

Cellular Chaperonin CCT γ Contributes to Rabies Virus Replication during Infection

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Rabies, as the oldest known infectious disease, remains a serious threat to public health worldwide. The eukaryotic cytosolic chaperonin TRiC/CCT complex facilitates the folding of proteins through ATP hydrolysis. Here, we investigated the expression, cellular localization, and function of neuronal CCT γ during neurotropic rabies virus (RABV) infection using mouse N2a cells as a model. Following RABV infection, 24 altered proteins were identified by using two-dimensional electrophoresis and mass spectrometry, including 20 upregulated proteins and 4 downregulated proteins. In mouse N2a cells infected with RABV or cotransfected with RABV genes encoding nucleoprotein (N) and phosphoprotein (P), confocal microscopy demonstrated that upregulated cellular CCT γ was colocalized with viral proteins N and P, which formed a hollow cricoid inclusion within the region around the nucleus. These inclusions, which correspond to Negri bodies (NBs), did not form in mouse N2a cells only expressing the viral protein N or P. Knockdown of CCT γ by lentivirus-mediated RNA interference led to significant inhibition of RABV replication. These results demonstrate that the complex consisting of viral proteins N and P recruits CCT γ to NBs and identify the chaperonin CCT γ as a host factor that facilitates intracellular RABV replication. This work illustrates how viruses can utilize cellular chaperonins and compartmentalization for their own benefit.

Rabies virus (RABV), the prototype of the *Lyssavirus* genus, is a neurotropic virus that almost exclusively invades neurons (1) and causes rabies disease that is fatal in almost 100% of cases. As the oldest known infectious disease, rabies remains incurable but preventable by pre- and postinfection vaccine therapy (2). Currently, approximately 15,000,000 individuals receive postexposure prophylaxis annually for rabies, which is responsible for over 55,000 human deaths globally each year (3). Therefore, rabies remains a serious threat to public health worldwide, especially in developing countries.

The life cycle of RABV progresses in the cytoplasm with transcription of the five viral genes encoding nucleoprotein (N), matrix protein (M), phosphoprotein (P), glycoprotein (G), and large protein (L). Aggregates called Negri bodies (NBs) form during RABV replication, as first described in 1903 by Adelchi Negri, an Italian pathologist and microbiologist (4). NBs resemble the intranuclear Cowdry bodies in herpesvirus-infected cells (5) and Guarnieri bodies in the cytoplasm of poxvirus-infected cells (6). Several research groups reported that NBs are characterized by the accumulation of viral nucleocapsid proteins (7–9) and also contain endothelial nitric oxide synthase (eNOS) (10) and Hsp70 (11). Recently published data demonstrated that NBs are sites of viral transcription and replication (12).

Researchers are paying increasingly close attention to the role of host factors in virus replication by using comparative proteomic approaches. Host factors related to infection have been identified for approximately 20 viruses. Several critical host proteins, such as BAG3 (13), cyclophilin A (14), Hsp90 (15), RACK1 (16), and desmin (17), have been shown to be involved in virus replication and carcinogenesis. In the past decade, multiple studies have been carried out in different laboratories to investigate the host response to RABV infection at the transcriptional (18–20) and translational levels (21–24). The cellular chaperone protein Hsp70 was reported to be incorporated into rabies virions (25), and TLR3 is a major host molecule involved in the spatial arrange-

ment of RABV-induced NBs and viral replication (26). However, less is known about whether other host factors are involved in RABV replication.

The eukaryotic cytosolic chaperonin TRiC/CCT is a large complex comprised of two stacked rings of eight subunits (CCT1 to -8) each that facilitates the folding of proteins through ATP hydrolysis (27). The folding of several mammalian proteins mediated by TRiC/CCT, such as tubulin, actin, WD-repeat proteins, cyclin E, Cdc20, and von Hippel-Lindau tumor suppressor protein, has been identified. Chaperonin CCT also plays critical roles in the nervous system and is ultimately required for the morphogenesis and survival of sensory neurons of the retina (28). In zebrafish, chaperonin CCT γ specifically controls retinotectal development (29). With regard to functions in the virus life cycle, t-complex polypeptide 1 (TCP-1) may be involved in hepatitis B virus capsid assembly (30), while CCT5 participates in hepatitis C virus (HCV) RNA replication and virion production by its interaction with NS5B (31). CCT β is known to associate with influenza virus RNA polymerase subunit PB2 (32), and the chaperonins TCP-1 γ and ϵ interact with the Gag polyprotein of retrovirus type D and nuclear protein EBNA-3 of Epstein-Barr virus, respectively (33, 34). However, the exact mechanism of CCT function in the RABV life cycle has yet to be determined, and the intrinsic *in vivo* events leading to the recruitment of CCT to the aggregation site remain poorly understood.

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In the present study, we analyzed the differentially expressed host protein profiles using mouse N2a cells as a RABV infection model and further investigated the association between NBs involved in viral transcription and replication and the chaperone protein CCT γ , as well as the functions of this host factor.

MATERIALS AND METHODS

Cells and viruses. Mouse neuroblastoma N2a cells (a gift of professor Xiaofeng Guo, South China agricultural University, Guangzhou, People's Republic of China) and 293T cells (ATCC, Rockville, MD, USA) were cultured at 37°C with 5% CO₂ in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% heat-inactivated fetal bovine serum (Gibco/Invitrogen, Carlsbad, CA, USA). All media were supplemented with penicillin (100 U/ml) and streptomycin (100 μ g/ml). The RABV HEP-Flury strain was stored in our laboratory and propagated in N2a cell monolayers for two-dimensional electrophoresis (2-DE) sample preparation.

Antibodies and reagents. Rabbit polyclonal antibodies (PAb) against TCP-1 γ (H-300, CCT γ), Hsp90, and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) were purchased from Santa Cruz Biotechnology (Santa Cruz, CA, USA), Abcam (Cambridge, MA, USA), and Hangzhou GoodHere Biotechnology Co. Ltd. (Hangzhou, China), respectively. Mouse monoclonal antibody (MAb) anti-Flag M2 antibody was purchased from Sigma-Aldrich (Mission collection, St. Louis, MO, USA). Mouse MAbs to N and P proteins of RABV, as well as rabbit PAb to N protein of RABV, were prepared in our laboratory. Urea, thiourea, sodium dodecyl sulfate (SDS), glycine, Tris, Biolyte (pH 3 to 10), iodoacetamide, dithiothreitol (DTT), low-melting-temperature agarose, CHAPS {3-[(3-cholamidopropyl)-dimethylammonio]-1-propanesulfonate}, and 24-cm ReadyStrip immobilized pH gradient (IPG) strips (linear, pI 5 to 8) were all from Bio-Rad Laboratories (Hercules, CA, USA).

In vitro cell infection and sample preparation. Sample preparation was carried out according to a previously described method (35, 36). Briefly, N2a cells were grown in T-flasks (75 cm², 15-ml working volume) to 80 to 90% confluence before being infected with the RABV Flury strain at a multiplicity of infection (MOI) of 1. The cells were harvested at 12, 48, and 96 h postinfection (h.p.i.) with a cell scraper and centrifuged at 20,000 \times g for 5 min. The pellet was washed 3 times with prechilled 0.25 M sucrose Tris buffer (pH 7.0).

The cell pellet was disrupted in lysis buffer, which contained 7 M urea, 2 M thiourea, 4% (wt/vol) CHAPS, and 65 mM DTT and was supplemented with 0.2% Biolyte 3/10, 1 mM phenylmethylsulfonyl fluoride (PMSF), 20 units/ml DNase I, and 0.25 mg/ml RNase A before use. The mixture was vortexed for 1 h and sonicated on ice for 20 min, followed by centrifugation at 20,000 \times g for 1 h at 4°C. The supernatant was kept at -80°C until used. The protein concentration of all samples diluted 1:10 was determined by the Bradford method (37) with 0.5 mg/ml bovine serum albumin (BSA) as the standard for quantitation.

2-DE and image analysis. 2-DE procedures were carried out as described previously (35) with minor modifications. The first dimension of 2-DE was performed in the Protean isoelectric focusing (IEF) cell (Bio-Rad) with commercial 24-cm-long ReadyStrip IPG strips (pH 5 to 8, linear; Bio-Rad) according to the manufacturer's instructions. A total of 250 μ g of protein was loaded onto each IPG strip. The following parameter settings and conditions for active rehydration of the IPG strips and IEF were modified from a previously described method (35, 38): constant temperature at 17°C and (i) 50 V for 14 h; (ii) 250 V, linear, for 1 h; (iii) 1,000 V, rapid, for 1 h; (iv) 10,000 V, linear, for 5 h; and (v) 10,000 V, rapid, for 90,000 V-h. After the IEF, the isoelectric-focused strips were incubated for 15 min in an equilibration buffer (6 M urea, 30% glycerol, 2% SDS, and 0.375 M Tris, pH 8.8) containing 1% DTT and then incubated for 15 min in an equilibration buffer containing 2.5% iodoacetamide. The equilibrated IPG strips were sealed with 1% low melting-point agarose in SDS running buffer (supplemented with 0.001% bromophenol blue marker) on top of the gels and further resolved by 11% SDS-poly-

acrylamide gel electrophoresis (PAGE) at 80 V for 45 min and then 200 V until the dye front reached the end of the gels. The gels were stained by the modified silver-staining method compatible with mass spectrometry (MS) (39) and scanned at a resolution of 500 dots per inch using the ImageScanner III (GE Healthcare, USA). Spot detection, spot matching, and quantitative intensity analysis were carried out with PDQuest 2-D analysis software 8.0.1 according to the manufacturer's protocol (Bio-Rad). The gel images were normalized according to the total quantity in the analysis set. Comparison of the relative intensities of the spots between RABV-infected and mock-infected groups at three time points (four replicates for each group) was analyzed by Student's *t* test. Only the spots with intensity ratios (infected versus uninfected group) larger than 2.0 ($P \leq 0.05$) or smaller than 0.5 ($P \leq 0.05$) were considered to have significantly changed due to infection and subjected to identification by MS.

