# Proteomic Profiling of Lipid Droplet Proteins in Hepatoma Cell Lines Expressing Hepatitis C Virus Core Protein

Shigeko Sato<sup>1</sup>, Masayoshi Fukasawa<sup>1,\*</sup>, Yoshio Yamakawa<sup>1</sup>, Tohru Natsume<sup>2</sup>, Tetsuro Suzuki<sup>3</sup>, Ikuo Shoji<sup>3</sup>, Hideki Aizaki<sup>3</sup>, Tatsuo Miyamura<sup>3</sup> and Masahiro Nishijima<sup>1,†</sup>

<sup>1</sup>Department of Biochemistry and Cell Biology and <sup>3</sup>Department of Virology II, National Institute of Infectious Diseases, Tokyo 162-8640; and <sup>2</sup>National Institute of Advanced Industrial Science and Technology (AIST), Biological Information Research Center, Tokyo 135-0064

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Hepatitis C virus (HCV) core protein has been suggested to play crucial roles in the pathogeneses of liver steatosis and hepatocellular carcinomas due to HCV infection. Intracellular HCV core protein is localized mainly in lipid droplets, in which the core protein should exert its significant biological/pathological functions. In this study, we performed comparative proteomic analysis of lipid droplet proteins in core-expressing and non-expressing hepatoma cell lines. We identified 38 proteins in the lipid droplet fraction of core-expressing (Hep39) cells and 30 proteins in that of non-expressing (Hepswx) cells by 1-D-SDS-PAGE/MALDI-TOF mass spectrometry (MS) or direct nanoflow liquid chromatography-MS/MS. Interestingly, the lipid droplet fraction of Hep39 cells had an apparently lower content of adipose differentiation-related protein and a much higher content of TIP47 than that of Hepswx cells, suggesting the participation of the core protein in lipid droplet biogenesis in HCV-infected cells. Another distinct feature is that proteins involved in RNA metabolism, particularly DEAD box protein 1 and DEAD box protein 3, were detected in the lipid droplet fraction of Hep39 cells. These results suggest that lipid droplets containing HCV core protein may participate in the RNA metabolism of the host and/or HCV, affecting the pathopoiesis and/or virus replication/production in HCV-infected cells.

# Key words: ADRP, DEAD box protein, hepatitis C virus, lipid droplet, TIP47.

Abbreviations: HCV, hepatitis C virus; HCC, hepatocellular carcinoma; MS, mass spectrometry; DNLC, direct nanoflow liquid chromatography; HRP, horseradish peroxidase; ADRP, adipose differentiation-related protein; DDX1, DEAD box protein 1; DDX3, DEAD box protein 3.

Hepatitis C virus (HCV) is a major causative agent of chronic hepatitis (1, 2). Persistent HCV infection, which occurs in more than 70% of infected patients, is strongly associated with the development of liver steatosis, which involves the accumulation of intracellular lipid droplets, cirrhosis, and hepatocellular carcinomas (HCC) (3, 4). Since more than 170 million people in the world are currently infected with HCV (1), and there is no cure that is completely effective, understanding the mechanism by which HCV induces serious liver diseases is one of the most important global public health issues. HCV, a member of the Flaviviridae family, possesses a single-stranded, positive-sense RNA genome of  $\sim 9.6$  kb (5). The HCV genome has a single open reading frame that codes for a large precursor polyprotein of  $\sim$ 3,000 amino acids that is processed into at least 10 individual proteins by host and viral proteases (6).

HCV core protein, the product of the N-terminal portion of the polyprotein, generated upon cleavage at the endoplasmic reticulum by signal peptidase and signal peptide peptidase (7, 8), forms the nucleocapsid of an HCV virion (9). Interestingly, in addition to its function as a structural protein, the core protein exhibits activities leading host cells to lipogenic and malignant transformation in vitro (10–12). Moreover, transgenic mice expressing HCV core protein developed liver steatosis and HCC (13, 14), suggesting an important role of the core protein in these diseases. Many studies have shown that HCV core protein substantially affects various cellular regulatory processes, such as gene transcription (15–17) and signal transduction pathways (12, 18-23), and interacts with a variety of host proteins (12, 18, 19, 22, 24-34), but it is not clear what activities/molecules are practically relevant to the pathogeneses of HCV (core)-derived liver steatosis and HCC. Extensive screenings for genes/proteins exhibiting differences in cellular expression by cDNA microarray (35-40) or proteome analysis (41, 42) have also been tried for HCV-related HCC. Although various genes/ proteins were identified, further studies are required to identify the molecules eventually involved in the pathogenesis of HCV-related HCC.

