## Full-length article

# Proteomic analysis of HuH-7 cells harboring *in vitro*-transcribed fulllength hepatitis C virus 1b RNA<sup>1</sup>

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### Key words

proteomics; hepatitis C virus; *in vitro*transcribed RNA; HuH-7-hepatitis C virus; HuH-7 mock cells

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#### Abstract

Aim: The present study examined the differential expression of proteins in HuH-7 cells and HuH-7 cells harboring in vitro-transcribed full-length hepatitis C virus 1b RNA (HuH-7-HCV), and elucidated the cellular responses to HCV replication. Methods: The protein profiles of matched pairs of HuH-7–HCV cells and HuH-7 mock cells were analyzed by 2-D electrophoresis (2DE). Solubilized proteins were separated in the first dimension by isoelectric focusing, and by 12.5% SDS-PAGE in the second dimension. The differential protein expression was analyzed by use of image analysis software to identify candidates for HCV infection-associated proteins. Results: In total, 29 protein spots showed increases and 25 protein spots showed decreases in signal in HuH-7-HCV cell 2DE profiles as compared with HuH-7 mock cells. In the next step, the 10 spots showing the greatest increase and the 10 spots showing the greatest decrease were excised from gels and the proteins present were identified by Matrix-Assisted Laser Desorption/Ionization Time of Flight Mass Spectrometer (MALDI-TOF MS) or MALDI-TOF/TOF MS. In total, 13 proteins were identified successfully. The potential significance of the differential expression due to HCV replication was discussed. Conclusion: Our study identifies changes in the proteome of HuH-7 cells in the presence of HCV replication and yields information of the mechanism of HCV pathogenesis. These results will be useful for the identification of HCV infection-associated proteins that could be molecular targets for treatment.

## Introduction

Hepatitis C virus (HCV) is a major worldwide health problem. In total, 70%–90% of people who become infected fail to have the virus removed and remain chronically infected with the risk of developing liver cirrhosis and hepatocellular carcinoma<sup>[1]</sup>. Thus far there is no prophylactic vaccine to prevent HCV infection, and the current treatment,  $\alpha$ -interferon in combination with ribavirin, is not satisfactory because of significant side-effects and resistance.

The genome of HCV is a single-stranded, positive-sense RNA molecule approximately 9600 nucleotides in length. It consists of a 5' untranslated region (UTR), a single large open reading frame, and a 3' UTR<sup>[2]</sup>. The 5' UTR is required for translation of positive-strand viral RNA and replication

of negative-strand RNA. The single open reading frame encodes a large viral polyprotein of ~3000 amino acids<sup>[3]</sup> which is cleaved co- and post-translationally into at least 10 individual polypeptides with the following gene order: 5'-C-E1-E2-p7-NS2-NS3-NS4A-NS4B-NS5A-NS5B-3'<sup>[4]</sup>. The 3' UTR is closely related to virus replication and assembly<sup>[5]</sup> and is comprised of 3 domains: a variable, type-specific region followed by a (U) C-rich sequence of variable length, and a highly-conserved 98 nucleotide sequence designated the X tail at the 3' end<sup>[6]</sup>. *In vivo* studies have demonstrated that the X tail is essential for replication<sup>[7]</sup>.

Because of the lack of efficient and accurate *in vitro* models, basic studies defining the role of regulatory elements for replication and testing novel anti-HCV substances have

been significantly hampered. Although infection of primary cells or cell lines with high-titer HCV-containing sera is possible, replication levels in these cultures are too low to allow detailed studies of HCV replication<sup>[8]</sup>. As an alternative, the development of selectable subgenomic HCV replicons has enabled the study of viral RNA replication in cell culture<sup>[9]</sup>. The selection of these RNA, which lack the complete structural region from gene C up to either p7 or NS2, was made possible by the insertion of the gene encoding neomycin phosphotransferase (neo) downstream of the HCV IRES, with translation of the HCV NS proteins directed by the IRES of Encephalomyocarditis virus. Although replication levels of these RNA were high, subgenomic HCV replicons were insufficient to study what happens in HCV-infected cells due to the lack of the HCV structural proteins and potential effects on cellular metabolism that could be associated with their presence. A genome-length HCV-RNA replication system, however, would allow in-depth study of the alterations that HCV-infected cells undergo. Several genomelength HCV-RNA replication systems, using the N, Con-1, O, and H77 strains, have been reported<sup>[10-12]</sup>. However, at present, the 3' end of the HCV genome sequence in GenBank generally terminates at the polyadenylation site. Therefore, previous work has not taken into account the final 3' 98 conserved nucleotides in the construction of cDNA of the HCV genome. This may be one of the main causes as to why whole HCV replicon cannot be efficiently replicated in such experiments.

