Quantitative Analysis of Severe Acute Respiratory Syndrome (SARS)-associated Coronavirus-infected Cells Using Proteomic Approaches

IMPLICATIONS FOR CELLULAR RESPONSES TO VIRUS INFECTION*

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We present the first proteomic analysis on the cellular response to severe acute respiratory syndrome-associated coronavirus (SARS-CoV) infection. The differential proteomes of Vero E6 cells with and without infection of the SARS-CoV were resolved and quantitated with two-dimensional differential gel electrophoresis followed by ESI-MS/MS identification. Moreover isotope-coded affinity tag technology coupled with two-dimensional LC-MS/MS were also applied to the differential proteins of infected cells. By combining these two complementary strategies, 355 unique proteins were identified and quantitated with 186 of them differentially expressed (at least 1.5-fold quantitative alteration) between infected and uninfected Vero E6 cells. The implication for cellular responses to virus infection was analyzed in depth according to the proteomic results. Thus, the present work provides large scale protein-related information to investigate the mechanism of SARS-CoV infection and pathogenesis. Molecular & Cellular Proteomics 4:902–913, 2005.

A new type of coronavirus was reported as the causal agent of severe acute respiratory syndrome (SARS)¹ in April 2003, and the genome of the SARS-CoV was sequenced by several groups (1–3). The properties of the SARS-CoV genome was analyzed in depth by bioinformatic tools (4, 5). In addition, several important works on the proteins of SARS-CoV have been reported recently, including the identification of SARS-CoV 3C-like protease structure (6) and the identification of angiotensin-converting enzyme 2 as a functional receptor for the spike protein (7). In our recent work, we identified all of the predicted SARS-CoV structural proteins, nucleocapsid (N), membrane (M), spike (S), and envelope (E), using proteomic approaches and found a novel protein, SARS-CoV 3a (8, 9).

To uncover the mechanisms of cellular responses to the virus infection and identify potential drug targets of antiviral treatment, it is very useful to study the molecular profiling of virus-infected cells with high throughput and quantitative approaches. Analysis of gene expression profiles during viral infection is one of the powerful approaches to probe potential cellular genes involved in viral infection and pathogenesis (10). The recent development of proteomic analytic technology such as differential gel electrophoresis (DIGE) (11, 12) and ICAT (13, 14) also provides new tools for such studies.

As a method based on two-dimensional (2D) electrophoresis, DIGE allows two or three independent samples labeled with different fluorescent dyes such as cyanine-2 (Cy2), cyanine-3 (Cy3), and cyanine-5 (Cy5) to be run in one gel simultaneously and viewed individually using the different fluorescent properties of Cy2, Cy3, and Cy5, circumventing some of the reproducibility problems associated with 2D electrophoresis and providing more accurate quantitative information compared with other staining methods such as silver staining with the dynamic range over 3–4 orders of magnitude (11, 12). More recently, the combination of stable ICAT, LC, and MS/MS has emerged as an alternative gel-free quantitative proteomic technology (13, 14). In ICAT analysis, two pools of proteins are labeled respectively with isotopically light and heavy ICAT regents, which are chemically identical and therefore serve as a good internal standard for accurate quantification. Although LC-MS provides quantitative information based upon the relative abundances of the heavy and light peptides, LC-MS/MS provides qualitative information based upon the peptide molecular mass and amino acid sequence information. These two technologies have been proved to be accurate.

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¹ The abbreviations used are: SARS, severe acute respiratory syndrome; SARS-CoV, SARS-associated coronavirus; 2D, two-dimensional; DIGE, differential gel electrophoresis; hnRNP, heterogeneous nuclear ribonucleoprotein; 2DE, two-dimensional electrophoresis; Cy, cyanine; DMEM, Dulbecco’s modified Eagle’s medium; FBS, fetal bovine serum; RP, reverse phase; MHV, mouse hepatitis virus; EF, elongation factor; eIF, eukaryotic initiation factor.

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provide 65 mM DTT to reduce the labeled samples for 15 min at room temperature. Then the samples were subjected to 2D-PAGE in the dark. The DIGE gels were scanned using Typhoon Variable Mode Imagers 9400 (Amersham Biosciences). DeCyder (Amersham Biosciences) software was used for image analysis.

