presence of random hexamers (Promega) and deoxynucleoside triphosphate. The reaction mixture was incubated for 5 min at 70°C and then for 1 h at 37°C. The reaction was stopped by heating to 95°C for 5 min. SYBR-PCR was performed using 3H-AdoMet for 2 h at 37°C. The reaction was then stopped by adding 20 μl of sample loading buffer, and the reaction mixture was boiled for 5 min and separated on an 8% SDS-polyacrylamide gel. The upper part of the gel was fixed for 30 min in a solution of isopropanol:water:acetic acid (25:65:10) and then immersed in Amplify (Amersham) with gentle agitation for 30 min. The gel was dried and then exposed to X-ray Hyperfilm (Amersham) for 5 days. The lower part of the gel was cut out and stained with Coomassie blue to check for equal loading.

Immunoprecipitation and immunoblotting. Cell lysates were incubated with anti-PP2Ac (Upstate), anti-PRMT1 (Abcam), anti-BID (BD Biosciences, Basel, Switzerland), anti-NS4B (made by Darius Moradpour), or anti-GST (Amersham) antibodies overnight at 4°C. Protein A-Sepharose (Sigma) was added, and the samples were incubated for 3 h at 4°C on a rotating wheel. After sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis and transfer onto a nitrocellulose membrane (Schleicher & Schuell, Dassel, Germany), proteins were detected with anti-PP2Ac (Upstate) or anti-His antibodies (Santa Cruz). For loading amount control, the nitrocellulose membrane was stained using Bio-Imob FastStain (GenoTech, Cell Concepts GmbH, Umkirch, Germany) according to the manufacturer’s instructions.

GST-PRMT1 binding to PP2Ac. One ml of bacterial cell lysates expressing GST-PRMT1 was loaded onto a GST column (Amersham) and incubated for 10 min at room temperature. After centrifugation at 2,500 rpm for 1 min, 40 μl of total proteins from Huh7 cells was loaded onto the column to a final volume of 120 μl, and then the binding reaction was performed overnight at 4°C on a rotating wheel. The column was then washed four times with cold phosphate-buffered saline. For elution, 120 μl of reduced glutathione was used for 1 h at 4°C on a rotating wheel. To verify the specificity of the binding of PP2Ac to GST-PRMT1, the membrane was stripped and reblotted for BIP.

Methylation assay. An in vitro methylation assay was performed according to a protocol previously described by Mowen et al. in 2001 (30) with some modifications. For Fig. 1C, 50 μg of Huh7 lysate was incubated in the presence of 6 μg of GST-PRMT1 and 1 μM of 16°C-AdoMet in a final reaction volume of 80 μl. After 1 h of reaction followed by heating to 95°C, the reaction was stopped by adding 20 μl of sample loading buffer and boiling for 5 min. The proteins were separated on an 8% SDS-polyacrylamide gel. The lower part of the gel was cut out and stained with Coomassie blue to check for equal loading.

For methyltransferase activity measurements in UHCV-57.3 cells (Fig. 2C), the reaction was performed by using 10 μg of whole-cell lysate in the presence of 3 μl of 16°C-AdoMet for 2 h at 37°C. The reaction was then stopped by adding 20 μl of sample loading buffer, and the reaction volume was boiled for 5 min and separated on an 8% SDS-polyacrylamide gel. The upper part of the gel was fixed for 30 min in a solution of isopropanol:water:acetic acid (25:65:10) and then immersed in Amplify (Amersham) with gentle agitation for 30 min. The gel was dried and then exposed to X-ray Hyperfilm (Amersham) for 5 days. The lower part of the gel was cut out and stained with Coomassie blue to check for equal loading.

For in vitro methylation of purified NS3 helicase (Fig. 3), 30 μg of His-Hel-His (13) was incubated with 2 μg of GST-PRMT in the presence of 6 μl of 16°C-AdoMet for 2 h at 37°C. The reaction was then stopped by adding 20 μl of sample loading buffer, and the reaction volume was boiled for 5 min and separated on an 8% SDS-polyacrylamide gel.