In-gel tryptic digestion. Differentially expressed protein spots were excised from the silver-stained gels, transferred to a V-bottom 96-well microplate, and washed with 100 μ l 50% (vol/vol) acetonitrile (ACN) in 25 mM ammonium bicarbonate for 1 h. The following procedure was performed as previously described (35, 36). Briefly, after being dehydrated with 100 μ l of 100% ACN for 20 min, gel plugs were thoroughly dried in a SpeedVac concentrator (Thermo Savant, Holbrook, NY, USA) for 30 min. The dried gel particles were treated with 2 μ l/well of trypsin (Promega, Madison, WI) in 25 mM ammonium bicarbonate at 4°C for 45 min for rehydration and then incubated at 37°C for 12 h. Following the trypsin digestion, the peptide mixtures were extracted by 8 μ l/well of extraction solution (0.5% TFA, 50% ACN) at 37°C for 1 h. Finally, the extracts were dried under the protection of N₂.

MS analysis and database searches. Matrix-assisted laser desorption ionization-time of flight (MALDI-TOF) MS and tandem TOF (TOF/TOF) MS were performed at the Proteomic Core Facility, Fudan University, China, as described previously (35, 36). Mixtures of the peptide were redissolved in 0.8 μ l of matrix solution (α -cyano-4-hydroxycinnamic acid [Sigma, St. Louis, MO, USA] in 0.1% TFA, 50% ACN) and then spotted on the MALDI plate. After air drying, samples were analyzed on a 4700 Proteomics Analyzer (Applied Biosystems, Foster City, CA, USA). The trypsin-digested peptides of myoglobin with internal calibration mode to calibrate the mass instrument were added to the six calibration spots on the MALDI plate. An accelerating voltage of 20 kV was applied to the sample plate. The UV laser was operated at a 200-Hz repetition rate with a wavelength of 355 nm. Data interpretation of acquired MS and tandem MS (MS/MS) peptide spectra were carried out with Data Explorer software version 4.5 (Applied Biosystems) in default mode. Parent mass peaks with a mass range of 700 to 3,200 Da and minimum signal-to-noise ratio of 20 were selected for TOF/TOF analysis. The MS and MS/MS spectra generated were subsequently submitted to MASCOT (version 2.1, Matrix Science, London, United Kingdom) by GPS Explorer software (version 3.6, Applied Biosystems). The parameters for searches were as follows: National Center for Biotechnology Information nonredundant (NCBIInr) database (release date 2009), taxonomy of mouse and viruses, carbamidomethylation selected as a fixed modification, peptide mass tolerance of 100 ppm; MS/MS ion mass tolerance of 0.6 Da, maximum of one missed cleavage; and tryptic digestion. Modifications included methionine oxidation. Known contaminant ions (human keratin and tryptic autodigest peptides) were excluded. In total, 11,505,486 sequences and 3,925,745,078 residues in the database were actually searched. MASCOT protein scores (based on combined MS and MS/MS spectra) of greater than 64 were considered statistically significant ($P \leq 0.05$). The individual MS/MS spectrum with a statistically significant (confidence interval of $\geq 95\%$) ion score (based on MS/MS spectra) was accepted. Only proteins with a significant ion score or an ion score above 21 were reported. To eliminate the redundancy of proteins that appeared in the database under different names and accession numbers, the single protein member belonging to the *Mus musculus* species or with the highest protein score (top rank) was separated from the multiprotein family.

TABLE 1 Primers designed for quantitative real-time PCR

Gene	Forward primer (5'→3')	Reverse primer (5'→3')	Length (bp)	GenBank accessionno.
HSP90	ATAGGCTTGTGTCTTCACCCTGCT	AGCCCATGTAGAGTTGTCTCGCA	111	S46109
CCT γ	GCAAGTGTGCCGAATGTTCTACT	TGTTCCACACCAGTCATGGCCTTA	117	NM_009836
CCT θ	GCAGGCACTGGTGCAAATGTCATA	TGTCCCATTTCTTCTTGGACGGGA	188	AK172867
Prph1	AAAGACGACTGTGCCTGAGATGGA	AAGACTTGTCCAGGTCAGTGTGCT	143	BC046291
ACTB	TGTGATGGTGGGAATGGGTCAGAA	TGTGGTGCCAGATCTTCTCCATGT	140	X03765
Srm	ACCAGCTCATGAAGACAGCACTCA	TGCTACACAGCATGAAGCCGATCT	189	L19311
EIF2S2	TTGCACCCACCAGGATAAGGACAT	ACAGCAGGGCTAACTGACTGTTCT	101	NM_026030
DLST	TGGTGTGAGCATGTTCTGGGAAGA	ATGACACCATCATCTCGTCGCTGT	166	NM_030225.4
OAS1B	ATGCCTTCATCCTCCCTTCCAGTT	TTAGCCATGGTTCGGATCACACA	100	AM887912
PSMA6	ACGGAAGCATTGGCTGTGTGATG	TCATACAACAACCGAGTGGCCTCA	199	NM_011968
SEPT2b	TGTGCTTTCATTGCTGCAGGAGG	TGCAATGTGAAGAGGCAAAGCGAG	168	NM_001159718
HNRPD	TTGGAGGCTTGTGCTGGGATACAA	ATCTTCCCGTGACTGGGTCTGTTT	108	BC021374
SET	ACGAAGCGAGGATGATTAGCACA	ACAGGGAGGAAAGGACTGCAACTT	119	BX005298.9
GAPDH	TCAACAGCAACTCCCCTCTTCCA	ACCCTGTTGCTGTAGCCGATTCA	115	NM_008084

Quantitative real-time PCR. To validate the results of the differential proteome analysis at the mRNA level, specific primers (Table 1) were designed to simultaneously amplify various target genes corresponding to the MS/MS-identified proteins and RABV-encoded proteins. Quantitative real-time PCR was performed using cDNA from N2a cells (12, 48, and 96 h.p.i.) as the template with a RevertAid first strand cDNA synthesis kit (Fermentas). cDNA synthesis was performed with 1 μ g RNA using oligo(dT) primers (100 ng). The quantitative real-time PCR was performed on the 7500 real-time PCR sys-

tem (Applied Biosystems) in a total volume of 20 μ l containing 200 ng of cDNA template, 200 nM each primer, and 1 \times SYBR premix *Ex Taq* (perfect real time, TaKaRa, Dalian, China). The target genes were amplified in triplicate with the following conditions: an initial denaturation at 95°C for 10 min and 40 cycles of 95°C for 10 s, 60°C for 30 s, and 72°C for 10 s. Melting curves were obtained, and quantitative analysis of the data was performed using the 7500 system SDS software version 1.3.1 with a relative quantification (cycle threshold [$\Delta\Delta C_T$]) method (Applied Biosystems). The uniqueness and sizes of PCR prod-