In order to ensure that HCV-RNA was able to replicate, transcribe, and synthesize proteins effectively in cell culture, the complete genome of HCV was used, including the 98 nucleotides at the 3' end in the present study. Our study is to discuss the proteomic techniques have been applied to globally analyze the protein expression profiles of a human hepatocarcinoma cell line HuH-7 harboring *in vitro*transcribed HCV 1b full length RNA (HuH-7–HCV) including the X tail. This work was aimed at elucidating the components of HCV replication and the cellular responses to HCV replication.

#### Materials and methods

**Cells** HuH-7 cells were grown in the presence of 5% CO<sub>2</sub> in Dulbecco's modified Eagle's medium with high levels of glucose (0.45%) supplemented with 10% fetal calf serum and antibiotics (100 U/mL penicillin and 100 mg/mL streptomycin).

**RNA preparation and transfection** A recombinant expression plasmid, pSP72-HCV, containing the full-length HCV 1b genome, including the 98 nucleotides at the 3' end was

constructed as described<sup>[13]</sup>. The plasmid DNA was extracted by phenol/chloroform extraction and ethanol precipitation. The HCV genomic RNA was transcribed by using MEGAscript TM T7 kit (Ambion, USA) in accordance with the manufacturer's protocol. For a 20  $\mu$ L reaction mixture, transcription was stopped by the addition of 4 U of DNase I and the mixture was incubated for 45 min at 37 °C. RNA was extracted with 1 mL TRIzol reagent (Invitrogen, USA) and precipitated with 500  $\mu$ L isopropanol. RNA pellets were washed once with 75% ethanol and dissolved in RNase-free water. To remove the residual amount of template DNA, RNA preparations were extracted once with acid phenol, precipitated with ethanol, and resuspended in RNase-free water.

HuH-7 cells were transfected with the RNA by electroporation and designated as HuH-7-HCV. Briefly, 5 µg RNA was mixed with  $2 \times 10^6$  cells suspended in 500 µL phosphate-buffered saline (PBS) in a cuvette with a gap width of 0.2 cm (Bio-Rad, Hercules, CA, USA). Electroporation consisted of 2 pulses of currents delivered by the Gene pulser II electroporation device (Bio-Rad, USA), which was set at 1.5 kV, 25 µF and maximum resistance. Mock-transfected (electroporation without HCV genomic RNA) cells were used as a control and designated as HuH-7 mock. The culture media of both HuH-7-HCV and HuH-7 mock cells were removed completely after 24 h and stored at 4 °C for subsequent assays. The cells were washed twice with PBS prior to refeeding with fresh culture media. The culture media of HuH-7-HCV and HuH-7 mock cells were collected at 48, 72, and 96 h post-transfection respectively. At each time point, 3 samples from different plates were collected.

**Detection of HCV accumulation in culture media by FQ– PCR and Western blotting** HCV-RNA titers in the culture media were analyzed by a HCV fluorescence PCR diagnostic kit (DaAn Gene, China).

HCV core and the NS3 protein were detected by Western blotting. The 96-h post-transfection culture media of both HuH-7–HCV and HuH-7 mock cells were separated by 10% SDS–PAGE, electrotransferred to nitrocellulose membranes (Millipore, USA), and blocked with 5% non-fat milk in TBST. The membranes were immunoblotted with rabbit anti-HCV core (1:200) and an anti-HCV–NS3 polyclonal antibody (1: 200; BIOS, China). Secondary antirabbit antibodies conjugated to horseradish peroxidase (BIOS, China) were applied at a dilution of 1:2000. The signals were detected by an ECL detection reagent (Pierce, USA).