DNA-DNA substrate. To prepare the double-stranded DNA (dsDNA) substrate for the unwinding assay, a short DNA oligonucleotide, 5′-TGG TAC TCC TCA CAC CTG GGC GCC GGC GGT TAA-3′, was 32P radiolabeled using the T4 polynucleotide kinase (Promega). Unincorporated ATP was removed through a Mini Quick Spin column. The labeled oligonucleotide was mixed with an equal amount the unlabeled complementary strand, 5′-GAC TAC GTA CTG TTA ACC GCC GCC CAG GTG TGA GGA GTA CCA GCC CAG ATC TGC-3′. The mixture was heated to 95°C for 3 min and left to cool slowly at room temperature.
DNA helicase assay. Purified methylated (Met-NS3h) or unmethylated (NS3h) His-Hel-His (100 nM) (13) was incubated with the double-stranded DNA substrate (10 nM) in reaction buffer (25 mM MOPS [morpholinepropanesulfonic acid; pH 6.2], 0.1% Tween 20, 3 mM MgCl₂) for 15 min at 23°C. The reactions were started by the addition of 5 mM ATP and then stopped at the indicated time points by the addition of a glycerol loading buffer containing 50 nM of a capture oligonucleotide (5’/H11032-TTA ACC GCC GCC CAG GTG TGA GGA GTA CCA-3’/H11032), 20 mM EDTA, and 0.5% SDS. The samples were analyzed by 8% non-denaturating polyacrylamide gel electrophoresis and quantitated by use of a PhosphorImager (Molecular Dynamics).

Purified methylated NS3h was obtained by incubating 30 μg of His-Hel-His (13) with 1 μg, 2 μg, or 3 μg of purified GST-PRMT in the presence of 9 mM AdoMet for 2 h at 37°C. To prepare the unmethylated NS3h, 30 μg of His-Hel-His was incubated with 2 μg of GST-PRMT1 for 2 h at 37°C (but without AdoMet).

ATPase assay. Reactions were done at 37°C in reaction buffer (25 mM MOPS [pH 6.2], 0.1% Tween 20, 3 mM MgCl₂, 5 mM ATP) with 100 nM of methylated or unmethylated NS3h. ATP quantification was done after 10 min with an ATP determination kit (Biaffin GmbH & Co KG, Kassel, Germany) according to the manufacturer’s instructions.

Treatment of HCV replicon cell lines with the PP2A inhibitor okadaic acid. HuH-7.5 cells harboring a subgenomic HCV replicon (3, 4) (kindly provided by Charles M. Rice, The Rockefeller University, New York, NY) were treated for 18 h with 5 μl/ml dimethyl sulfoxide, 25 nM OA, 2 μM 2’-O-methyladenosine (6) (a specific inhibitor of the HCV RNA-dependent RNA polymerase; kindly provided by Steven S. Carroll, Merck & Co., Inc., West Point, PA), or 1,000-U/ml human IFN-β (Roferon-A; Roche). Preliminary dose titrations indicated that 25 nM OA strongly inhibited PP2A activity with only minimal toxicity after 18 h. Subsequently, total cellular RNA was extracted with Trizol (Invitrogen) following the manufacturer’s instructions. Denaturing agarose gel electrophoresis and Northern blot analyses were performed according to standard protocols. 32P-labeled cDNA fragments representing the entire HCV nonstructural region and a glyceraldehyde-3-phosphate dehydrogenase (GAPDH) probe (12) as an internal control were employed simultaneously for Northern blot hybridization.

PP2Ac siRNA treatment of HuH-7.5 cells harboring a subgenomic HCV replicon. HuH-7.5 cells (10⁶) were transfected using Lipofectamine 2000 from Invitrogen according to the manufacturer’s instructions with 10 μl of a 20-pmol/μl solution of PP2Ac SMARTpool small interfering RNA (siRNA) duplexes from Dharmacon or with a corresponding amount of a nonsilencing siRNA duplex from Qiagen. Four hours after the transfection, cells were recovered for 48 h in Dulbecco’s minimal essential medium supplemented with 10% fetal calf serum. Total RNA was isolated and reverse transcribed. RNA quantification was done at 37°C in reaction buffer (25 mM MOPS [pH 6.2], 0.1% Tween 20, 3 mM MgCl₂, 5 mM ATP) with 100 nM of methylated or unmethylated NS3h. ATP quantification was done after 10 min with an ATP determination kit (Biaffin GmbH & Co KG, Kassel, Germany) according to the manufacturer’s instructions.

Treatment of HCV replicon cell lines with the PP2A inhibitor okadaic acid.