In host cells, HCV core protein is distributed mainly in lipid droplets and the endoplasmic reticulum (7, 10, 43-46), in which the core protein is predicted to exert its significant biological/pathological functions. In this study, we thus focused on HCV core protein and lipid droplets, and

<sup>\*</sup>To whom correspondence should be addressed. Tel: +81-3-5285-1111, Fax: +81-3-5285-1157, E-mail: fuka@nih.go.jp

<sup>&</sup>lt;sup>†</sup>Present address: Department of Clinical Pharmacy, Faculty of Pharmaceutical Sciences, Doshisha Women's College of Liberal Arts, Kyoto 610-0395.

performed comparative targeted proteomic analysis of the lipid droplet proteins in HCV core–expressing and non-expressing hepatoma cell lines using two strategies: conventional 1-D-SDS-PAGE/MALDI-TOF mass spectrometry (MS) and automated high-throughput direct nanoflow liquid chromatography (DNLC)–MS/MS. We found prominent differences in the protein compositions of lipid droplets between HCV core–expressing and non-expressing hepatoma cell lines.

#### MATERIALS AND METHODS

Cell Lines—The human hepatoma HepG2 cell line constitutively expressing HCV core protein (Hep39) was established as described previously (47). Another HepG2 cell line transfected with expression vector pcEF321swxneo without the HCV core protein insert (Hepswx) was used as a mock control (47). Both cell lines were plated on collagen-coated dishes (Asahi Techno Glass, Tokyo, Japan) and maintained in the normal culture medium [DMEM supplemented with 10% fetal bovine serum, 100 units/ml Penicillin G, 100 µg/ml streptomycin sulfate, and 1 mg/ml G418 (Sigma, St. Louis, MO, USA)] under a 5% CO<sub>2</sub> atmosphere at 37°C.

Lipid Droplet Preparation-Hepswx and Hep39 cells were seeded at  $4 \times 10^6$  cells/dish (150 mm, inner diameter) in 25 ml of normal culture medium and cultured for one day. For efficient formation of lipid droplets by cells, cholesterol (final 20 µg/ml) and oleic acid (final 400 µM)/fatty acid-free BSA (final 60 µM) complex, prepared as stock solutions of 5 mg/ml cholesterol in ethanol and 10 mM oleic acid/1.5 mM BSA in PBS, respectively, were added to the medium. Each cell line was further incubated for 48-72 h at 37°C. For proteomic analysis of lipid droplet proteins, confluent monolayers of Hepswx and Hep39 cells in fifteen cell culture dishes (150 mm, inner diameter) were harvested by scraping and pelleted by centrifugation  $(200 \times g \text{ for 5 min at } 4^{\circ}\text{C})$ . After being washed with PBS three times, each cell pellet was resuspended in 10 mM Tris-HCl buffer, pH 7.5, containing 0.25 M sucrose and Complete<sup>™</sup>, EDTA-free (Roche, Mannheim, Germany) to achieve a final volume equal to five times the volume of the cell pellet (*i.e.* a 20% cell suspension). The cell suspension was homogenized with a ball-bearing homogenizer (48), and then centrifuged at  $800 \times g$  for 5 min at 4°C. One milliliter of each post-nuclear supernatant fraction was layered under 2 ml of 10 mM Tris-HCl buffer, pH 7.5, containing 0.15 M NaCl (TN-buffer). After centrifugation at  $100,000 \times g$  for 60 min at 4°C, the lipid droplet fraction, *i.e.* the distinct white band on the top of the preparation, was collected with a pipetman. The floating lipid droplet fraction was diluted with 3.5 ml of TN-buffer and then re-purified by centrifugation  $(100,000 \times g \text{ for } 30 \text{ min at})$ 4°C). This washing step was repeated three times. Lipid droplets in the floating fractions in both cells were enriched up to more than 500-fold compared with those in the total cell lysates as estimated by their protein contents. The amounts of lipid droplets isolated from Hepswx and Hep39 cells were nearly the same. The purified lipid droplet fractions ( $\sim 0.1$  mg of protein per ml) were stored at -80°C until use. The purity of the lipid droplet fractions was verified by microscopic and immunoblot (Fig. 1) analyses. Adipose differentiation-related protein (ADRP), a