**2-D electrophoresis** At 96 h post-transfection, the HuH-7– HCV and HuH-7 mock cells were harvested and dissolved in lysis buffer [7 mol/L urea, 2 mol/L thiourea, 1% Triton X-100, 4% (*w/v*) CHAPS, 4% Tris-base, 40 mmol/L dithiothreitol, 2% pharmalyte, 1 mmol/L phenylmethyl-sulfonyl fluoride, 1 g/L DNase I and 0.25 g/L RNase A]. The lysates were vortexed and incubated at 37 °C for 1 h and centrifuged at  $1000 \times g$  for 30 min at 4 °C. First-dimension isoelectric focusing was carried out using an Ettan IPGphor isoelectric focusing system, as described by the manufacturer (GE Amersham). Samples containing up to 100 µg protein were added to 13 cm IPG strips (pH 3–10). The strips were then applied by rehydration at 30 V for 12 h, and 1 h at 500 V. A gradient was then applied at 1000 V for 1 h, and 8000 V for 8 h. The temperature throughout this process was maintained at 20 °C. Separation in the second dimension was carried out at a current setting of 15 mA/gel for the initial 20 min and 30 mA/gel until the bromophenol blue front reached the bottom of the gels. To account for experimental variation, 3 batches of total proteins extracted from cells were subjected to 2-D electrophoresis (2DE) and replicate gels were simultaneously run 3 times. The gels were stained, respectively, with Coomassie brilliant blue and silver nitrate, as described previously<sup>[14]</sup>. Protein patterns in the gels were recorded as digitalized images using UMax powerlook 2110XL (GE Amersham, USA) scanner.

**In-gel protein digestion** In-gel digestion of proteins was carried out as follows. The spots of interest were excised and put into 1.5-mL microtubes. Samples were destained with 200–400  $\mu$ Lof 100 mmol/LNH<sub>4</sub>HCO<sub>3</sub>/30% ACN and dried in a vacuum centrifuge. The dried gel pieces were rehydrated in 5  $\mu$ L of 10 ng/ $\mu$ L trypsin (TPCK-treated, proteomics grade, Promega, USA), and incubated at 4 °C for 30–60 min. After digestion, 30  $\mu$ L of 25 mmol/L NH<sub>4</sub>HCO<sub>3</sub> was added, and the tubes were incubated at 37 °C for 20 h. The supernatants were recovered, and the extractions were repeated with 100  $\mu$ L of 60% ACN/0.1% TFA. The extracted solutions were mixed and dried in a vacuum centrifuge.

MALDI-TOF MS analysis The peptide mixtures were

solubilized with 5 µL of 1% formic acid, then vortexed and centrifuged. Samples  $(2 \mu L)$  were spotted on the target and dried. In total, 0.5 µL saturated acyano-4-hydroxy-transcinnamic solution was spotted on the samples and air dried. Spots were then analyzed in a Bruker-Daltonics AutoFlex TOF-TOF LIFT mass spectrometer (Bruker, Germany). The parameters were set up as follows: positive ion-reflector mode, accelerating voltage 20 kV. A trypsin-fragment peak served as an internal standard for mass calibration. A list of the corrected mass peaks was obtained as PMF. Proteins were identified from PMF data using MASCOT software (http:// www.matrixscience.com). The following search parameters were applied: protein mass unrestricted; peptide mass tolerance±100 ppm, fragment mass tolerance±0.8 Da, number of missed cleavage sites allowed 1, cysteine residue modified as carbamidomethyl-cys, variable modifications oxidation (M), mass values MH+, and monoisotopic. NCBI were used as the protein sequence databases.