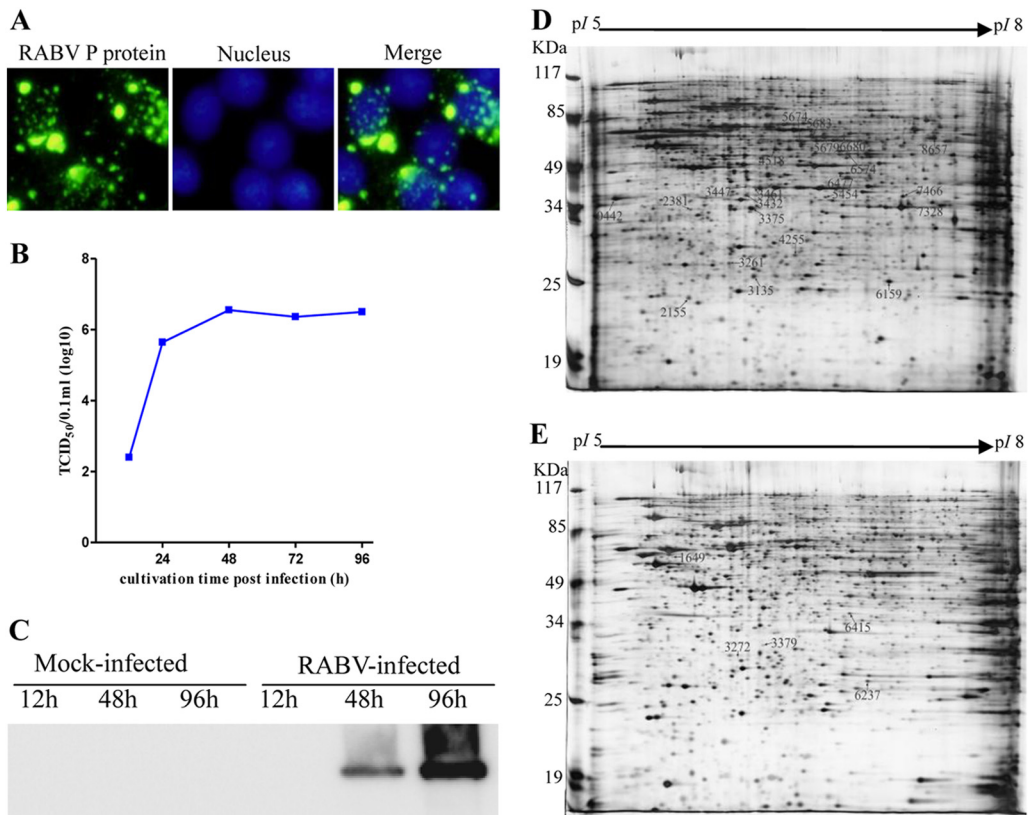


FIG 1 Identification of RABV-infected mouse N2a cells and representative 2-DE proteomic profiles from mock- and RABV-infected N2a cells. (A) Identification of RABV-infected mouse N2a cells by IFA using the MAb to RABV P. (B) RABV infection kinetics in N2a cells. (C) Detection of the viral N protein in RABV-infected and mock-infected N2a cells by Western blotting using a specific MAb. (D) 2-DE gel of RABV-infected N2a cells with upregulated protein spots labeled. (E) 2-DE gel of mock-infected N2a cells with downregulated protein spots marked.

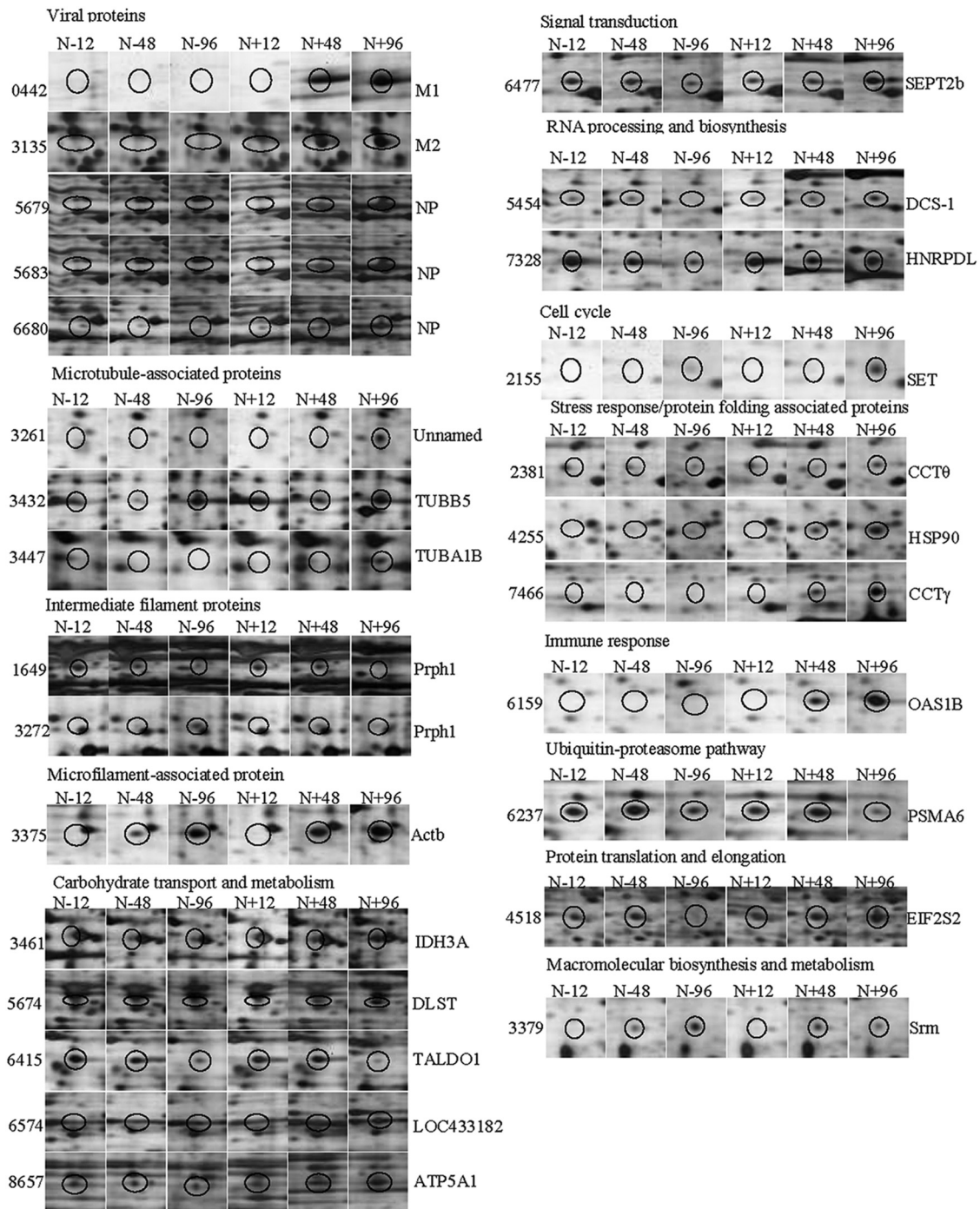


FIG 2 Dynamic 2-DE profiles of the differentially expressed proteins in RABV-infected N2a cells. Differentially expressed protein spots are circled. N+ and N- indicate the RABV-infected and mock-infected mouse N2a cells, respectively, and numbers indicate time (h) posttransfection.

ucts were confirmed by generating melting curves. N2a cells mock infected in parallel were used as controls.

Immunoblotting. Samples of RABV-infected and mock-infected N2a cells were lysed at 12, 48, and 96 h.p.i., and the protein concentrations were determined by the Bradford assay as described above. Equivalent amounts of cell lysates (80 μ g) were subjected to 12% SDS-PAGE and then transferred to nitrocellulose membranes (Hybond-C extra; Amersham Biosciences, USA). After blocking with 5% skimmed milk in phosphate-buffered saline (PBS) with 0.05% Tween 20 for 1 h, the membranes

were incubated with mouse MAb to RABV N or rabbit PAb against CCT γ at 37°C for 2 h, followed by horseradish peroxidase (HRP)-conjugated goat anti-mouse IgG or goat anti-rabbit IgG (Kirkegaard & Perry Laboratories, Inc. [KPL], Gaithersburg, MD, USA) at 37°C for 1 h. Finally, the blots were developed with SuperSignal West Femto maximum sensitivity substrate (Thermo Fisher Scientific, Rockford, IL, USA).

Double-staining immunofluorescence and nuclear staining. After RABV- and mock-infected N2a cells were cultured for 48 h, they were washed once with PBS, fixed with cold acetone-methanol (1/1) for 20 min

TABLE 2 Differentially expressed protein spots which were successfully identified by MALDI-TOF and MALDI-TOF/TOF