FIG. 1. PP2Ac binds directly to PRMT1 and inhibits its enzymatic activity. (A) Whole-cell lysate from Huh7 cells was incubated overnight at 4°C in a GST column that was left empty (left lane) or in one that was preloaded with GST-PRMT1 (right lane). The unbound proteins were recovered as flowthrough, and the bound proteins were eluted with reduced glutathione. PP2Ac was detected by Western blotting (upper panels). As a control, the membranes were then stripped and reprobed with antibodies against the immunoglobulin binding protein BIP (lower panels). The arrow with the asterisk shows the size of BIP. (B) Purified PP2Ac (0.2 units) or His-Bax (100 ng) was incubated with 10 μg of purified GST-PRMT1. Antibodies to GST were used to immunoprecipitate bound proteins. PP2Ac or His-Bax was detected by Western blotting (with anti-PP2Ac or anti-His, respectively). (C) Whole-cell lysate from Huh7 cells was incubated with 14C-AdoMet alone (lane 1), with 14C-AdoMet plus GST-PRMT1 (lane 2), or with 14C-AdoMet plus GST-PRMT1 plus 0.1 units (lane 3), 0.2 units (lane 4), 0.3 units (lane 5), or 0.4 units (lane 6) of purified PP2Ac. The proteins were separated on an SDS-polyacrylamide gel, and methylation of proteins was detected by autoradiography. To control for equal loading, a 1-cm strip at the bottom of the gel was cut and stained with Coomassie blue (middle panel). The strongest band in the autoradiography (indicated with an asterisk) was quantified for each lane by use of NIH imaging software (lower panel).
done by SYBR-PCR using specific primers for PP2Ac, HCV, and GAPDH. The primers for HCV were 5’ CGCTGCTTCTGCTTTCG 3’ and 5’ CACCCCTGCTATAACC 3’. The primers for GAPDH were 5’ CAAGCTGTGGGCAAGGT 3’ and 5’ GGAAGGCCATGCCAGTGA 3’. The primers for PP2Ac and further details of the method are described above (see “RNA isolation, reverse transcription, and SYBR-PCR.”)

**PP2A phosphatase activity assay.** Assessment of PP2A activity was performed according to the manufacturer’s instructions with whole-cell extracts by use of a

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FIG. 2. Expression of HCV proteins induces PP2Ac. UHCV-57.3 and UGFP-20 cells (controls that express green fluorescent protein) were cultured for 24 h in the presence or absence of tetracycline. (A) PP2Ac mRNA was quantified with real-time RT-PCR. The ratio of the amounts of PP2Ac mRNA in derepressed (minus tet) and repressed cells (plus tet) is shown. Expression of viral proteins increases PP2Ac mRNA levels about twofold. The graph shows the mean (with the standard error of the mean) of two independent experiments. No error bar is shown for UHCV-57.3 cells, because the two values (2.22 and 2.21) were too close. (B) PP2Ac protein (arrow) expression in UHCV-32 cells was examined by Western blotting (left panel). Cells were cultured in presence (left lane) or absence (right lane) of tetracycline. Membranes were probed with a mixture of antibodies to PP2Ac (arrow) and NS4B (asterisk; faint upper band in right lane). The PP2Ac bands were quantified with NIH imaging software (lower panel). As a loading control, the membrane was stained with Blot-FastStain (right panel). (C) UHCV-57.3 cells were cultured in presence (left lane) or absence (right lane) of tetracycline for 24 h. Whole-cell lysates were then incubated for 2 h with radioactively labeled AdoMet. Protein methylation was detected by autoradiography. Equal loading was controlled by cutting the lower part of the gel and staining with Coomassie blue (lower panel). (D) Whole-cell extracts were prepared from two culture plates each of Huh7 cells and of Huh7 cells harboring an HCV replicon (Huh7.5 cells). The samples were analyzed by Western blotting using an antibody against PP2Ac (arrow). Loading was controlled by staining the membrane with Blot-FastStain (lower panel). Huh7.5 cells (lanes 3 and 4) had increased PP2Ac protein expression compared with Huh7 cells (lanes 1 and 2).
AdoMet with 6/H9262 containing 250 mM imidazole (pH 7.2), 1 mM EGTA, 0.1% serine/threonine phosphatase assay system (Promega) with PPTase-2A buffer inhibits the enzymatic activity of PRMT1. Ten/H9262 and 0.5 mg/ml bovine serum albumin.