In-gel Trypsin Digestion—Changes observed in 2D-DIGE images were aligned with Phastgel Blue R-stained protein patterns in preparative gels. Spots of interest were cut manually. Gel pieces were digested as described by Yu et al. (17).

**RP-HPLC-ESI-MS/MS Analysis**—RP-HPLC was performed using a Surveyor LC system (Thermo Finnigan, San Jose, CA) on a C18 column (RP, 180 μm × 150 mm, BioBasic® C18, 5 μm, Thermo Hypersil-Keystone). The pump flow rate was split 1:120 to achieve a column flow rate of 1.5 μl/min. Mobile phase A was 0.1% formic acid in water, and mobile phase B was 0.1% formic acid in acetonitrile. The tryptic peptide mixtures were eluted using a gradient of 2–98% B over 60 min.

The mass spectral data shown here were acquired on an LTQ linear ion trap mass spectrometer (Thermo Finnigan) equipped with an electrospray interface operated in positive ion mode. The temperature of the heated capillary was set at 170 °C. A voltage of 5.4 kV applied to the ESI needle resulted in a distinct signal. Normalized collision energy was 35.0. The mass spectrometer was set so that one full MS scan was followed by three MS/MS scans on the three most intense ions from the MS spectrum with the following Dynamic Exclusion™ settings: repeat count, 2; repeat duration, 0.5 min; exclusion duration, 2.0 min.

**ICAT Analysis**—ICAT analysis was performed using the Cleavable ICAT™ reagent kit (Applied Biosystems, Foster City, CA) according to the manufacturer’s guidelines with some modifications. For ICAT analysis, the cellular samples were precipitated and resolubilized in denaturing buffer (6 M guanidine hydrochloride, 100 mM Tris-Cl, pH 8.3). 100 μg of the E6 or E6-V protein sample in 80 μl of denaturing buffer were reduced at 37 °C for 2 h with 5 mM tributylphosphine (Bio-Rad) and alkylated at 37 °C for 2 h in the dark with ICAT-light and ICAT-heavy reagent, respectively. The samples were digested with trypsin at 37 °C for 20 h. Then the ICAT-labeled peptides were purified using the kit of ICAT™ Avidin Buffer Pack and Avidin Affinity Cartridge (Applied Biosystems) according to the manufacturer’s guidelines.

**2D LC-MS/MS**—Orthogonal 2D LC-MS/MS was performed using a ProteomeX work station (Thermo Finnigan). The system was fitted with a strong cation exchange column (320 μm inner diameter × 100 mm, DEV SCX, Thermo Hypersil-Keystone) and two C18 RP columns (180 μm × 100 mm, BioBasic® C18, 5 μm, Thermo Hypersil-Keystone). The salt steps used were 0, 25, 50, 75, 100, 150, 200, 400, and 800 mM NH4Cl synchronized with nine 140-min RP gradients. RP solvents were 0.1% formic acid in either water (A) or acetonitrile (B). The setting of the LCO Deca Xplus ion trap mass spectrometer is as follows. One full MS scan was followed by three MS/MS scans on the three most intense ions from the MS spectrum according to the following Dynamic Exclusion settings: repeat count, 1; repeat duration, 0.5 min; exclusion duration, 3.0 min.

**Data Base Searching**—The acquired MS/MS spectra were automatically searched against the nonredundant human protein data base (NCBI) (www.ncbi.nlm.nih.gov), December 4, 2003 release) using the TurboSEQUEST program in the BioWorks™ 3.1 software suite. For protein spot identification, an accepted SEQUEST result had to have a Xcorr score of at least 0.1 (regardless of charge state). Peptides with a +1 charge state were accepted if they were fully tryptic and had a cross correlation (Xcorr) of at least 1.8. Peptides with a +2 charge state were accepted if they had an Xcorr >2.5. Peptides with a +3 charge state were accepted if they had an Xcorr >3.7. For ICAT analysis, protein identification and quantification were achieved by using SEQUEST and EXPRESS software tools. Peptides with a +1...
charge state were accepted if they were fully tryptic and had an Xcorr of at least 1.5. Peptides with a +1 charge state were accepted if they had an Xcorr > 1.5. Then the confirmation of protein identification and quantification of the peptides was further analyzed manually as described by Han et al. (14). Protein abundance ratios larger than 1.5 or smaller than 0.67 were set as a threshold indicating significant changes (18).