Type of protein and spot no. ^a	Protein name	Abbreviation	Accession no. ^b	M_r (pred/exp ^c)	pI (pred/exp)	Matched/unmatched ^d	Abundance ratio in infected/uninfected cells (mean \pm SD) at indicated time (h) ^e			Coverage (%) ^f	Protein/best ion score ^g	Peptides identified ^h
							12	48	96			
RABV viral proteins												
0442	Phosphoprotein	M1	gi 70724981	33.43/40.3	4.8/5.0	14/61	NI	NI	42	311/105	IFVNSAIR AGLADLEMAEETVD LINR NIEDNOAHLQGEPI EVDNLPEDMR	
3135	M2 protein	M2	gi 9627200	23.14/25.1	6.34/6.1	6/48	NI	NI	25	114/55	TLIFQWADSR	
5679	Nucleoprotein	NP	gi 18478987	50.71/55.5	6.37/6.7	20/43	NI	NI	52	204/47	EALLYFFHK	
5683	Nucleoprotein	NP	gi 39930275	49.15/55.6	6.04/6.6	20/84	NI	NI	49	256/52	FLAGTYDMFFSR	
6680	Nucleoprotein	NP	gi 38017865	50.73/55.3	6.08/6.8	13/75	NI	NI	28	91/20	FLAGTYDMFFSR	
Microtubule-associated proteins												
3261	Unnamed protein product	Unnamed	gi 26355849	32.24/26.9	5.56/5.9	7/32	2.59 \pm 0.35***	NI	17	67/23	FPGQINADLR	
3432	Tubulin, beta 5	TUBB5	gi 7106439	49.64/39.6	4.78/6.1	12/59	0.63 \pm 0.19	NI	20	131/44	FPGQINADLR	
3447	Alpha-tubulin isotype M-alpha-2	TUBA1B	gi 202210	50.13/41.8	4.94/6.1	8/23	NI	NI	24	162/50	AVFVDLEPTVIDEVR	
Intermediate filament proteins												
1649	Peripherin	Prph1	gi 2253159	52.65/56.5	5.36/5.5	28/27	1.07 \pm 0.37	2.10 \pm 0.16**	57	228/38	FLEQQNAALR	
3272	Peripherin	Prph1	gi 2253159	52.65/31.4	5.36/6.0	18/30	2.06 \pm 0.81	0.46 \pm 0.09***	38	136/36	LLGSGPSSAR	
Microfilament-associated proteins												
3375	Putative beta-actin (amino acids 27–375)	Actb	gi 49868	39.16/37.1	5.78/6.1	9/37	2.24 \pm 0.51**	1.29 \pm 0.05**	25	203/70	GYSFTTAAER QEYDESGPSIVHR	
Energy and metabolism												
3461	Isocitrate dehydrogenase 3 (NAD ⁺) alpha isoform	IDH3A	gi 148693874	39.30/41.3	5.88/6.1	6/27	1.51 \pm 0.68*	2.82 \pm 0.91**	18	73/31	IAEFAFEYAR	
5674	Dihydropyrimidinase 2 component of 2-oxoglutarate complex	DLST	gi 21313536	48.96/52.1	9.11/6.5	14/117	0.91 \pm 0.34	2.71 \pm 0.57**	31	176/42	VEGGTFLTLR NVETMNYADIER	
6415	Transaldolase 1, isoform CRA_e	TALDO1	gi 148686116	31.51/39.1	7.66/6.8	12/100	0.83 \pm 0.42	0.74 \pm 0.23	38	108/23	FAADAIKLER	
6574	Hypothetical protein LOC433182	LOC433182	gi 70794816	47.11/49.7	6.37/6.9	9/113	NI	NI	21	117/49	AAVPSGASTGIYEALELR	
8657	ATP synthase, H ⁺ transporting, mitochondrial F1 complex, alpha subunit, isoform 1, isoform CRA_h	ATP5A1	gi 148677504	54.74/55.8	9.36/7.5	13/29	1.13 \pm 0.16	2.31 \pm 0.64**	25	98/24	TGAIVDVPVGEELLGR	
Signal Transduction												
6477	Septin 2 b	SEPT2b	gi 228480253	36.95/44.3	5.74/6.8	11/111	0.61 \pm 0.22*	1.21 \pm 0.12	31	175/51	STLINSFLTDLYPER	

RNA processing and biosynthesis																					
5454	Chain A, crystal Structure Of mRNA decapping enzyme (Dcps) from <i>Mus musculus</i> at 1.83-Å resolution	DCS-1	gi 52696229	40.42/38.6	6.25/6.7	11/57	0.71 ± 0.33	1.20 ± 0.12	2.29 ± 0.33**	32	103/33	DLTPEHLPLLR									
7328	Heterogeneous nuclear ribonucleoprotein D-like	HNRPDL	gi 81917948	33.54/38.1	6.85/7.3	12/76	1.02 ± 0.23	1.63 ± 0.21*	2.94 ± 0.51***	30	120/32	GFGEVLFKDLTEYLSR									
Cell cycle																					
2155	SET translocation	SET	gi 123295280	23.11/24.91	5.6/5.43	12/39	0.23 ± 0.08*	2.09 ± 0.36**	2.09 ± 0.36**	57	137/24	EQQEAIHIDEVQNEIDR									
Stress response/protein folding associated proteins																					
2381	mKIAA0002 protein	Cctθ	gi 50510319	60.19/37.7	5.44/5.6	6/20	NI	NI	2.17 ± 0.69*	11	64/32	FAEAFEAIIPR									
4255	HSP90	HSP90	gi 13752262	52.56/28.0	8.81/6.4	16/54	NI	NI	NI	36	88/0										
7466	Chaperonin containing Tcp1, subunit 3 (gamma)	CCTY	gi 6753320	60.59/39	6.28/7.3	15/95	NI	NI	NI	31	65/21	TAVETAVLLLR									
Immune response																					
6159	Putative oligoadenylate synthetase 1b	OAS1B	gi 159031195	43.86/24.1	9.46/7.2	11/22	NI	NI	NI	24	64/0										
Ubiquitin-proteasome pathway																					
6237	Proteasome alpha 6 subunit	PSMA6	gi 6755198	27.35/26.7	6.34/7.0	9/63	0.80 ± 0.16	0.91 ± 0.40	0.50 ± 0.14**	39	256/72	HITTESPEGRAINOGGLTSVAVR ILTEAEDIAHLVALAERD									
Protein translation and elongation																					
4518	Eukaryotic translation initiation factor 2, subunit 2 (beta)	EIF2S2	gi 14149756	38.07/51.4	5.61/6.3	10/25	0.95 ± 0.30	1.13 ± 0.02	18.98 ± 4.63***	22	76/28	DYTYEELLNR									
Macromolecular biosynthesis and metabolism																					
3379	Spermidine synthase	Srm	gi 6678131	33.97/33.7	5.3/6.2	7/40	1.11 ± 0.26	0.40 ± 0.05***	0.40 ± 0.05***	18	122/31	YQDILVFRVLIIGGGGGVLR									

^a Spot no. is the unique sample spot protein number that refers to the labels in [Figure 2](#).

^b Accession no. is the MASCOT result of MALDI-TOF/TOF searched from the NCBItr database.

^c pred/exp, predicted/experimental.

^d Number of peaks that match/do not match the trypsin peptides.

^e Protein spots with an abundance ratio for infected/uninfected cells of less than 0.5 were downregulated, and spots with a ratio greater than 2 were upregulated. Mean, the average abundance ratio for paired protein samples; SD, standard deviation of abundance ratios of the one certain protein spot; NI, newly induced proteins in RABV-infected N2a cells; ND, protein spot was not detectable in RABV-infected N2a cells. *P* values from paired *t* test are shown as follows: * *P* ≤ 0.05; ** *P* ≤ 0.01; *** *P* ≤ 0.001.

^f Sequence coverage (%) is the number of amino acids spanned by the assigned peptides divided by the sequence length.

^g Protein score (based on combined MS and MS/MS spectra) and best ion score (based on MS/MS spectra) were from MALDI-TOF/TOF identification. The proteins that had a statistically significant protein score of great than 64 (*P* ≤ 0.05) were considered successfully identified.

^h Peptides identified by MALDI-TOF/TOF with a statistically significant ion score (confidence interval, ≥95%), or ion score above 21.

at -20°C , and then allowed to be air dried. The fixed cells were incubated with a mixture of MAbs against the proteins N and P of RABV, rabbit anti-N PAb and PAb to CCT γ (Santa Cruz), or PAb to Hsp90 (Abcam). Nuclei were stained with 4',6-diamidino-2-phenylindole (DAPI) for indirect immunofluorescence assay (IFA). Viral proteins were detected by incubation with tetramethyl rhodamine isocyanate (TRITC)-conjugated goat anti-mouse secondary antibody (KPL), and the host proteins were detected by incubation with fluorescein isothiocyanate (FITC)-conjugated goat anti-rabbit secondary antibody (KPL). The triple-stained cells were washed 5 times with PBS-Tween 20 and subsequently examined under a laser confocal microscope. Confocal images were acquired using an inverted Leica TCS SP5 microscope with proper excitation laser wavelengths.

Construction of plasmids and DNA transfection. The full-length N and P genes of RABV and the CCT γ gene were amplified via PCR from the cDNA of RABV-infected N2a cells and then cloned into pCI-neo (Promega, WI, USA), pCMV-N-Flag (Clontech, CA, USA), and pCMV-N-HA (Clontech, CA, USA) by using the XhoI/XbaI, EcoRI/SalI, and EcoRI/XhoI cloning sites, respectively. The specific primer sequences for the P gene are as follows: for the pCI-neo construct, forward primer 5'-GCCG AATTCATGAGCAAGATCTTTGTC-3' and reverse primer 5'-GACGTC GACTTAGCAAGATGTATAGCG-3' (EcoRI and SalI restriction sites are underlined), and for the pCMV-N-Flag construct, forward primer 5'-GG AATTCGGATGAGCAAGATCTTTGTTAATCCGAGTG-3' and reverse primer 5'-CCCTCGAGTTAGCATGATTTGTAGCGATCCAAGTCA T-3' (EcoRI and XhoI restriction sites are underlined). The specific primer sequences for the N gene are as follows: for the pCI-neo construct, forward primer 5'-GAACTCGAGATGGATGCCGACAAGATT-3' and reverse primer 5'-GCTCTAGATTATGAGTCACTCGAATACGTT-3' (XhoI and XbaI restriction sites are underlined), and for the pCMV-N-Flag construct, forward primer 5'-GGAATTCGGATGGATGCCGACAA GATTGTGTT-3' and reverse primer 5'-CCCTCGAGTTATGAGTCACT CGAATACGTTTGT-3' (EcoRI and XhoI restriction sites are underlined). The specific primer sequences for the CCT γ gene are as follows: for the pCI-neo construct, forward primer 5'-GGAATTCATGATG GGCCACCGTC-3' and reverse primer 5'-GCTCTAGACTCACTCTGG CCAGCAT-3' (EcoRI and XbaI restriction sites are underlined), and for the pCMV-N-HA construct, forward primer 5'-GGAATTCGGATGATG GGCCACCGTCCAGTGTCT-3' and reverse primer 5'-CCCTCGAGTCA CTCTGGCCAGCATCTGGAG-3' (EcoRI and XhoI restriction sites are underlined). All constructs were verified by DNA sequencing.