PP2Ac directly binds and inhibits PRMT1. Purified PP2Ac (Fig. 1C, lanes 3 to 6). We conclude that can be inhibited in a dose-dependent way by the addition of GST-PRMT1 increases the methylation of proteins (Fig. 1C, polyacrylamide gel (Fig. 1C, lane 1). The addition of purified cellular proteins by endogenous PRMT1 can be detected by Western blotting. As shown in Fig. 1A, PP2Ac was specifically bound by PRMT1 (whereas the ubiquitously expressed protein BIP was not). This interaction was direct and did not require additional proteins present in the whole-cell extract, because purified PP2Ac was also bound by GST-PRMT1 (Fig. 1B). As a control, purified His-Bax fusion protein was incubated with GST-PRMT1 but could not be coimmunoprecipitated (Fig. 1B).

Next, we tested if PP2Ac binding to PRMT1 inhibits its enzymatic activity. We have shown previously that expression of a constitutive active form of PP2Ac in Huh7 cells results in a strongly reduced methylation of STAT1 (10). Since STAT1 methylation is catalyzed by PRMT1 (30), we first wanted to test if PP2Ac can bind to PRMT1. To that end, whole-cell extracts from Huh7 cells were loaded onto a GST-PRMT1 column, and the flowthrough as well as the bound proteins were analyzed by Western blotting. As shown in Fig. 1A, PP2Ac was specifically bound by PRMT1 (whereas the ubiquitously expressed protein BIP was not). This interaction was direct and did not require additional proteins present in the whole-cell extract, because purified PP2Ac was also bound by GST-PRMT1 (Fig. 1B). As a control, purified His-Bax fusion protein was incubated with GST-PRMT1 but could not be coimmunoprecipitated (Fig. 1B).

Next, we tested whether GST-PRMT1 can methylate NS3. To this end, we expressed and purified a NS3 helicase protein (NS3h). Incubation of NS3h with purified GST-PRMT1 in the presence of 14C-AdoMet led to methylation of NS3h (Fig. 3, lane 2). This result is in agreement with previous report that NS3 is methylated on arginine 467 of the NS3 helicase domain (corresponding to position 1493 of the polyprotein) (31). Importantly, the addition of purified PP2Ac to the in vitro methylation assays inhibited the arginine methylation of the NS3 helicase domain (Fig. 3, lane 3).

**RESULTS**

**PP2Ac binds directly to PRMT1 and inhibits its enzymatic activity.** We have shown previously that expression of a constitutive active form of PP2Ac in Huh7 cells results in a strongly reduced methylation of STAT1 (10). Since STAT1 methylation is catalyzed by PRMT1 (30), we first wanted to test if PP2Ac can bind to PRMT1. To that end, whole-cell extracts from Huh7 cells were loaded onto a GST-PRMT1 column, and the flowthrough as well as the bound proteins were analyzed by Western blotting. As shown in Fig. 1A, PP2Ac was specifically bound by PRMT1 (whereas the ubiquitously expressed protein BIP was not). This interaction was direct and did not require additional proteins present in the whole-cell extract, because purified PP2Ac was also bound by GST-PRMT1 (Fig. 1B). As a control, purified His-Bax fusion protein was incubated with GST-PRMT1 but could not be coimmunoprecipitated (Fig. 1B).

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**Methylation of NS3 inhibits its helicase activity.** To assess the effect of NS3 methylation on its helicase activity, we used methylated and unmethylated purified NS3 helicase (amino acids 166 to 631 of NS3) in a helicase assay (22). The substrate for the reaction was a radioactively labeled double-stranded DNA. Heating the dsDNA at 95°C for 5 min or unwinding the dsDNA by enzymatic activity of NS3 helicase produced a single-stranded DNA (Fig. 4). Methylated NS3 helicase was obtained by incubating purified NS3 helicase with 1 μg, 2 μg, or 3 μg of purified PRMT1 in the presence of the methyl donor AdoMet (9 mM) for 2 h at 37°C. To exclude an unspecific inhibition of NS3 helicase by PRMT1, the sample with the unmethylated NS3 helicase was also incubated with purified PRMT1 (2 μg) without the methyl donor AdoMet. When tested in the helicase assay, the unmethylated NS3 helicase was more active than the methylated samples (Fig. 4A, lane 2 versus lanes 3 to 5). The difference in the levels of unwinding activity was evident at all time points during a time course experiment (Fig. 4B). We then measured the levels of ATP consumption by the methylated and unmethylated NS3 helicases during a 10-min helicase assay experiment. The more active unmethylated NS3 helicase consumed 65% of the ATP, whereas the less active methylated NS3 helicase samples con-
We conclude that methylation of NS3 by PRMT1 inhibits its helicase activity. Theoretically, this conclusion could be tested more vigorously by doing in vitro helicase assays with a mutated NS3 that cannot be methylated on arginine 467 and therefore should not be inhibited by preincubation with PRMT1. However, the mutation of arginine 467 in the NS3 helicase domain leads to a complete loss of helicase activity (16). Because PP2Ac can inhibit PRMT1, as shown above, the upregulation of PP2Ac by HCV ultimately may result in enhanced helicase activity of NS3.