Western Blotting—SDS-PAGE-separated proteins were transferred onto a nitrocellulose membrane (Protran, Schleicher & Schuell) on a Mini Trans-Blot Cell (Bio-Rad). The transferred membrane was blocked with 1 × NET-Gelatin (150 mM NaCl, 5 mM EDTA, pH 8.8, 50 mM Tris-HCl, 0.05% Triton X-100, 8.3% gelatin) at room temperature for 1 h and then was incubated with corresponding primary antibodies (anti-HSP90 (rabbit), Santa Cruz Biotechnology, 1:5,000; anti-α-tubulin (mouse), Sigma, 1:600,000; anti-stathmin (rabbit), Calbiochem-Novabiochem, 1:5,000; anti-vimentin (mouse), Sigma, 1:1,000) for 1 h at room temperature, and then the membranes were incubated with the corresponding secondary antibodies. After being detected with ECL PLUS (Amersham Biosciences) according to the manufacturer's instructions, the membrane was scanned using a Typhoon Scanner 9400 (Amersham Biosciences).

Bioinformatic Annotation—The theoretical pI and molecular mass values of proteins were defined by the program pepstats (www.hgmp.mrc.ac.uk/Software/EMBOSS). The protein function and subcellular location annotation was from the Swiss-Prot and TrEMBL protein data base (us.expasy.org/sprot/).

RESULTS

2D-DIGE Analysis of the SARS-CoV-infected and Uninfected Vero E6 cells—2D-DIGE as a qualitative and quantitative proteomic approach was performed to determine the differential proteomes of the SARS-CoV-infected and uninfected Vero E6 cells. The lysate from infected cells was labeled with Cy5, whereas the lysate from uninfected cells was labeled with Cy3. After electrophoresis and imaging, the Cy3 and Cy5 images were false colored in red and green, respectively, and two images were overlapped (Fig. 1). The biological
The variation analysis mode of DeCyder was used for comparing the overlay image. The standardization was done by the comparison of normalized Cy3 and Cy5 protein spot volumes with the corresponding Cy2 standard spot volumes within each gel. Protein spots with an average ratio value greater than 1.2-fold and a t test p value <0.05 were selected for mass spectrometric identification. A total of 63 proteins belonging to 48 unique gene products were identified with ESI-LC-MS/MS (Supplemental Table I). Among those proteins, 17 proteins (21 protein spots) were down-regulated, whereas 13 proteins (15 protein spots) were up-regulated by at least 1.5-fold in SARS-CoV-infected cells (Table I). The identified protein spots with at least a 2.0-fold quantitative alteration in SARS-CoV-infected Vero E6 cells are listed in Table II. The differentially expressed proteins are involved in various functions including cytoskeleton, actin-associated network, metabolic enzymes, and signal transduction (Table II and Supplemental Table I).