One day prior to DNA transfections, N2a cells and 293T cells were seeded into 96-well plates (Corning, NY, USA) or 35-mm glass bottom dishes (Shengyou Biotechnology Co. Ltd., Hangzhou, China), and the resultant vectors were cotransfected into the N2a cells and 293T cells using Lipofectamine 2000 reagent (Invitrogen) for transient protein expression. Two days later, cells were washed twice with PBS and fixed with ice-cold acetone-methanol (1/1) for subsequent use in IFA. N2a cells and 293T cells transfected with empty vectors pCI-neo, pCMV-N-HA, and pCMV-N-Flag were used as negative controls.

Coimmunoprecipitation assay. The coimmunoprecipitation assay was conducted with a minor modification as stated previously (36). Briefly, at 48 h after infection or cotransfection, N2a cells inoculated with the Flury strain of RABV and 293T cells cotransfected with the vectors pCMV-HA-CCT γ and pCMV-Flag-P or pCMV-Flag-N were lysed with NP-40 lysis buffer (Beyotime, Jiangsu, China). The lysate supernatant was incubated with the MAb against RABV viral protein N or P, anti-Flag, and anti-hemagglutinin MAbs overnight at 4°C , respectively. Immune complexes were precipitated by incubation with protein A/G Plus-agarose (Santa Cruz) for 4 h at 4°C . After being washed five times, the immunoprecipitated proteins were analyzed by Western blotting using anti-CCT γ MAb, anti-Flag MAbs, or MAb to RABV viral protein N or P, respectively.

Knockdown of cellular CCT γ by lentivirus transduction and establishment of a stable cell line expressing CCT γ shRNA. pLKO.1-based lentiviral particles for knockdown of CCT γ (catalog number

TRCN0000120448, targeting sequence CCAAGTACCATTTCGTCTGCTTA) and Mission nontarget small hairpin RNA (shRNA) control transduction particles (catalog number SHC002V, targeting sequence CAACA AGATGAAGAGCACCAA) were purchased from Sigma-Aldrich (Mission Collection, St. Louis, MO, USA). The N2a cells were plated in a 24-well plate with DMEM overnight, and lentivirus transduction was carried out according to the manufacturer's instructions. Hexadimethrine bromide (Sigma-Aldrich) was added to the medium (final concentration, 8 $\mu\text{g}/\text{ml}$) to enhance transduction efficiency. Lentiviral particles (MOI = 0.5) were added, and the plates were swirled gently for mixing. Puromycin dihydrochloride (5 $\mu\text{g}/\text{ml}$) was added 48 h later for 10 to 14 days of selection. After the transduction of lentiviral vectors to target cells, the validation of shRNAs was performed by Western blotting and real-time comparative quantitative PCR. The viability of N2a cells expressing CCT γ shRNA was determined using the CCK-8 assay according to the manufacturer's protocol (Sigma-Aldrich). To further analyze the function of CCT γ , a stable N2a cell line expressing CCT γ shRNA was infected with the Flury strain of RABV at an MOI of 0.1, and the supernatants were collected at 24, 36, and 48 h.p.i. The titers of viruses in the culture supernatants were determined in BHK-21 cells, and 50% tissue culture infective dose (TCID₅₀) values were calculated by the Reed-Muench method (40). The transcript levels of the RABV genomic RNA in the infected cells were quantified by comparative and absolute real-time PCR using the specific primers 5'-AAGGAGTTGAATGACAGGGTGCCA-3' and 5'-ACTTGG GATGGTTCGAAAGGAGGA-3'. Viral protein N was analyzed in RABV-infected cells by Western blotting assay.

To investigate which step of the RABV life cycle is affected, the stable N2a cells were treated with or without cycloheximide (CHX; Beyotime, Shanghai, China) at a dose of 100 $\mu\text{g}/\text{ml}$ as described previously (41) and simultaneously infected with RABV at an MOI of 5. At 8 and 12 h.p.i., cellular total RNA was isolated from the resultant cells using TRIzol reagent following the manufacturer's protocol. Finally the viral mRNA of the RABV N gene was quantified with a relative quantification ($\Delta\Delta C_T$) method using the specific primers 5'-AGCAGCAATGCAGTTCCTTGA GGG-3' and 5'-TTGTCAGTTCATGCCTCCTGTCA-3'.

Statistical analysis. Statistically significant differences between groups were determined using the Student *t* test. A *P* value of less than 0.05 was considered significant.

RESULTS

2-DE protein profiles of RABV infection in N2a cells. To provide a detailed profile of changes in protein expression, RABV-inoculated N2a cell monolayers were evaluated at 12, 48, and 96 h.p.i. by indirect immunofluorescence assay (IFA) and immunoblotting. Successful virus replication was confirmed by IFA analysis of RABV-inoculated N2a cells at 48 h.p.i. (Fig. 1A). Based on the growth curve, RABV replication reached a maximum at about 48 h.p.i. with a titer of $10^{6.55}$ TCID₅₀ (Fig. 1B). In Western blot analysis, the N protein of RABV was only detected at 48 and 96 h.p.i. from RABV-infected N2a cells (Fig. 1C). These data showed that infectious RABV progeny were generated in the N2a cell monolayers.

Subsequently, the cellular proteins of RABV-infected N2a cells extracted at 12, 48, and 96 h.p.i. were subjected to 2-DE analysis on 24-cm gels (pI 5 to 8), and a total of 27 protein spots were found to be dynamically changed in the N2a cells based on the average intensity ratios of proteins detected (Fig. 1D and E). Among the upregulated protein spots in infected N2a cells (Fig. 2 and Table 2), 10 protein spots were newly induced at 48 h.p.i., and 10 protein spots were newly induced at 96 h.p.i. Among the downregulated protein spots (Fig. 2 and Table 2), 1 protein spot disappeared at 96 h.p.i., and 4 protein spots decreased in intensity at 96 h.p.i. In essence, most of the differentially expressed proteins were found at 48 and 96 h.p.i.

Mass spectrometry (MS) identification of differentially expressed proteins. To identify the differentially expressed proteins in RABV-infected N2a cells, 27 protein spots with a threshold greater than 2-fold were selected for in-gel trypsin digestion and subsequent matrix-assisted laser desorption ionization–tandem time of flight (MALDI-TOF/TOF) identification, resulting in the successful identification of 27 protein spots corresponding to 24 proteins (MS and tandem MS [MS/MS] spectra are shown in [Table 2](#); see also Fig. S1 in the supplemental material). Among these were 22 significantly upregulated protein spots matching 20 proteins (ratio for infected/mock-infected cells of ≥ 2 , [Table 2](#)) and 5 significantly downregulated protein spots matching 4 proteins (ratio for infected/mock-infected cells of ≤ 0.5 , [Table 2](#)). According to the annotations from the UniProt Knowledgebase (Swiss-Pro/TrEMBL) and Gene Ontology Database, the differentially expressed proteins that were identified are related to the cytoskeleton, macromolecular biosynthesis and metabolism, stress response, energy metabolism, immune response, protein translation and elongation, RNA processing and biosynthesis, signal transduction, and the ubiquitin-proteasome pathway.

Interestingly, several protein spots (e.g., nucleoprotein and peripherin) were identified as the product of the same gene, suggesting that they may have different isoforms and/or posttranslational modifications. Of note, some of the cellular proteins, such as hypothetical protein LOC433182, Hsp90, CCT γ , and OAS1B, were newly induced in RABV-infected N2a cells.