Fig. 1. Immunoblot analysis of lipid droplet fractions in Hepswx and Hep39 cells using antibodies against various organelle markers. Total cell lysates and lipid droplet fractions (1.5  $\mu$ g of protein per lane in the case of anti-ADRP; 5  $\mu$ g of protein per lane in others) from Hepswx and Hep39 cells were analyzed by immunoblotting with the indicated antibodies.

known lipid droplet protein, was significantly enriched in the lipid droplet fractions of Hepswx and Hep39 cells (Fig. 1). Other organelle marker proteins, such as nucleoporin p62 for the nucleus, prohibitin for the mitochondria, fatty acid synthase for the cytoplasm, GM130 for the Golgi apparatus, EEA1 for the early endosome, or annexin II for the plasma membrane, were not detected in the lipid droplet fractions of either cells (Fig. 1). Small amounts of calnexin, a marker of the endoplasmic reticulum, which is a major organelle, were detected in the lipid droplet fractions of both cells to a similar extent. Although we did not detect calnexin in the lipid droplet fractions by proteomic analysis (see Tables 1 and 2), the lipid droplet fractions of both cells could be contaminated with a small amount of endoplasmic reticulum.

1-D-SDS-PAGE/MALDI-TOF MS Analysis—The lipid droplet fraction (30  $\mu$ g protein) of each cell line was fractionated in a 10% SDS-polyacrylamide gel, and the gel was stained with Coomassie Brilliant Blue. The protein bands were excised from the gel and subjected to in-gel trypsin digestion. The tryptic peptide mixtures were analyzed by MALDI-TOF MS as described previously (49). Prior to MALDI-TOF MS analysis, the peptide mixtures were desalted using C18 Zip Tips (Millipore, Billerica, MA, USA) according to the manufacturer's instructions. The peptide data were collected in the reflection mode and with positive polarity, using a saturated solution of

Table 1.	Lipid di	roplet pro	oteins identi	fied in Hepsv	vx and Hep39	cells by	means of 1-	D-SDS-PA	GE/MALDI	-TOF MS
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Dectain	$\mathbf{M}_{\mathbf{r}}$	A	SDS band No. <sup>a</sup>	
Protein	Molecular mass (RDa) (calc.)	Accession No.	Hepswx	Hep39
PAT family proteins				
Adipose differentiation-related protein (ADRP)	48.1	34577059	5	21
Cargo selection protein/TIP47	47.0	20127486		22
Lipid metabolism				
Acyl CoA synthetase long chain family member 3	80.4	42794752	4	18
Cytochrome $b_5$ reductase	34.3	4503327	9	26
Lanosterol synthase	83.3	4808278	4	18
NAD(P)-dependent steroid dehydrogenase-like; H105e3	41.9	8393516	8	25
Retinal short-chain dehydrogenase/reductase retSDR2	33.0	7705905	10	27
Cytosolic phospholipase A <sub>2</sub>	85.2	1352707		14
Rab GTPases				
Rab1A	22.7	4758988	13	30
Rab1B	22.2	23396834	13	30
Rab5C	23.5	38258923	11	28
Rab7	23.5	34147513	12	29
RNA metabolism/binding				
DEAD box protein 1 (DDX1)	82.9	6919862		16
DEAD box protein 3 (DDX3)	73.2	3023628		18
HC56/gemin 4	118.8	10945430		15
Other/unknown proteins				
BiP protein	70.9	14916999	3	17
CGI-49 protein	46.9	7705767	6	23
Heat shock protein gp96 precursor	90.2	15010550	2	14
Ancient ubiquitous protein 1	41.4	31712024	7	
Major vault protein	99.3	15990478	1	
Apoptosis-inducing factor–homologous	40.5	13543964		24
mitochondrion-associated inducer of death				
KIAA0887 protein	52.4	4240263		21
Protein disulfide-isomerase [EC 5.3.4.1] ER60 precursor	56.7	1085373		20
Transport-secretion protein 2.1	57.7	9663151		19
HCV core protein	20.6	974345		31

<sup>a</sup>Band numbers correspond to those in Fig. 2.