**ICAT Analysis of the SARS-CoV-infected and Uninfected Vero E6 Cells—** Recently a new proteomic approach, ICAT, has been developed to quantitatively analyze the protein differential expression, which is faster than the gel method. In the present work, the ICAT method coupled to 2D-LC-MS/MS was used for quantitative comparison of differential proteome profiles between SARS-CoV-infected cells and uninfected cells. Cysteines were labeled with light (12C) and heavy (13C) ICAT reagent, and labeled peptides were affinity-purified using an avidin column. The resultant peptides were first fra-
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| Protein description | GI no. | Subcellular location | Peptide hits | Unique peptides | Ratio, ICAT (L:H; E6:E6-V) | Peptide sequence (charge/Xcorr/ΔCn) |
|---------------------|-------|----------------------|--------------|-----------------|-----------------------------|-------------------------------------|
| Actin network protein | gi[31657094] | Cytoskeletal | 1 | 1 | 1:0.25 | R.FGERC*OEHSKESPAR.S (2+/2.05/0.20) |
| α-tubulin | gi[728751] | Cytoskeletal | 1 | 1 | 1:0.38 | K.C*QLEINFNTLQTK.L (2+/3.16/0.36) |
| Destrin (actin-depolymerizing factor) | gi[5802966] | Cytoskeletal | 4 | 1 | 1:2.77 | K.HEC*QANGPEDLNR.A (3+/3.34/0.37) |
| Channel or transporter protein | | | | | | |
| Potassium channel-modulatory factor 1 | gi[31543383] | No annotation | 1 | 1 | 1:2.59 | R.HEVHSV*AC*LK.G5 (2+/2.90/0.43) |
| Potassium channel subfamily K member 15 | gi[24636282] | Membrane | 1 | 1 | 1:2.97 | K.VFC*MFYALLGIPITLVFTQOSLGE.L (3+/2.52/0.11) |
| Sodium bicarbonate transporter 1 isoform a | gi[15042959] | No annotation | 1 | 1 | 1:8.84 | R.FFGGCL*LDIK.R (2+/2.19/0.20) |
| Chromosome-associated protein | | | | | | |
| DEK protein | gi[544150] | Nuclear | 1 | 1 | 1:12.45 | K.C*PEILSDESSSDEDK.K (3+/2.90/0.20) |
| High mobility group protein 4 | gi[20138144] | Nuclear | 1 | 1 | 1:2.20 | K.RPPSGLFLC*SEFRPK.L (3+/3.70/0.34) |
| Condensin subunit 2 | gi[30172801] | Cytoplasmic and nuclear | 1 | 1 | 1:2.21 | K.TAASFDCEC*STAGVFLSTHC*QDYR.S (3+/4.73/0.54) |
| Heat shock protein | | | | | | |
| HSP 90-α | gi[123678] | Cytoplasmic | 3 | 1 | 1:2.08 | K.HGLQIEVIMELIDKCY*VQQL.K (3+/2.58/0.11) |
| Stress-induced phosphoprotein 1 | gi[400042] | No annotation | 2 | 1 | 1:2.18 | K.DC*EEC*IQUEFTIK.G (2+/2.54/0.25) |
| Heterogeneous nuclear ribonucleoprotein | | | | | | |
| hnRNP D0 | gi[13124489] | Nuclear | 4 | 1 | 1:5.15 | R.GFC*FITFK.E (2/2.27/0.41) |