Transcriptional profiles of differentially expressed proteins during RABV infection. To determine the reliability of our MS analysis, the genes encoding the 13 differentially expressed proteins in RABV-infected cells were selected for validation by real-time PCR ([Fig. 3B](#)) using primers shown in [Table 1](#). The *GAPDH* mRNA transcript was used as an internal control housekeeping gene. Generally, the changes in abundance of most of the mRNA transcripts were consistent with those of the corresponding proteins in the 2-DE gels ([Fig. 3A](#) and [Table 2](#)). Among the genes selected for validation, the abundance of the *Hsp90*, *CCT γ* , *ACTB*, and *OAS1B* transcripts increased from 12 to 96 h.p.i., while the transcript abundance of *HNRPD* only increased at 96 h.p.i. Intriguingly, a fluctuating expression pattern was observed for some gene transcripts. For example, *CCT θ* , *Prph1*, *EIF2S2*, *DLST*, *PSMA6*, *SEPT2b*, and *SET* were upregulated at 12 and 96 h.p.i. but downregulated at 48 h.p.i. The *Srm* mRNA was elevated at 12 h.p.i. but then became significantly downregulated at 48 h and 96 h.p.i. Gene expression was affected by many factors, and we have long known that the correlation between mRNA and protein abundance is typically weak. In this study, although gene transcript levels and protein expression were not absolutely consistent, the results provide information complementary to the differential proteomics data obtained with 2-DE and MS/MS.

Cellular chaperonins are recruited to the viral N/P protein complex. The 2-DE analysis showed that RABV replication induced the persistent upregulation of cellular chaperonins CCT γ and Hsp90 in mouse N2a cells ([Fig. 2](#) and [3A](#)). Quantitative real-time PCR also revealed that the *CCT γ* and *Hsp90* transcripts were upregulated in N2a cells during RABV infection ([Fig. 3B](#)) compared with their levels in the mock-infected cells. Therefore, the relevance of CCT γ and Hsp90 to the localization of N or P protein of RABV was then analyzed by confocal microscopy. In RABV-infected N2a cells, the hollow ringlike inclusions containing the N ([Fig. 4A](#)) or P ([Fig. 4B](#)) viral proteins were colocalized with intra-

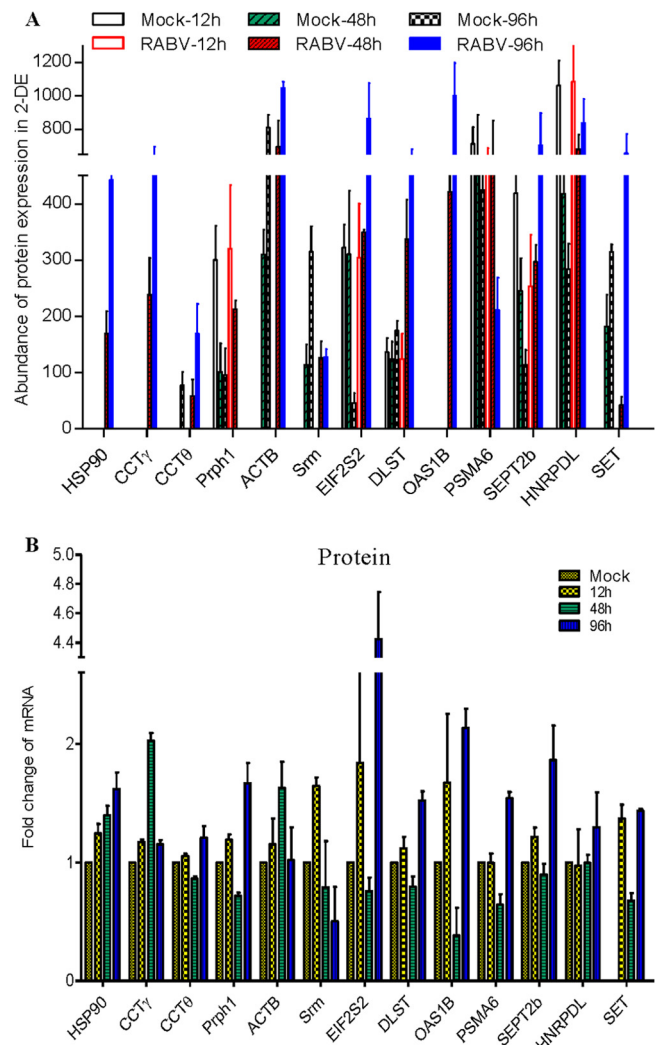
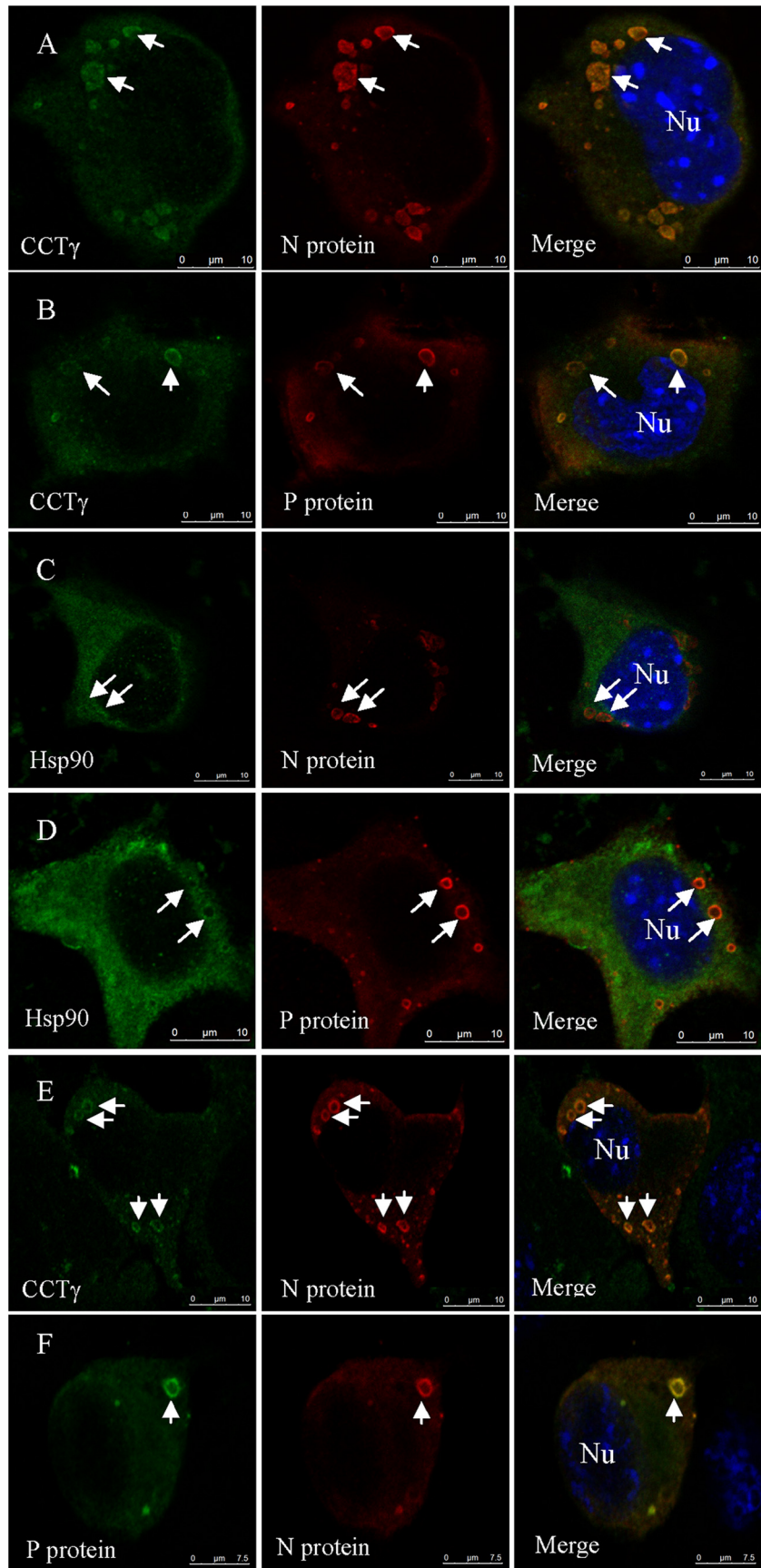


FIG 3 Translational and transcriptional profiling of altered proteins in RABV-infected mouse N2a cells. (A) Translational profiling of the differentially expressed proteins in 2-DE. The spot densities (total pixel intensity within spot boundaries) in 2-DE maps were carried out with PDQuest 2-D analysis software 8.0.1. (B) Total cellular mRNA of mouse N2a cells with or without RABV infection was analyzed by quantitative real-time PCR. All samples were normalized with the results for the *GAPDH* gene as an internal control and with the results for mock-infected N2a cells at each time point as the reference. The values are quantitation ratios of fold increase or decrease relative to the results for mock-infected cells, and mock-infected-cell transcripts were normalized to 1. Error bars represent standard deviations.

cellular CCT γ and were confirmed to contain the viral protein N and P (data not shown). Meanwhile, cytosolic Hsp90 showed some colocalization with the hollow ring-like inclusions containing N ([Fig. 4C](#)) and P ([Fig. 4D](#)). However, in mock-infected N2a cells, CCT γ and Hsp90 were homogeneously distributed in the cytoplasm of N2a cells (data not shown). Similarly, confocal microscopy also showed that CCT γ colocalized with the viral protein N as a hollow circle-like structure in mouse N2a cells cotransfected with the viral N and P genes ([Fig. 4E](#)). Notably, there was no colocalization of CCT γ and Hsp90 with viral proteins in N2a cells that only expressed either N or P alone (data not shown). To further investigate whether the hollow ring-like inclusions are composed of the viral proteins N and P, N2a cells cotransfected



with the viral N and P genes were labeled with MAbs to viral proteins N and P. Confocal microscopy analysis showed that the colocalization of the viral proteins N and P formed a ring-like inclusion in the cotransfected N2a cell (Fig. 4F). These results indicate that the viral N/P protein complex formed a ring-like structure similar to NBs and recruited the aggregation of intracellular chaperonin CCT γ .