Real-time RT-PCR and Western blotting analysis for differentially expressed proteins To validate the MS results, another set of cells was isolated for total mRNA and subsequent real-time RT-PCR. The cDNA was synthesized by random hexamer primed reverse transcription using the RevertAid first strand cDNA synthesis kit (Fermentas, Canada). Real-time PCR was performed using the ABI 7500 real-time PCR system and SYBR premix ex Taq (perfect real time; TaKaRa, Dalian, China), primarily following the manufacturers' protocol. In brief, the reaction mixture (25 µL total volume) contained 500 ng cDNA, primers (Table 1) at 0.2 µmol/L final concentration and 12.5 µL SYBR premix ex Taq. Thermal cycling conditions were 95 °C for 10 s and 40 cycles at 95 °C for 5 s and 60 °C for 34 s. The experiments were performed in triplicate in the same reaction. Human GAPDH gene was amplified as an internal control in the same reac-

Table 1.	Primers	and	PCR	condition	for	each	gene.
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Gene name	Primers (5'-3')	Annealing/°C	Product/bp	
GRP 78	Sense: CTGTGCAGCAGGACATCAAGTTC	64	92	
	Antisense: TGTTTGCCCACCTCCAATATCA			
SLP-2	Sense: GCTGAACAGATAAATCAGGCAGCA	64	142	
	Antisense: GCTCGGCCACAGTCAGTGAA			
Keratin 9	Sense: CCTCAATGACATGCGTCAGGA	64	138	
	Antisense: TTGGCACTGGACTGCACCTC			
Regucalcin	Sense: CACCAAGCAAGTACAGCGAGTGA	64	148	
	Antisense: ATCCACCGTGGCCAAGACA			
GAPDH	Sense: GCACCGTCAAGGCTGAGAAC	64	138	
	Antisense: TGGTGAAGACGCCAGTGGA			

tion with target genes. Comparative quantification was determined using the  $2^{\Delta Ct}$  method<sup>[15]</sup>.

An equal quantity of protein was used for the Western blotting analysis of the glucose-regulated protein 78 kDa (GRP 78) and keratin 9 protein expressions. Cells were lysed in RIPA buffer with protease inhibitors. Western blot analysis was performed as described earlier, and  $\beta$ -actin was used as an internal control. Rat anti-GRP 78 and goat anticytokeratin 9 (Santa Cruz, Santa Cruz, CA, USA) were diluted at 1:500, while a rabbit anti- $\beta$ -actin polyclonal antibody (BIOS, China) was diluted at 1:200. Secondary antirat, antigoat, and antirabbit antibodies conjugated to horseradish peroxidase (BIOS, China) were applied at 1:2000 dilution.

**Statistical analysis** The expression levels of the proteins were quantified by analyzing the intensity of each spot with PDQuest software (Bio-Rad, USA). The differences in the protein expression levels between HuH-7–HCV and HuH-7 mock cells were analyzed by Student's *t*-test or Mann-Whitney *U*-test. Probability values of less than 0.05 were considered significant. All analyses were carried out using SPSS 10.0 statistical software (SPSS, Chicago, IL, USA).

#### Results

HCV-RNA and protein accumulation in HuH-7–HCV culture media Following transfection, increasing titers of HCV-RNA ranging from a few copies to 1×10<sup>6</sup> copies/mL were detected in the HuH-7–HCV cell culture media by fluorescence RT–PCR assays. These levels increased significantly over with time post-transfection. In contrast, in the HuH-7 mock culture supernatants, the HCV titers were <80 copies/mL at 24, 48, 72, and 96 h post-transfection, respectively (Figure 1A). Similarly, HCV core and the NS3 protein were detected in the culture media of HuH-7–HCV by Western blotting, but not in that of HuH-7 mock cells (Figure 1B). These results demonstrated that HCV-RNA was efficiently replicated in the *in vitro* culture system.

Proteomic analysis of differentially-expressed proteins The proteome of the HuH-7–HCV and HuH-7 mock cells was compared using 2DE (Figure 2). Reproducible highresolution 2DE profiles were obtained. The average numbers of spots representing either individual or similarly migrating protein species obtained with both HuH-7–HCV and HuH-7 mock cells were  $2618\pm111$  and  $2410\pm248$ , respectively. Fifty-four spots were found to be significantly altered (*P*< 0.05). Among them, the intensities of 29 protein spots were increased and the intensities of 25 protein spots were decreased in HuH-7–HCV cell extracts as compared to those obtained with HuH-7 mock cell extracts. The 10 spots show-