| hnRNP U | gi[6228894] | Nuclear | 4 | 1 | 1:1.77 | K.TC*NC*EDTVGKEF (2+/2.68/0.37) |
| hnRNP core protein A1 | gi[133254] | Nuclear | 6 | 1 | 1:1.90 | K.YHVTNGHC*EVRL.S (3+/3.58/0.24) |
| hnRNP I | gi[131528] | Nuclear | 2 | 1 | 1:1.95 | K.LSLDGNINYAC*TRIL.R (2+/3.37/0.41) |
| hnRNP K isoform a | gi[14165437] | Cytoplasmic and nuclear | 4 | 1 | 1:2.00 | K.LFQEC*PHSTDR.V (2+/3.55/0.45) |
| hnRNP A0 | gi[8134660] | Nuclear | 1 | 1 | 1:2.04 | R.GHFAEGITLDC*VVNVNPK.T.R (3+/2.98/0.43) |
| hnRNP D-like | gi[4885423] | No annotation | 8 | 1 | 1:2.18 | R.GFC*FITFDEEPVKK.L (3+/4.30/0.43) |
| hnRNP E2 | gi[6707736] | Nuclear | 1 | 1 | 1:2.27 | K.INSIGC*PERI.L (2+/2.84/0.29) |
| Intracellular trafficking protein | | | | | | |
| Translocation protein SEC63 homolog | gi[18203500] | Endoplasmic reticulum | 1 | 1 | 1:0.50 | R.APTLSLENCMKLQSMAV-OQLQFQP.S (3+/2.74/0.29) |
| Synaptophysin-like protein isoform a; pantophysin | gi[5803185] | Vesicular | 1 | 1 | 1:2.28 | K.GOTEIUVNC*PPAVTKNTATGYPF.R.L (3+/2.71/0.21) |
| α-Soluble NSF attachment protein | gi[6226705] | Membrane | 1 | 1 | 1:9.51 | K.C*LXXVAGYAALEQQY.K (3+/2.66/0.13) |
| Metabolic enzyme | | | | | | |
| Vacular ATP synthase catalytic subunit A | gi[22096378] | Endoplasmic reticulum | 1 | 1 | 1:0.36 | R.FCFPYYKTGMSINIAFMYD.MARRA. (3+/2.26/0.21) |
| Pyruvate dehydrogenase E1 component α subunit | gi[129063] | Mitochondrial | 1 | 1 | 1:0.36 | K.LPCIFICENR.Y (2+/2.13/0.23) |
| FK506-binding protein 4 | gi[399866] | Cytoplasmic and nuclear | 2 | 1 | 1:0.41 | K.VGEVCHITCKPEYAYGSAGSP.PK.I (3+/2.86/0.23) |
| Alanine-glyoxylate aminotransferase 2 | gi[17432913] | Mitochondrial | 1 | 1 | 1:0.45 | K.CLOHFNTGFGMNPMACIGSAVL.DIK.E (3+/2.77/0.16) |
| GMP reductase 2 | gi[25008511] | No annotation | 1 | 1 | 1:2.10 | K.GHIISSDG.C*CGQDVAK.A (3+/2.63/0.16) |
| Succinyl-CoA synthetase, α chain | gi[20141765] | Mitochondrial | 6 | 1 | 1:2.16 | R.LIGPN*CQPGVING.EK.I (2+/3.78/0.52) |
| Thioredoxin-like protein p46 | gi[29839560] | Endoplasmic reticulum | 19 | 2 | 1:2.44 ≥ 0.45 | K.VDC*TASHDSVC*SAGQVR.G (2+/4.18/0.50); K.VDC*TOTHYEC*SGQVR.G (3+/4.27/0.35) |
| UDP-glucose 6-dehydrogenase Protein degradation | gi[6175086] | No annotation | 1 | 1 | 1:3.56 | R.AVQAC.L*AYEHWVPR.E (3+/2.60/0.30) |
| Ovochymase | gi[34419641] | No annotation | 1 | 1 | 1:2.03 | R.YLDDYRGREC*SWLVR.V (3+/2.62/0.17) |
| Proteasome subunit P50 | gi[20532406] | Cytoplasmic and nuclear | 3 | 1 | 1:2.22 | R.C*TDDFNGAQQ*C.K.A (2+/3.16/0.45) |
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TABLE III—continued