To detect whether CCT γ interacts directly with viral protein N or P of RABV, the coimmunoprecipitation assay was performed in the N2a cells infected with RABV and cotransfected with CCT γ and the viral gene encoding N or P. Regrettably, the coimmunoprecipitation analysis could not confirm the interaction of CCT γ and the viral protein N or P in either RABV-infected N2a cells or cotransfected N2a cells (data not shown), suggesting that interactions between CCT γ and viral protein N or P were either weak or indirect at the lowest detectable levels.

Knockdown of CCT γ results in reduction of infectious RABV progeny. To examine the potential role of CCT γ in the life cycle of RABV, we further performed small interfering RNA (siRNA)-mediated knockdown of CCT γ . The CCT γ -specific shRNA or nontargeting control shRNA were transfected into N2a cells by lentivirus-mediated shRNA transfer to construct cell lines stably expressing the shRNAs. Western blot analysis confirmed that CCT γ expression was significantly decreased in the N2a cells transfected with CCT γ shRNA (Fig. 5A). To exclude the possibility that the viability of CCT γ -silenced cells is affected, the viability of shCCT γ -transfected N2a cells was detected by CCK-8 assay. The results showed that the viability of shCCT γ -transfected N2a cells was decreased slightly in comparison with that of N2a and nontargeting-shc002v-transfected N2a cells ($P > 0.05$) (Fig. 5B). Subsequently, CCT γ -silenced N2a cells were inoculated with the Flury strain of RABV at a multiplicity of infection of 0.1, and RABV replication was assessed by determining the virus titer (TCID₅₀) and performing real-time reverse transcription (RT)-PCR. The data showed that the virus titer and viral genomic RNA of RABV were significantly decreased ($P < 0.01$) in CCT γ -silenced N2a cells (Fig. 5C and E) compared to the results for the parental N2a cells and N2a-shc002v cells (transfected with nontargeting shRNA). Also, Western blot analysis showed that the expression of RABV N protein in CCT γ -silenced N2a cells was significantly decreased ($P < 0.01$) in comparison with its expression in the parental N2a cells and N2a-shc002v cells (Fig. 5D). Further transcriptional analysis of the viral N gene demonstrated that the mRNA of the RABV N gene was significantly downregulated in CCT γ -silenced N2a cells treated with or without CHX ($P < 0.01$) (Fig. 5F), indicating that CCT γ affected RABV replication at the mRNA level. Collectively, our data show that CCT γ is a host factor that is required for the transcription and replication of RABV. Correspondingly, RABV replication was monitored by performing quantitative real-time PCR and determining the virus titer for CCT γ -overexpressing N2a cells. As shown by the results in Figure 6, the virus titer in the cell supernatants and cellular viral

genomic RNA had no significant changes ($P > 0.05$) in CCT γ -overexpressing N2a cells, indicating that RABV replication was not upregulated in CCT γ -overexpressing N2a cells.

DISCUSSION

Our study demonstrates colocalization of CCT γ with hollow ring-like structures of NBs containing viral proteins N and P in N2a cells infected by RABV or cotransfected with N and P expression vectors (Fig. 4). Meanwhile, the colocalization of viral proteins N and P was confirmed in N2a cells infected by RABV or cotransfected with N and P expression vectors. The N protein was previously reported to interact with P of RABV (42). However, we found that in N2a cells transfected only with the RABV viral N or P gene, CCT γ was not colocalized with either expressed viral protein (data not shown), indicating that the colocalization and interaction of viral proteins N and P is crucial to NBs. Several research teams reported that neurotropic RABV induces the formation of cytoplasmic inclusion bodies (NBs), which are sites of RABV genomic transcription and replication (12) and have been demonstrated to contain Toll-like receptor 3 (TLR3) (26), Hsp70 (11, 26), viral proteins N and P, and eNOS (10). Taken together, these data confirmed that only the RABV N/P complex and not the individual viral proteins could recruit CCT γ molecules to NBs and that the subcellular distribution of CCT γ was altered in RABV-infected cells. Indeed, our further investigation also showed that the infectious RABV progeny and mRNA were decreased in CCT γ -knockdown N2a cells inoculated with RABV (Fig. 5). Moreover, knockdown of CCT γ did not affect the formation of the NB-like structures (data not shown). These findings demonstrate that CCT γ plays a positive regulatory role in RABV genome replication and that CCT γ is a component but not a core factor of NBs. However, the exact mechanism of CCT γ function in the RABV life cycle needs further investigation.

In the present study, CCT γ overexpression did not show markedly enhanced virus replication in RABV-infected cells as determined by detection of viral genomic RNA and viral titer (Fig. 6). Similarly, in a recent research report (11), the investigators demonstrated that Hsp70 played a proviral role in RABV infection and also found that Hsp70 overexpression only resulted in a slight increase of viral protein synthesis or viral production. Cellular chaperones were involved primarily in assisting the folding of newly synthesized and stress-denatured polypeptides, and they also assisted in the assembly of folded subunits into oligomeric structures (43). As a central mediator of cytosolic protein folding and assembly, CCT provides a physically defined compartment inside which a protein domain or a complete protein can fold while being sequestered from the cytosol (32). Hsp70 is known to be upstream from CCT, is the first target chaperone to bind when the nascent polypeptide chain comes out from the ribosome to help attain a quaternary structure, and will then be transferred to chaperonin CCT for its final folding (44, 45). In *in vivo* and *in vitro* experiments, Hsp70 was found to associate with chaperonin CCT,

FIG 4 Subcellular distribution of CCT γ and Hsp90 in N2a cells by confocal microscopy. Immunostaining was conducted using MAbs to RABV N and P proteins followed by TRITC-conjugated IgG (red) and FITC-conjugated IgG (green) and counterstaining using rabbit antibodies to CCT γ and Hsp90 followed by FITC-conjugated IgG (green). Nuclei (Nu) were stained with DAPI (blue). The triple-stained cells were observed by confocal microscopy. (A and B) Colocalization of CCT γ with viral proteins N and P, revealing a hollow ring-like structure in RABV-infected N2a cells. (C and D) Hsp90 has some colocalization with the hollow ring-like structure containing viral N and P proteins in RABV-infected N2a cells. (E) CCT γ colocalized with viral protein N, revealing a hollow ring-like structure in mouse N2a cells cotransfected with RABV N and P genes. (F) Colocalization of viral proteins P (green) and N (red) formed a hollow ring-like structure (Merge) in N2a cells cotransfected with RABV N and P genes. Scale bars = 10 μ m.