**Figure 1.** Profile of HCV-RNA accumulation in the HuH-7–HCV culture media. (A) HuH-7 cells were transfected with HCV 1b fulllength RNA including the 3' X tail. Culture media of HuH-7–HCV and HuH-7 mock cells were harvested at 24, 48, 72 °C and 96 h posttransfection and analyzed for HCV-RNA titer by fluorescence RT– PCR. HCV-RNA titers in HuH-7–HCV culture media ranged from a few copies to  $1\times10^6$  copies/mL. HCV titers in HuH-7 mock culture supernatants were <80 copies/mL at all time points. (B) 96-h posttransfection culture media of HuH-7–HCV and HuH-7 mock were examined for the presence of HCV core and NS3 protein by Western blot analysis. Both HCV proteins were detected the HuH-7–HCV cell supernatant, but not in that of HuH-7 mock-transfected cells.

ing the greatest increase in signal and the 10 spots showing the greatest decrease in signal were chosen for further analysis.

The spots of interest were excised from the 2D gels and subjected to MALDI–TOF MS or MALDI–TOF/TOF MS. The resulting PMF data were used to search the NCBI databases via MASCOT software. The identity of specific proteins was determined by comprehensively comparing the corresponding experimental pI, Mr, the number of matched peptides, and the sequence coverage. Upregulated proteins in HuH-7–HCV cells included keratin 9, keratin 8 isoform CRA\_a, keratin 1, p47 protein isoform c, stomatin (EPB72)-like 2 (SLP-2), and regucalcin. Downregulated proteins included heat shock 70 kDa protein 5, heat shock 70 kDa protein 5 glucose-regulated protein 1 isoform CRA\_c, MTHSP75, and Annexin V (Table 2).

Verification of specific differentially-expressed proteins by real-time RT–PCR and Western blotting The differential expression of the genes encoding GRP 78, stomatin-like



Figure 2. Representative 2-D gel analyses of HuH-7-HCV and HuH-7 mock transfected solubilized protein extracts.

Table 2.	Differentially-expressed	proteins in	HuH-/-HCV	cells identified	by MALDI-	TOF or MALDI-I	OF/TOF MS.

Spot no.	Protein name	Accession (Gi no.)	Predicted mass (Da)	Predicted pI	Score <sup>a</sup>	Alteration of protein expression
2410	Keratin 9	gi 55956899	62255	5.5	91	+
923	Keratin 8, isoform CRA a	gi 119617057	57829	6	172	+
1667	Keratin 1	gi 11935049	66198	6	66	+
1855	p47 protein isoform c	gi 33286434	28505	6.5	67	+
1405	Stomatin (EPB72)-like 2	gi 7305503	38624	5.5	123	+
1661	Regucalcin	gi 4759036	33802	6.5	87	+
652	Heat shock 70 kDa protein 5	gi 16507237	72402	5.5	73	_
656	Heat shock 70 kDa protein 5	gi 16507237	72402	5.5	233	_
1137	Heat shock 70 kDa protein 5 glucose-regulated protein 78 kDa, isoform CRA b	gi 119608027	51197	5.5	112	-
2031	BiP protein	gi 6470150	71002	5	94	
752	Heat shock 60 kDa protein 1 (chaperonin), isoform CRA c	gi 119590557	41067	5.5	152	-
639	MTHSP75	gi 292059	74019	6	84	
1842	Chain, Annexin V (Lipocortin V, endonexin ii, placental anticoagulant protein)	gi 999926	35839	5.5	94	_

<sup>a</sup>Protein scores greater than 65 are significant (P < 0.05).

2, keratin 9, and regucalcin were validated by real-time RT– PCR (Figure 3A). The differential expression of the GRP 78 (Figure 3B) and keratin 9 (Figure 3C) proteins was verified by Western blotting. The changes in the expression levels of these selected proteins were consistent with the 2DE results. Such results demonstrate that the proteomic analysis of HuH-7–HCV and HuH-7 mock cells can reliably identify cellular proteins that are differentially expressed as the result of the replication and expression of the HCV genome.

# Discussion

In order to establish protein expression profiles and to characterize HCV infection-related proteins, HuH-7 cells were transfected with *in vitro*-transcribed HCV full length RNA. The differential expression proteins of HuH-7 harboring HCV-RNA and HuH-7 mock cells were analyzed by 2DE and MS.