| Protein description | GI no. | Subcellular location | Peptide hits | Unique peptides | Ratio, ICAT (L:H; E6:E6-V) | Peptide sequence (charge/Xcorr/ΔCn)a |
|---------------------|--------|---------------------|--------------|----------------|-----------------------------|--------------------------------------|
| Ribosomal proteinS | gi[4506725] | Ribosomal 11 2 | 1:1.52 ± 0.33 | K.LREC*PLPLIFLR.N (3/5/3.12/0.32) |
| 60 S ribosomal protein | gi[15431248] | Ribosomal 6 1 | 1:1.58 | K.FSVN*VLGDOQH*DEAK.A (2/2.37/0.27) |
| 40 S ribosomal protein | gi[14165646] | Ribosomal 1 1 | 1:1.61 | K.CV*GIVSP.R (2/2.10/0.11) |
| 40 S ribosomal protein | gi[4506699] | Ribosomal 2 1 | 1:1.65 | K.TYAI*CRAIL.R (2/2.27/0.25) |
| 40 S ribosomal protein | gi[15011946] | Ribosomal 6 1 | 1:1.66 | K.LHYVCS*SAH.*K.5 (3/10/0.53) |
| 40 S ribosomal protein | gi[14277700] | Ribosomal 9 2 | 1:1.66 ± 0.12 | K.LVEAL*C*AHQINLK.V (2/4.17/0.19); R.KVGC*S*C*VWV.D (2/2.72/0.21) |
| 40 S ribosomal protein | gi[4506681] | Ribosomal 13 1 | 1:2.03 | R.DWQVENVTVG*CRPLSK.T (2/4.65/0.53) |
| 40 S ribosomal protein | gi[125969] | Ribosomal 4 1 | 1:2.06 | R.ADHQPLEASYVNLIALC*NTDPLSR.Y (3/15/0.49) |
| 40 S ribosomal protein | gi[4506713] | Ribosomal 7 2 | 1:2.18 ± 0.23 | K.C*ILTC*FNKE*DK. (2/3.35/0.40); R.EC*PSDE*GAQVFMASHFDR.H (3/4.17/0.45) |
| Signaling protein | Cell division protein kinase 6 | gi[266423] | No annotation 1 1 | 1:0.17 | R.ADOQYECYAEIGEGYAG.K.V (3/2.51/0.13) |
| Ectodysplasin A receptor-associated adapter protein | gi[21382527] | Cytoplasmic 1 1 | 1:0.40 | K.NECTCSSC.PR.A (2/2.25/0.15) |
| Transducin β chain 5 | gi[38258891] | No annotation 1 1 | 1:0.45 | R.YPPSGFASAGSSD*A*LDR.A (3/5/0.22) |
| Phosphoinositol-3-kinase, regulatory subunit 4, p150 | gi[23943912] | No annotation 1 1 | 1:0.49 | K.PIPV*ST*PSTY*RR*T.C (3/5.52/0.12) |
| Galectin-1 | gi[216155] | No annotation 23 2 | 1:2.07 ± 0.35 | K.DSNLC*LH*FNPR.F (2/3.34/0.32); R.FNAGHDAIV*VN.KD (2/4.70/0.59) |
| Receptor tyrosine kinase | MuSK | gi[5031927] | No annotation 1 1 | 1:2.18 | R.VEY*C*LVKELFC*AKELVME.TK (3/2.82/0.15) |
| Fibroblast growth factor-9 | gi[544290] | Secreted 1 1 | 1:2.42 | R.QLQYORTG*FHILEP*NIGTQGR.K (3/2.78/0.22) |
| Apoptotic chromatin condensation inducer in the nucleus | gi[7662238] | Nuclear 1 1 | 1:2.47 | K.FLC*ADYAEQD*LHYR.G (3/3.36/0.42) |
| Testin | gi[17380320] | No annotation 2 1 | 1:2.52 | K.NHA*V*QQ*G*H*NADP*VEQ.V (3/3.03/0.37) |
| Tenascin X precursor | gi[9087217] | Secreted 1 1 | 1:3.68 | R.VRGGEE*S*TVGGL*PC.K (3/2.68/0.11) |
| Transcription or replication factor | DNA ligase I | gi[118773] | Nuclear 1 1 | 1:0.38 | K.GLFVA*S*HR*SFARI.S (3/2.65/0.22) |
| Cleavage- and polyadenylation-specific factor 6, 68-kDa subunit | gi[5901928] | Nuclear 1 1 | 1:2.01 | K.RELHM*G*VP*TK*N.KO (3/2.69/0.40) |
| DNA helicase homolog | gi[5523990] | Nuclear 1 1 | 1:4.31 | R.QOLPLQLAYAMSHIKSSQGM*MTLD*C*VEISL-GR.V (3/3.08/0.12) |
| Translation factorb | elf-1 | gi[1174483] | No annotation 8 2 | 1:1.56 ± 0.08 | K.FAC*NGTV*E*HEP*Y*G*V*ILQG*GDOR.K (3/4.16/0.38); K.NICQFL*VEQ*LAK.D (2/2.20/0.15) |
| elf-5 | gi[27735202] | No annotation 1 1 | 1:1.60 | K.FVLCP*EC*PETDL.V*NP.KK (3/2.93/0.27) |
| elf-5A2 protein | gi[9966867] | No annotation 1 1 | 1:1.76 | K.KYEDIC*P*SNH*MDV*PK.N (3/3.42/0.41) |
| EF-Tu, mitochondrial precursor | gi[1706611] | Mitochondrial 4 1 | 1:2.17 | R.HYAT*DC*P*G*HADYVK.N (3/3.96/0.49) |
| Other function protein | WD repeat domain 17 isoform 1 | gi[31317311] | No annotation 1 1 | 1:0.42 | R.IWVTDQD*IC*A*L*H*TAP*GR.V (3/2.52/0.15) |
| Ran-binding protein 2 | gi[1709217] | Nuclear 3 1 | 1:2.03 | K.C*IA*Q*NP*G*Q (2/2.86/0.43) |
| Hyaluronan-binding protein 4 | gi[24307947] | No annotation 1 1 | 1:2.06 | R.YGGNDK*I*V*RTED*MMG*SC.G*G*V*GR.T (3/2.80/0.17) |
| Metallothionein II | gi[127397] | No annotation 1 1 | 1:2.51 | K.CAQG*GIC*G (2/2.20/0.15) |