 $\alpha$ -cyano-4-hydroxycinnamic acid (Sigma) in 50% acetonitrile and 0.1% trifluoroacetic acid as the matrix. Spectra were obtained using a Voyager DE-STR MALDI-TOF mass spectrometer (PE Biosystems, Foster City, CA, USA). Internal calibration was performed with adrenocorticotropic hormone, fragment 18–39 (Sigma), and bradykinin fragment (Sigma). The data base–fitting program MS-Fit available at the WWW site of the University of California, San Francisco (prospector.ucsf.edu/ucsfhtml3.4/msfit.htm) was used to interpret the MS spectra of protein digests (50).

DNLC-MS/MS Analysis—The lipid droplet fraction (10 µg protein) of each cell line was first delipidated by chloroform—methanol extraction as originally described (51). Two volumes of chloroform and 1 volume of methanol were mixed with 0.8 volume of the lipid droplet fraction. Then, 1 volume of chloroform and 1 volume of water were added to the mixture, and the mixture was vortexed for 30 s, and centrifuged at 10,000 × g for 5 min at room temperature. The resulting organic (lower) phase was removed. The aqueous (upper) phase and interface, containing all the lipid droplet proteins, was lyophilized. The delipidated lipid droplet proteins were digested with endoproteinase Lys-C, and the resulting peptides were analyzed by DNLC-MS/MS as described (52, 53).

Cell Fractionation-All manipulations were performed at 4°C or on ice. After being washed with PBS, confluent monolayers of Hepswx and Hep39 cells were harvested by scraping and pelleted by centrifugation  $(200 \times g, 5 \text{ min})$ . The precipitated cells were homogenized with a ballbearing homogenizer in 10 mM Tris-HCl buffer, pH 7.5, containing 0.25 M sucrose, and Complete<sup>™</sup>, EDTA-free. After centrifugation of the lysate at  $800 \times g$  for 5 min, the cytosolic fraction  $(100,000 \times g \text{ supernatant})$  and membrane fraction  $(100,000 \times g \text{ precipitate})$  were separated from the post-nuclear supernatant fraction by centrifugation at  $100,000 \times g$  for 60 min. The membrane fraction was resuspended in TN-buffer and then re-purified twice by centrifugation. The protein concentrations of these preparations were determined with BCA protein assay reagents (Pierce Biotechnology, Rockford, IL, USA) using BSA as a standard.

Immunoblot Analysis—Equivalent amounts of proteins from Hepswx and Hep39 cells were separated in a 10 or 12.5% SDS-polyacrylamide gel and then electrophoretically transferred to a polyvinylidene difluoride membrane. The membranes were blocked overnight at 4°C or 30 min at room temperature in TBS containing 0.1% Tween 20 and 5% skim milk. The blots were probed with a mouse