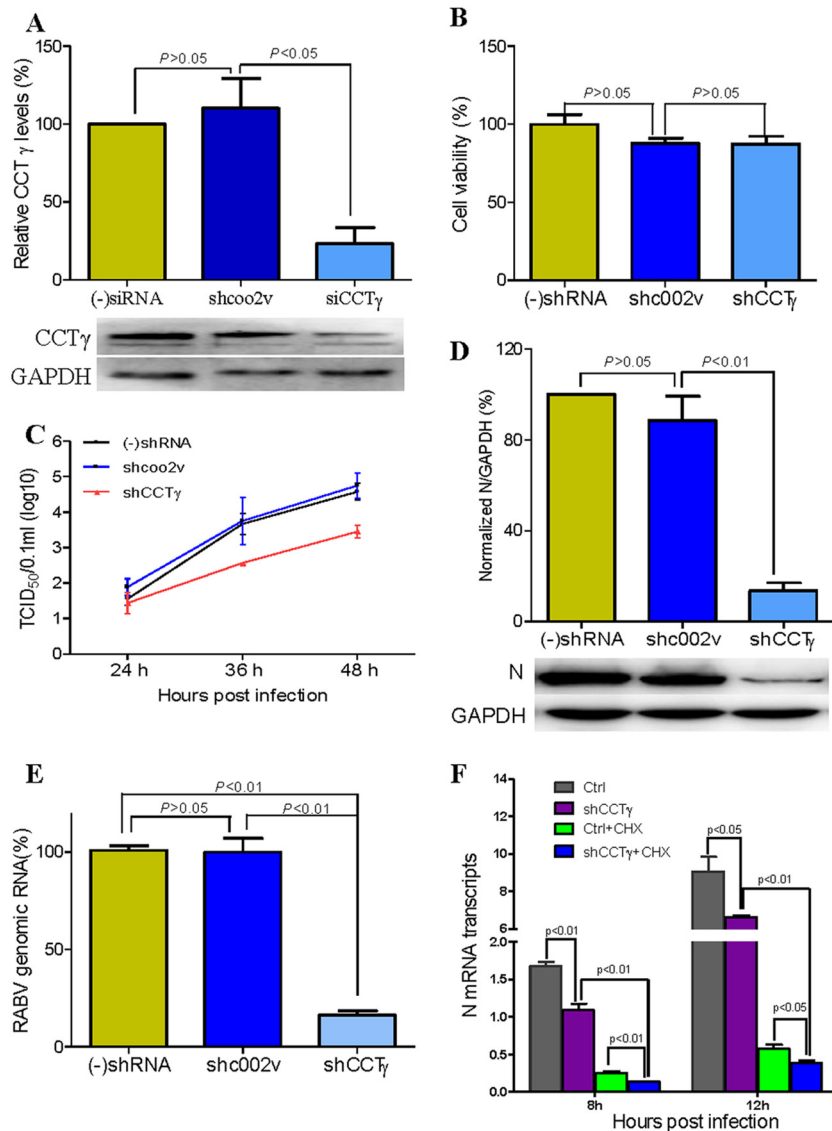


FIG 5 CCT γ knockdown inhibits RABV replication and transcription. (A) Western blotting of CCT γ -silenced N2a cells. (-)shRNA, no shRNA. (B) Viability assay of CCT γ -silenced N2a cells. Cell viability is expressed as the percentage of (-)shRNA N2a cells. (C) Growth curve of RABV in CCT γ -silenced N2a cells. N2a cells only transfected with nontargeting shRNA shcoo2v or CCT γ -specific shRNA, as well as no shRNA, used as positive control [(-)shRNA], were infected at an MOI of 0.1. At 48 h.p.i., the virus titer in cells transfected with CCT γ -specific shRNA decreased nearly 20-fold in comparison with that in cells transfected with nontargeting shRNA shcoo2v. (D) Expression of RABV N protein in N2a cells as determined by Western blotting. (E) Quantitation of RABV genomic RNA by quantitative real-time PCR. The genomic RNA of RABV in shRNA-silenced cells was decreased \sim 80% ($P < 0.01$) in comparison with that in cells transfected with nontargeting shRNA shcoo2v. (F) The mRNA transcripts of RABV N gene as determined by real-time comparative quantitative PCR. The transcription of RABV N was decreased significantly after silencing of CCT γ , although CHX interfered significantly in the N2a cells transfected with and without CCT γ -specific shRNA. Values are means \pm standard deviations of three independent experiments.

suggesting their cooperation in the process of protein folding (43, 46, 47). Here, a reasonable hypothesis is that the functions of CCT γ in the RABV life cycle may need other cellular factors' cooperation. Interestingly, our 2-DE data also showed that chaperonins Hsp90, CCT γ , and CCT θ were upregulated in RABV-infected cells. Meanwhile, Hsp90 is involved in intracellular transport, complex assembly and disassembly, and protein maturation (24) and is reported to be upregulated in the brainstem and hippocampus of dogs naturally infected by RABV (24) and in RABV-infected mouse brain (18). Therefore, the proteomic data seems to provide further evidence to support our suggestion.

As for the interaction of CCT γ with other viral proteins, a previous study reported that a yeast two-hybrid screening experiment revealed that type D retrovirus p4 specifically interacted with CCT γ (33), and CCT γ also associated with high specificity with the Mason-Pfizer monkey virus pp23/16-p12 domain and human immunodeficiency virus p6. In addition, the crystal structure of the mouse CCT γ apical domain has been determined, and the signature region on the CCT γ apical domains is expected to be involved in the specific binding of its appropriate substrate, partially folded tubulin (48). A recent study demonstrated the interaction of Hsp70 with viral protein N of RABV (11). Unfortu-

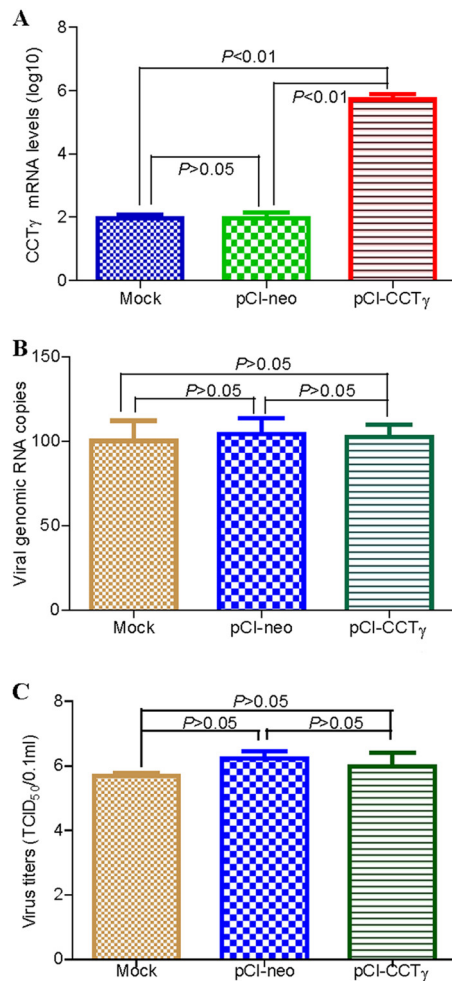


FIG 6 RABV replication in CCT γ -overexpressing N2a cells. (A) CCT γ mRNA transcript. CCT γ mRNA was detected from the transfected N2a cells for 48 h by comparative quantitative PCR (qPCR). *GAPDH* gene was used as a control. (B) Detection of cellular RABV genomic RNA. The cellular RABV genomic RNA was quantified by absolute qPCR from the RABV-infected N2a cells for 48 h. The RABV genomic copies in the mock N2a cells were normalized to be 100%. (C) Virus titer of RABV. The virus titer in cell supernatants is represented as TCID₅₀.

nately, our attempts to coimmunoprecipitate viral proteins N and P and chaperonin CCT γ have failed (data not shown) in N2a cells infected with RABV and cotransfected with N and P expression vectors. Whether CCT γ being recruited to NBs interacts directly or indirectly with RABV proteins remains unclear, and further detailed studies are needed to clarify the exact mechanism involved in this process.

Notably, except for altered cellular chaperone expression, in 2-DE coupled with MALDI-MS/MS identification, we demonstrate that cytoskeletal protein profiles were altered by RABV infection in N2a cells. Microtubule-associated proteins (TUBB5 and TUBA1B) and a microfilament-associated protein (Actb) were identified as being upregulated in RABV infection. In contrast, the intermediate filament protein peripherin1 (Prph1) was downregulated in the RABV-infected N2a cells (Fig. 2). Peripherin is a type III intermediate filament protein that is expressed nearly exclusively by peripheral neurons and some central nervous system

neurons. In both herpes simplex virus and developmental studies, motor neurons were identified by their expression of peripherin (49). Previous *in vivo* studies have shown that peripherin is the specific interaction partner of BPAG1-n (50). Taken together, our data indicate that RABV remodels the expression profile of the cytoskeletal proteins in host cells to favor its own replication.

In addition, in dynamic 2-DE imaging of RABV-infected cells, we observed that several differentially expressed spots on the gel represented the same protein (Fig. 2), including peripherin and viral nucleoprotein, and this phenomenon has also been observed in avibirnavirus-infected CEFs (35) and porcine circovirus 2-infected PK15 cells (36). A previous report (51) suggested that the P protein of RABV exists as five isoforms (P, P2, P3, P4, and P5), and not only are their subcellular distributions different, P and P3 interact directly with promyelocytic leukemia protein. Whether the migration of proteins at different spots is the result of post-translational modification or different isoforms arising during the course of RABV replication and assembly needs further investigation.

In this work, RABV infection resulted in the upregulation of a series of energy metabolism-related proteins, including IDH3A, DLST, LOC433182, and ATP5A1, while TALDO1 was downregulated. Isocitrate dehydrogenase (IDH) is a member of the β -decarboxylating dehydrogenase family of enzymes and catalyzes the oxidative decarboxylation of isocitrate to 2-oxoglutarate (52). NAD⁺-dependent isocitrate dehydrogenases catalyze the allosterically regulated rate-limiting step of the tricarboxylic acid cycle. DLST (E2) is the structural and catalytic core of the 2-oxo acid dehydrogenase multienzyme complex that catalyzes lipoic acid-mediated and coenzyme A- and NAD⁺-linked decarboxylation of 2-oxo acids (53). As part of this complex, the induction of DLST may be regarded as an attempt to compensate for the total energy loss in the cell due to the synthesis and transport of viral components.

In summary, our data indicate that the host protein CCT γ is associated with but not a core factor of NBs in RABV-infected N2a cells and contributes to RABV genomic replication. Although further studies are needed to determine the exact mechanism of CCT γ function in the RABV life cycle, the analyses performed in this study highlight the potential of host proteins as useful new targets for drug development.

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