a Select list of the peptides with the best assignment scores for protein identification and quantification by ICAT analysis. C/C*, ICAT light/heavy reagent-labeled cysteine-including peptide, respectively.

b For heterogeneous nuclear ribonucleoproteins, ribosomal proteins, and translation factors, proteins with at least 1.5-fold alteration are listed. The MS and MS/MS spectra for identification and quantitation of these proteins are shown in Supplemental Figs. 4–24.
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...tivated with nine salt steps, and each step was further separated using a reverse phase column and analyzed with ESI-MS/MS. The relative quantitation of proteins in infected and uninfected cells was determined from the relative abundance ratio of labeled peptides. By this approach, the relative abundance of 322 proteins was measured (Supplemental Table I). Individual peptide sequences and their probability scores for identification and quantitation of those proteins are listed in Supplemental Table II. Among these cellular proteins identified from the infected cells, 167 proteins had at least 1.5-fold quantitative alterations with 119 proteins up-regulated and 48 proteins down-regulated (Table I). Some selected proteins with quantitative alterations are listed in Table III. In addition, two SARS proteins, M and S, were observed to be significantly increased in the infected cells (data not shown).

Comparison and Validation of the Quantitation of Differential Proteins—We used 2D-DIGE and ICAT methods to investigate the differentially expressed proteins in Vero E6 cells infected with SARS-CoV. These two methods are both more sensitive and accurate than traditional methods such as silver staining (data not shown). We further used Western blot assay to confirm the differential expression of the proteomes identified by those two methods. The Western blotting results showed that the ratios of four representative proteins, α-tubulin, HSP90, OP18 stathmin, and vimentin, between the infected and uninfected cells were in agreement with those obtained from 2D-DIGE or ICAT approaches (Fig. 2 and Supplemental Figs. 1–3).

It was observed that the overall expression alterations were similar with both quantitative proteomic approaches, although only 15 differential proteins were identified by both 2D-DIGE and ICAT approaches (Table IV). In addition, the DIGE method detected more isoforms of proteins than the ICAT approach did (Table IV).

Subcellular Location and Function Classification of the Differential Proteins—In the present work, a total of 355 unique gene products of SARS-CoV-infected and uninfected cells were identified and quantitated with either the 2D-DIGE or ICAT method of which 186 proteins had at least 1.5-fold quantitative alterations (Table I and Supplemental Table I). Among those 186 differentially expressed proteins, 60 proteins were down-regulated in infected cells, whereas 126 proteins were up-regulated. These proteins were further classified according to their subcellular locations (Fig. 3). Interestingly, the up-regulated proteins in infected cells were mainly located in the nuclei (about 25%, see Fig. 3A), whereas down-regulated proteins distributed within the cells (Fig. 3B).

Those 186 proteins with at least 1.5-fold differential expression were further classified according to their functions (Supplemental Table I). Fig. 4 presents the functional categories of these proteins. In these identified SARS-CoV-infected cellular proteins, the enzymes, signal proteins, ribosomal proteins, and heterogeneous nuclear ribonucleoproteins (hnRNPs) consisted of about 45% of the total up-regulated proteins (Fig. 4A). On the other hand, more than half of the down-regulated proteins of the infected cells were involved in the enzymatic reactions, signal transduction, immune responses, and actin networks (Fig. 4B).

DISCUSSION

From the literature, very few studies have been performed to analyze the interaction between coronavirus and host cells with proteomic approaches. In the present work, we used two quantitative proteomic assays, DIGE and ICAT, to determine the differentially expressed protein profiles of SARS-CoV-infected and uninfected cells. It was noted that only a few proteins were identified by both DIGE and ICAT approaches, suggesting that these two methods are complementary to each other. In addition, the main difference of those two assays is that ICAT only determines the overall expression level, whereas DIGE can detect and quantitate protein isoforms in a gel. Although the DIGE method can detect protein isoforms possibly caused by post-translational modifications, it has limitations in identification of proteins with very high or low molecular weights, extreme acidic/alkaline proteins, and low abundance proteins. In our present work, 45 proteins that have a molecular mass greater than 100 kDa were identified among which only two proteins were detected by the DIGE approach. Moreover all 22 identified proteins with pI > 10
were contributed by the ICAT method (Supplemental Table I). Concerning the capacity of the protein identification, the DIGE approach identified 48 unique proteins, whereas the ICAT approach obtained 322 proteins; the ICAT approach especially identified more signal proteins, which usually are low abundance proteins in a cell.