If PP2A enhances NS3 helicase activity, then inhibition of PP2A should have consequences for HCV replication. We therefore used HuH-7.5 cells harboring a subgenomic HCV replicon (3, 4) and the PP2A inhibitor OA to test this hypothesis. OA was added to the cells at a final concentration of 25 nM. This concentration was used because it significantly inhibited the phosphatase activity of PP2A with only minimal toxicity (data not shown). Treatment of replicon cells with OA indeed inhibited the replication of the HCV replicon (Fig. 5, lane 3). This inhibition was not as strong as that seen after treatment with IFN-α or with the HCV polymerase inhibitor 2′-C-methyladenosine (Fig. 5, lanes 4 and 5) but was still highly significant.

To confirm these results, PP2Ac expression was inhibited in HuH-7.5 cells by use of siRNA (Fig. 6). Again, a reduction of replicon RNA to 70% of the initial value found for HuH-7.5 cells was observed.

DISCUSSION

Given the limited coding capacity of a 9.6-kb RNA genome, HCV has an astonishing ability to evade the powerful immune system of the host and to establish a persistent infection over decades. In its interference with host defense mechanisms, HCV therefore probably targets a limited number of key enzymes of the cells in which it replicates. Here we provide evidence that PP2A is such a key enzyme. We have found that HCV infection or expression of HCV proteins induces the expression of PP2Ac in liver biopsy samples from patients with chronic hepatitis C (10), in liver cells of HCV transgenic mice (10), and in UHCV-57.3 and UHCV-32 cells. PP2A is a widely expressed serine/threonine phosphatase involved in a wide range of cellular processes (15, 27, 34). It can be expected that the induction of such an important phosphatase will have a number of effects on the host cell. Here we have concentrated our efforts on the study of the effect of PP2A on PRMT1, an important arginine methyltransferase. During previous studies, we found that HCV interferes with IFN-α-induced signaling by inhibiting the methylation of STAT1, an important signal transduction pathway. This interference is likely to contribute to the evasion of the immune response by HCV.
transducer of IFN-α. Since STAT1 methylation is catalyzed by PRMT1, we hypothesized that PP2A directly or indirectly inhibits the enzymatic activity of PRMT1. We show here with purified proteins that PP2Ac binds directly to PRMT1 and, furthermore, that PP2Ac binding inhibits the methyltransferase activity of PRMT1 in vitro. Therefore, as a first and important consequence of PP2Ac induction by HCV, STAT1 methylation is reduced. Unmethylated STAT1 is bound by its inhibitor PIAS1, and IFN-α-induced induction of target genes important for cellular defense against HCV is impaired.

Surprisingly, the same mechanism of interference with the host cell is exploited twice by HCV. It has been shown recently that NS3, an HCV protein with protease and helicase activity, can be modified by arginine methylation through the cellular enzyme PRMT1 (31). The functional relevance of this modification was not known, but mutational analysis of arginine residues in the helicase domain provided strong evidence that these arginines are important for the enzymatic activity (16). For example, the mutation of arginine residue 467 of NS3 helicase (arginine 1493 in the polyprotein) to lysine disrupts the enzymatic activity completely (16). Here we show that methylation of NS3 reduces its helicase activity but does not completely abrogate it. Given the direct interaction of PP2Ac with PRMT1, it was not surprising that we did find an inhibition of NS3 methylation by PP2Ac. We conclude that, as a
second important consequence of PP2Ac induction by HCV, NS3 helicase activity is increased.

The characterization of PP2A as a key target of HCV interference with the host cell identifies PP2A as a potential target of treatment strategies against the virus. In a proof-of-concept experiment, we treated replicon cells with OA, a reasonably specific inhibitor of PP2A (27), and found a significant inhibition of replication. The same degree of inhibition of the replicon was found after decreasing the expression of PP2Ac by siRNA. In our model (Fig. 7), inhibition of PP2A by OA results in an enhanced enzymatic activity of PRMT1. NS3 helicase is one of the substrates of PRMT1 and becomes methylated on critical arginine residues. This methylation reduces the helicase activity of NS3, leading to a decrease of viral (or replicated) RNA.

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