**Figure 3.** (A) Verification of GRP 78, SLP-2, keratin 9, and regucalcin gene expression in 96 h post-transfection HuH-7 mock and HuH-7– HCV cells by real-time RT–PCR. Relative quantification of each gene was calculated by the  $2^{ACt}$  method where the HuH-7 mock control levels were normalized to 1. (B) Western blot analysis of the GRP 78 protein expression in HuH-7 mock and HuH-7–HCV cells. Results show that the expression of the GRP 78 protein decreased in HuH-7– HCV cells. (C) Western blot analysis of the keratin 9 protein expression in HuH-7 mock and HuH-7–HCV cells. Results show that the keratin 9 expression was increased in HuH-7–HCV cells.  $\beta$ -Actin was used as an internal loading control.

The identified differentially-expressed proteins included proteins associated with the host cell cytoskeleton, intracellular trafficking, molecular chaperones, apoptosis, protein degradation, oxidative and endoplasmic reticulum (ER) stress, cell cycle progression, antigen presentation, and calcium homeostasis.

Several members of the keratin family (keratin 1, 9, and 8) were upregulated after HCV-RNA transfection. Keratins are proteins that form the intermediate filament cytoskeleton and are essential for normal tissue structure and function. Keratin 8 is a type II keratin and occurs in many epithelial cells, including hepatocytes. Several isoforms of keratin 8 generated by differential splicing have been identified. The majority of keratin 8 is assembled with its partner, keratin 18, into highly insoluble 10 nm filaments that extend from the nucleus to the internal leaflet of the plasma membrane<sup>[16]</sup>. The function of keratin 8/18 in the liver is protection from mechanical and non-mechanical forms of stress<sup>[17]</sup>, which is supported by the association of several K8/18 mutations with cryptogenic and non-cryptogenic forms of human liver disease<sup>[18]</sup>. It has been documented that keratin 8/18 has been linked to tumor necrosis factor-induced and Fas-mediated apoptosis<sup>[19]</sup>. There is increasing evidence suggesting that liver cell damage in chronic HCV infection is mediated by the induction of apoptosis<sup>[20]</sup>. Engulfment of apoptotic bodies by hepatic

stellate cells stimulates the fibrogenic activity of these cells and may be one mechanism by which hepatocyte apoptosis promotes fibrosis<sup>[21]</sup>. Thus our findings indicated that the upregulation of keratin 8 agreed with this proposed mechanism.

HCV utilizes the host cell ER as the main site for replication and protein synthesis. Recent studies have been demonstrated that HCV proteins have the propensity to induce distinct alterations of ER membranes<sup>[22]</sup> and homeostasis, and cause some level of ER stress. The ER stress response works at the level of protein translation and appears to be regulated by the level of stress. Mild ER stress inhibits protein synthesis initiation and can slow cell growth, while prolonged ER stress can lead to apoptosis. These events correlate with the transcriptional induction of the ER chaperone GRP 78, a typical marker of ER stress. Heat shock 70 kDa protein 5 GRP 78 is also known as heat shock 70 kDa protein 5 and immunoglobulin binding protein (BiP). It belongs to the Hsp70 multigene family. In humans, the Hsp70 multigene family consists of at least 4 members: Hsp70, GRP 78, MTHSP75, and Hsc70. The molecular chaperone GRP 78 is important in the folding, maturation, and transport of proteins out of the cell, and GRP 78 is critical for the unfolded protein response required to alleviate ER "stress", maintain ER function, and protect cells against death<sup>[23]</sup>. It works as a coreceptor for viruses and MHC I antigen presentation, and as a receptor for activated forms of the plasma proteinase inhibitor  $\alpha_2$ -macroglobulin<sup>[24]</sup>. GRP 78 is also identified as a keratin-associated protein. The GRP 78-keratin 8/18 association is induced by tunicamycin or glucose starvation. The physiological relevance of keratin 8/18-GRP 78 binding remains to be determined. The location of GRP 78 in the ER makes any association of K8/18 with GRP 78 significant only under conditions of ER damage, or export of resident luminal ER proteins into the cytosol or the cytoplasmic resident soluble keratins into the ER<sup>[25]</sup>. The synthesis of HCV proteins may influence cell fate through the regulation of the ER stress signaling pathway and apoptotic response.