Based on the identified proteins in the present work, we can gain an overall insight into the altered protein expression of the host cell responding to SARS-CoV infection. Among the differentially expressed host proteins, many of them participated in viral RNA replication and translation (Tables II and III and Supplemental Table I). It is known that positive-strand RNA viruses recruit normal components of host cellular RNA-processing or translation machineries for the viral RNA synthesis and protein synthesis (19, 20). Our data here indicate that SARS-CoV virus uses a similar strategy when infecting the host cells.
hnRNPs are described as a major group of nuclear RNA-binding proteins that function in transcription, RNA processing, mRNA translation, and turnover (21, 22). In the present work, a total of eight kinds of hnRNP factors were identified to be up-regulated significantly in SARA-CoV-infected cells (Table III and Supplemental Figs. 4–11), suggesting that the virus...
requires the function of hnRNPs. Among these identified hnRNPs, hnRNP A1, hnRNP K, and poly(rC)-binding protein have been reported previously to participate in positive-strand virus genome replication. Moreover, the rest of the hnRNPs were described here for the first time to be involved in coronavirus infection (Table III).

HnRNP A1 has been extensively studied for its role in viral RNA replication. Some reports showed that hnRNP A1 could
bind the RNA of a mouse coronavirus (mouse hepatitis virus (MHV)) at the 3'-end of both plus and minus strands and modulate MHV RNA synthesis (21–23). However, a recent work argued that hnRNP A1 might not be necessary for MHV viral genome replication or transcription in vivo because the absence of hnRNP A1 in infected cells had no effect on the production of infectious MHV (24). In addition, another experiment revealed that a mouse erythroleukemia cell line, CB3, did not express hnRNP A1 but still supported MHV replication, whereas hnRNP A2/B1, hnRNP A/B, and hnRNP A3 could replace hnRNP A1 in cellular functions and viral infection (25). In the present study, hnRNP A1 has been identified and quantitated by ICAT analysis based on doubly charged and both ICAT light reagent- and heavy reagent-labeled peptide K.YHTVNGHNCER.K, a typical trypsin-digested peptide fragmentized at the lysine or arginine carboxyl end, which is marked with the period (Supplemental Fig. 4) and shows up-regulated expression (E6:E6-V, 1:1.90) in SARS-CoV-infected cells (Table III). The present work provides a new explanation for such controversial results on hnRNP A1, i.e. several different hnRNP factors in the infected cells may form a functional hnRNP complex participating in viral RNA metabolism in which one hnRNP factor can be substituted by another without disruption of the function of the hnRNP complex.

The viral genomic RNA of all positive-strand RNA viruses need to be translated by recruited host factors (26, 27). A recent report showed that the decrease of 60 S ribosome protein levels reduced a positive-strand virus (Brome mosaic virus)-directed expression in yeast cells (27). The quantitatively proteomic approaches used here revealed that about nine kinds of ribosomal proteins, including components of both 40 and 60 S ribosomal subunits, were up-regulated significantly (Table III and Supplemental Figs. 12–20), suggesting that overall up-regulation of the ribosomal protein expression is required for positive-strand RNA virus propagation in host cells.

Translation factors have been well documented to participate in the processes of the virus RNA and protein synthesis (26, 28). For example, the elongation factors EF-Tu and EF-Ts were found to bind tightly to the viral RNA-dependent RNA polymerase (29). We also identified the up-regulation of EF-Tu expression in SARS-CoV-infected cells (Table III and Supplemental Fig. 21). Some studies showed that eukaryotic translation initiation factor eIF-4 protein complex is involved in viral protein synthesis (26). On the other hand, the present studies revealed that the expression of eIF-1, eIF-5, and eIF-5A2 was increased in the infected cells, suggesting that these factors play a role in the process of SARS-CoV viral translation (Table III and Supplemental Figs. 22–24).

In summary, we present the first quantitative proteomic work on the cellular responses to SARS-CoV infection, establishing so far the most comprehensive differential proteomic index for SARS-CoV-infected cells. The identified differential profile derived from the infected cellular proteins gives the implications for the infectivity and pathogenesis of SARS-CoV and provides a valuable resource for diagnosis, drug development, and clinical treatment for SARS.

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