Table 2. Lipid drop	olet proteins identified in He	pswx and Hep39 cells b	v means of DNLC-MS/MS.
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Ductoin	Accession	Molecular mass	Matched peptide sequence				
Protein	No.	(kDa) (calc.)		Hepswx <sup>a</sup>	Hep39 <sup>a</sup>		
PAT family proteins Adipose differentiation-related protein (ADRP)	34577059	48.1	+	TITSVAMTSALPIIQK DAVTTTVTGAK EVSDSLLTSSK	+ TITSVAMTSALPIIQK DAVTTTVTGAK EVSDSLLTSSK		
Cargo selection protein / TIP47	20127486	47.0	+	VSGAQEMVSSAK	+ VSGAQEMVSSAK		
Lipid metabolism							
Acyl-CoA synthetase long-chain family member 3	42794752	80.4	+	VLSEAAISASLEK	+ ELTELARK		
Cytochrome $b_5$ reductase	4503327	34.2	+	DILLRPELEELRNK	+ SNPIIRTVK		
Gastric-associated differentially- expressed protein YA61P	6970062	14.9	+	AIGLVVPSLTGK	+ AIGLVVPSLTGK		
Retinal short-chain dehydrogenase/ reductase retSDR2	7705905	33.0	+	HGLEETAAK	+ FDAVIGYK		
Sterol carrier protein 2–related form, 58.85K	86717	58.8	+	LQNLQLQPGNAK	+ LQNLQLQPGNAK		
Acyl-CoA synthetase long-chain family member 4	4758332	74.4	+	SDQSYVISFVVPNQK			
Fatty acid binding protein 5	4557581	15.2	+	ELGVGIALRK			
Hydroxysteroid (17-beta) dehydrogenase 4	4504505	79.7			+ NHPMTPEAVK		
Rab GTPases							
Rab1A	4758988	22.6	+p	QWLQEIDRYASENVNK RMGPGATAGGAEK	+ RMGPGATAGGAEK		
Rab1B	23396834	22.1	$+^{\mathrm{b}}$	QWLQEIDRYASENVNK	+ RMGPGAASGGERPNLK		
Rab7	34147513	23.5	+	NNIPYFETSAK	+ ATIGADFLTK		
Rab18	20809384	22.9	+	HSMLFIEASAK	+ ILIIGESGVGK		
Rab10	12654157	22.5	+	LLLIGDSGVGK			
Rab11	4758986	24.5	+	VVLIGDSGVGK			
Rab8	539607	23.6			+ IRTIELDGK		
RNA metabolism/binding							
DEAD box protein 1 (DDX1)	6919862	82.4			+ FGFGFGGTGK		
DEAD box protein 3 (DDX3)	3023628	73.2			+ GVRHTMMFSATFPK		
IGF-II mRNA-binding protein 3	30795212	63.7			+ EGATIRNITK		
Ribosomal protein L29	14286258	17.8			+ AQAAAPASVPAQAPK		
Other/unknown proteins							
Apoptosis-inducing factor homologous mitochondrion-associated inducer of death	13543964	40.5	+	EVTLIHSQVALADK	+ EVTLIHSQVALADK		
BiP protein	14916999	72.3	+	SQIFSTASDNQPTVTIK	+ VYEGERPLTK		
Hypothetical protein DKFZp586A0522.1	7512845	28.2	+	LQHIQAPLSWELVRPH- IYGYAVK	+ RELFSNLQEFAGPSGK		
Prolyl 4-hydroxylase, beta subunit	20070125	57.1	+	VHSFPTLK	+ AEGSEIRLAK		
Ancient ubiquitous protein 1	31712024	41.4	+	GTQSLPTASASK			
Heat shock protein gp96 precursor	15010550	90.2	+	FAFQAEVNRMMK			
Hypothetical protein FLJ21820	11345458	37.3	+	DIYGLNGQIEHK			
Molecule possessing ankyrin repeats induced by lipopolysaccharide	38173790	78.1	+	CLIQMGAAVEAK			
Ubiquitin-conjugating enzyme E2G 2, isoform 1	15079469	18.6	+	RLMAEYK			
CGI-49 protein	7705767	46.9			+ AGGVFTPGAAFSK		
DILV594	37182139	31.4			+ RELFSQIK		
Hypothetical protein DKFZp564F0522.1—human (fragment)	7512734	33.1			+ ILRTSSGSIREK		
Hypothetical protein HSPC117	7657015	55.2			+ EQLAQAMFDHIPVGVGSK		
Tumor protein D52–like 2 isoform e	40805860	22.2			+ TQETLSQAGQK		
Vesicle amine transport protein 1	15679945	41.9			+ VVTYGMANLLTGPK		

<sup>a</sup>+, detected. <sup>b</sup>This peptide sequence is present in both Rab1A and Rab1B.

monoclonal anti-ADRP antibody (PROGEN Biotechnik GmbH, Heidelberg, Germany) (1:25), a guinea pig polyclonal anti-TIP47 antibody (PROGEN Biotechnik GmbH) (1:250), a mouse monoclonal anti-HCV core protein antibody (Anogen, Ontario, Canada) (1: 1,000), a mouse monoclonal anti-DDX1 antibody (Pharmingen, San Diego, CA, USA) (1:500), or a rabbit polyclonal anti-DDX3 antibody (antibody custom-made by Invitrogen, CA, USA) (1:500) for 90 min at room temperature. The blots were then incubated with horseradish peroxidase (HRP)-conjugated goat anti-rabbit IgG (BIO-RAD), HRP-conjugated goat anti-mouse IgG (BIO-RAD), or HRP-conjugated goat antiguinea pig IgG (ICN Pharmaceuticals, Aurora, OH, USA) at 1:2,000 dilution for 60 min. Detection of immunoreactive proteins was performed with an ECL system (Amersham Biosciences Corp., Piscataway, NJ, USA).