A mitochondrial chaperone, heat shock 60 kDa protein, was downregulated in the HuH-7–HCV cells. This chaperone may protect mitochondria from oxidative stress by facilitating the proper assembly of mitochondrial proteins<sup>[26]</sup>. The downregulation of this chaperone has been reported in various tumor series correlating with disease outcome<sup>[27]</sup>. Acute ablation of Hsp60 results in destabilization of survivin levels, and nearly complete and selective loss of the mitochondrial pool of survivin<sup>[28]</sup>, thus abrogating this anti-apoptotic response<sup>[29]</sup>. MTHSP75 is another mitochondrial chaperone for the import and folding of newly-synthesized nuclear and mitochondrial-encoded proteins<sup>[30]</sup>. It has been reported to be involved in antigen processing and presentation<sup>[31]</sup>. Therefore, the downregulation of MTHSP75 may contribute to the cytopathic effect on mitochondria induced by HCV and possibly impairs antigen presentation.

There is no previous report concerning the relationship between HCV infection and the SLP-2, regucalcin, Annexin V, and p47 proteins. SLP-2 is a novel stomatin homolog and is mainly localized in the cytoplasm with some distribution on the membrane<sup>[32]</sup>. SLP-2 is a cancer-related gene whose product promotes cell growth, tumorigenicity, and adhesion in human esophageal squamous cell carcinoma, endometrial adenocarcinoma, and laryngeal squamous cell carcinoma. Furthermore, the SLP-2 expression correlates with the clinical stage of carcinoma. Zhang et al showed that SLP-2 was overexpressed in all the tumor types they tested compared with their normal counterparts<sup>[33]</sup>. The etiology of the increased SLP-2 expression in human cancers is unknown. Possible mechanisms include point mutation, gene amplification, gene rearrangement, and insertion of a strong promoter or enhancer. Epigenetic modifications, including demethylation and deacetylation may also be responsible.

Regucalcin (senescence marker protein-30) is a multifunctional protein that is expressed in multiple tissues, including hepatocytes and renal tubular epithelia<sup>[34]</sup>. Regucalcin regulates calcium homeostasis for the maintenance of hepatocyte morphology and differentiation, inhibits cell death, and provides protection againt cell injury and oxidative stress by enhancement of  $Ca^{2+}$  efflux to either the extracellular space or intraorganellar spaces through membrane  $Ca^{2+}$  pump activity<sup>[35]</sup>. In the present study, the upregulation of regucalcin may be the response of the cell to the injury caused by HCV.

Annexin V is a member of the Annexin family of structurally-homologous proteins interacting with anionic phospholipid membranes in a reversible calcium-dependent manner<sup>[36]</sup>. It was shown to interact with membrane proteins and to regulate the sodium–calcium exchanger<sup>[37]</sup>. In a previous study, Annexin V was abundant in normal mucosa compared with neighboring tumor tissue<sup>[38]</sup>. P47 is an adaptor molecule of the cytosolic ATPase p97. The principal role of the p97–p47 complex is known to be involved in the heterotypic fusion of transport vesicles with their target membranes, the homotypic fusion of membrane compartments and ubiquitin-dependent protein degradation<sup>[39]</sup>. In the present study, the level of the p47 protein was higher than the control and it is worth further study to elucidate the role of p47 in HCV replication.

In summary, the progression of HCV infection involves various cellular and molecular mechanisms. These results

provide a further understanding of the molecular mechanism of cellular events and pathogenesis associated with viral progression, and are useful for the identification of HCV infection-associated proteins which will be molecular targets for treatment.

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#### Author contribution

Meng XUN, Si-hai ZHAO, Yong-lie CHU designed research; Meng XUN, Si-hai ZHAO, Chun-xia CAO, Juan SONG performed research; Meng XUN, Si-hai ZHAO, Yonglie CHU contributed new reagents or analytic tools; Meng XUN, Si-hai ZHAO, Ming-ming SHAO analyzed data; Meng XUN, Juan SONG, Yong-lie CHU wrote the paper.

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