#### RESULTS

Proteomic Analysis of Lipid Droplets by 1-D-SDS-PAGE/MALDI-TOF MS—Lipid droplet proteins from control (HCV core non-expressing) Hepswx cells and HCV core–expressing Hep39 cells were separated by 10% SDS-PAGE, and the protein bands were visualized by Coomassie Brilliant Blue staining (Fig. 2). In each cell



Fig. 2. Signature SDS-PAGE patterns of the lipid droplet fractions of Hepswx and Hep39 cells. Proteins in the purified lipid droplet fractions (30  $\mu$ g of protein per lane) of Hepswx cells and Hep39 cells were separated in a 10% SDS-polyacrylamide gel, and visualized by Coomassie Brilliant Blue staining. The 31 numbered bands were excised from the gel, subjected to in-gel trypsin digestion, and processed for MALDI-TOF-MS. Molecular weights (MW) are given to the left of the gel.

line  $\sim$ 30 bands were seen. The visible bands (areas) were excised from the gels, trypsinized, and analyzed by MALDI-TOF MS. Among the 31 bands, we identified 25 proteins: 15 proteins in Hepswx cells and 23 proteins in Hep39 cells (Fig. 2 and Table 1). Thirteen of the 25 proteins were detected in both types of cell. The lipid droplet proteins found in both Hepswx and Hep39 cells could be categorized into four groups: (1) PAT family proteins, *i.e.* ADRP and TIP47; (2) multiple molecules involved in lipid metabolism; (3) several Rab GTPases; and (4) other/unknown proteins (Table 1). In addition, Hep39 cells contained another group of proteins involved in RNA metabolism/binding (Table 1).

Proteomic Analysis of Lipid Droplets by DNLC-MS/ MS—Some protein bands in Fig. 2 could not be identified, probably due to the restricted separation capacity of 1-D-SDS-PAGE (*i.e.* multiple proteins migrating to the same area). We had, however, difficulty in applying 2-DE to the separation of lipid droplet proteins because of their hydrophobic characteristics. We then tried a new LC-based MS strategy. Lipid droplet fractions from Hepswx and Hep39 cells were delipidated and then digested with Lys-C. The resulting peptide mixtures were directly analyzed using a DNLC-MS/MS system (52). We identified 36 lipid droplet proteins: 24 proteins in Hepswx cells and 27 proteins in Hep39 cells (Table 2). Twenty-three lipid droplet proteins were newly identified with this system. Fifteen proteins detected in both cell lines were classified into four categories (Table 2) as in the case of 1-D-SDS-PAGE/ MALDI-TOF MS analysis. A group of proteins involved in RNA metabolism/binding was also found only in Hep39 cells (Table 2).

Proteins Exhibiting Differences in Their Association with Lipid Droplets Due to HCV Core Protein Expression—SDS-PAGE patterns of lipid droplet proteins were similar but revealed several distinct differences in protein composition between Hepswx and Hep39 cells (Fig. 2). The most remarkable differences were seen in the bands corresponding to PAT family proteins. The amount of ADRP, a major PAT family protein in lipid droplets in the liver (54, 55), and likely to be the most abundant lipid droplet protein in Hepswx cells (Fig. 2, band 5), seemed to be less in HCV core-expressing Hep39 cells (Fig. 2, band 21). On the other hand, TIP47, which is also known to be a PAT family protein in lipid droplets (56, 57), was detected as a major protein only in Hep39 cells (Fig. 2, band 22, and Table 1). To confirm these findings, the contents of ADRP and TIP47 in the lipid droplet fractions of Hepswx and Hep39 cells were examined by immunoblot analysis with specific antibodies. The lipid droplet fraction of HCV core-expressing Hep39 cells showed an apparently lower content of ADRP and a much higher content of TIP47 than the levels in Hepswx cells (Fig. 3).

Next we examined the cellular distributions of ADRP and TIP47 in Hepswx and Hep39 cells by cell fractionation. ADRP was highly concentrated in the lipid droplet fractions of both cells, even though the content in the lipid droplets was much lower in Hep39 cells than in Hepswx cells (Fig. 4). ADRP was not detected in post-nuclear supernatant fractions or in either the cytosol or membrane fractions, probably because of low expression levels in these cells or low affinity of the anti-ADRP antibody we used



Fig. 3. The lipid droplet fraction of Hep39 cells contains less ADRP, but more TIP47, than Hepswx cells. Lipid droplet fractions (1.5  $\mu$ g of protein per lane) from Hepswx and Hep39 cells were analyzed by immunoblotting with the indicated antibodies.



Fig. 4. Subcellular localization of ADRP and TIP47 in Hepswx and Hep39 cells. Hepswx and Hep39 cells were fractionated into post-nuclear supernatant (lane L),  $100,000 \times g$  precipitate (lane P),  $100,000 \times g$  supernatant (lane S), and lipid droplet (lane D) fractions as described in "MATERIALS AND METHODS." Ten micrograms of protein was processed for gel electrophoresis, and then analyzed by immuonoblotting with anti-ADRP and anti-TIP47 antibodies.

(Fig. 4). The mRNA expression level of ADRP in Hep39 cells was less than half that in Hepswx cells (data not shown), consistent with the immunoblot data shown in Fig. 4. These results suggest that the lower ADRP content in the lipid droplet fraction of Hep39 cells is due to a low expression level of ADRP. In contrast, Hep39 cells had much more TIP47 in the lipid droplet fraction (Figs. 3 and 4, lanes D), but the cellular TIP47 content of Hep39 cells was not more than that in Hepswx cells (Fig. 4, lanes L). Besides the lipid droplet fraction, the cytosolic fraction of Hepswx cells was found to contain TIP47 at a substantial level, while the cytosolic fraction of Hep39 cells did not (Fig. 4, lanes S). These results indicate that the intracelular distribution of TIP47 shifts drastically from the cytosol to lipid droplets in HCV core–expressing Hep39 cells.

Another obvious difference between Hepswx and Hep39 cells in Fig. 2 is the presence of a specific  $\sim$ 85 kDa band (Fig. 2, band 16) in Hep39 cells, which was identified as DEAD box protein 1 (DDX1), a DEAD box protein family member, by 1-D-SDS-PAGE/MALDI-TOF MS analysis (Table 1). DNLC-MS/MS analysis also supported the existence of DDX1 in the lipid droplet fraction of Hep39 cells (Table 2). In addition, DEAD box protein 3 (DDX3), another DEAD box protein family member, was also detected in the lipid droplet fraction of Hep39 cells by means of the two different strategies used for proteomic analysis (Tables 1 and 2), suggesting that DDX3 is a major lipid droplet protein in Hep39 cells. To verify the association of DDX 1 and DDX3 with lipid droplets in Hep39 cells, immunoblot analysis was carried out. Figure 5 shows that DDX 1 and DDX 3 exist in the lipid droplet fraction of HCV core-expressing Hep39 cells, but not Hepswx cells. These results imply the



Fig. 5. Hep39 cells, but not Hepswx cells, have DDX1 and DDX3 in the lipid droplet fraction. Lipid droplet fractions (0.5  $\mu$ g of protein per lane) in Hepswx and Hep39 cells were analyzed by immunoblotting with anti-DDX1 and anti-DDX3 antibodies.

special pathological functions of DDX1 and DDX3 in lipid droplets in HCV core–expressing cells.

### DISCUSSION

To analyze lipid droplet proteins, we performed proteomic analysis by means of 1-D-SDS-PAGE/MALDI-TOF MS and automated DNLC-MS/MS, and identified 25 and 36 proteins, respectively (Tables 1 and 2). Many more lipid droplet proteins were identified by DNLC-MS/MS, and 22 major proteins separated by 1-D-SDS-PAGE (Fig. 2, bands, 2, 3, 4, 5, 7, 9, 10, 12, 13, 16, 17, 18, 21, 22, 24, 26, 27, 29, and 30) and detected on MALDI-TOF MS analysis were also detected on DNLC-MS/MS analysis. These results indicate that DNLC-MS/MS is a very sensitive and reliable system as well as a high-throughput method. Particularly, DNLC-MS/MS would be a powerful system for exhaustive proteomic analysis of protein mixtures/ complexes (up to  $\sim$ 100 proteins) such as lipid droplets.

In our targeted proteomic study, we identified a total of 48 lipid droplet proteins: 30 proteins in control Hepswx cells, 38 proteins in HCV core-expressing Hep39 cells, and 20 proteins in both cell lines. The resident lipid droplet proteins were classified into four groups (Tables 1 and 2), consistent with the recently reported data obtained on proteomic analysis of lipid droplet proteins in other cell lines (58-60). In addition, multiple proteins, such as the sterol carrier protein 2-related form, fatty acid binding protein 5, and apoptosis-inducing factor homologous mitochondrion-associated inducer of death, were newly identified as lipid droplet proteins in this study. These accumulated data obtained on proteomic analysis will be useful for understanding the biogenesis and functions of lipid droplets about which little is yet known.

A prominent effect of the expression of HCV core protein on the composition of lipid droplet proteins was observed among the PAT family proteins, i.e. ADRP and TIP47. HCV core-expressing Hep39 cells contained much less ADRP in the lipid droplet fraction (Fig. 3), probably because of the lower cellular expression level and the lack of induction of expression upon lipid loading (data not shown). In contrast, a substantial amount of TIP47 was associated with the lipid droplet fraction of Hep39 cells (Fig. 3). Perilipin, a structural protein of lipid droplets in adipocytes, ADRP, and TIP47, termed PAT family proteins (61), share extensive amino acid sequence similarity (61–63), suggesting a common biological function in lipid droplet formation. For example, the transition in surface protein composition of lipid droplets from ADRP to perilipin occurs during adipocyte differentiation (64). Thus, TIP47 might replace ADRP

on the lipid droplets in Hep39 cells. Cellular TIP47 was not up-regulated in Hep39 cells, resulting in a reduction of TIP47 in the cytosolic fraction (Fig. 4). Since TIP47, originally identified as having the ability to interact with the mannose 6-phosphate/IGF-II receptor (63), appears to be essential for the endocytic recycling system (65-67), the altered distribution of cellular TIP47 in Hep39 cells could affect intracellular membrane trafficking pathways. Consistent with this assumption, our preliminary results showed that the rate of protein secretion from cells was apparently slower for Hep39 cells than Hepswx cells (unpublished data). Patients chronically infected with HCV (68) and HCV core-transgenic mice (69) exhibit decreased levels of plasma very low density lipoproteins secreted from the liver, also suggesting interference with intracellular membrane trafficking (secretion pathways) by HCV core proteins. We currently speculate that the reduction in cellular ADRP expression mediated by HCV core protein causes the accumulation of TIP47 in lipid droplets as a substitute, and that the resulting depletion of TIP47 in the cytosol could cause the impairment of intracellular membrane trafficking, followed by the cellular accumulation of membrane lipids and consequent lipid droplet formation. Although further studies remain to be done to confirm these possibilities, we suggest that HCV core protein influences not only the biogenesis of lipid droplets but also intracellular membrane trafficking.

Another interesting finding in this study is that Hep39 cells, unlike Hepswx cells, contain DEAD box proteins, DDX1 and DDX3, as major lipid droplet proteins (Figs. 2 and 5). On the basis of the results of studies involving yeast two-hybrid assays, DDX3 has been shown to be able to interact with HCV core protein, and studies involving immunofluorescent microscopy have revealed that DDX3 is distributed in cytosolic spots such as lipid droplets (27, 70, 71). These results, together with our present findings, suggest that DDX3 is associated with lipid droplets via HCV core proteins located on lipid droplets. In addition to DDX1 and DDX3, which possess ATPase/RNA helicase activities (27, 72, 73), several other proteins involved in RNA metabolism/binding, including HC56/gemin 4 and IGF-II mRNA-binding protein 3, were also detected in the lipid droplet fraction of HCV core-expressing Hep39 cells (Tables 1 and 2). Recently Dvorak et al. reported that RNA itself can be associated with lipid droplets in human mast cells (74). Taken together, these data strongly suggest that lipid droplets containing HCV core proteins may participate in the RNA metabolism of the host and/or HCV in HCV-infected cells. Furthermore, the findings that DDX1 is overexpressed in cell lines derived from tumors such as retinoblastomas and neuroblastomas (75), and that cellular expression of DDX3 induces anchorage-independent cell growth (76) suggest the involvement of DDX1 and DDX3 in carcinogenesis.

Some groups recently reported profiles of mRNAs upor down-regulated by expression of the HCV core protein (77–79), but these mRNAs included no molecules identified as lipid droplet proteins in this study. Since lipid droplets are a minor organelle in cells, it might be difficult to detect changes in the mRNA expression levels of lipid droplet proteins. The merits of targeted proteomic study are that it is possible to focus on minor cellular fractions, and also to detect changes in the intracellular distributions of proteins. Actually, the mRNA expression levels of TIP47, DDX1, and DDX3 did not change in Hep39 cells (data not shown).

We identified many other lipid droplet proteins found in either Hepswx or Hep39 cells, but their biological functions remain mostly unknown (Tables 1 and 2). Elucidation of the biological functions of these proteins will lead to an advanced understanding of the pathogeneses of HCV-derived liver diseases.

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