

Measles Virus Defective Interfering RNAs Are Generated Frequently and Early in the Absence of C Protein and Can Be Destabilized by Adenosine Deaminase Acting on RNA-1-Like Hypermutations

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ABSTRACT

Defective interfering RNAs (DI-RNAs) of the viral genome can form during infections of negative-strand RNA viruses and outgrow full-length viral genomes, thereby modulating the severity and duration of infection. Here we document the frequent *de novo* generation of copy-back DI-RNAs from independent rescue events both for a vaccine measles virus (vac2) and for a wildtype measles virus (IC323) as early as passage 1 after virus rescue. Moreover, vaccine and wild-type C-protein-deficient (C-protein-knockout [C^{KO}]) measles viruses generated about 10 times more DI-RNAs than parental virus, suggesting that C enhances the processivity of the viral polymerase. We obtained the nucleotide sequences of 65 individual DI-RNAs, identified breakpoints and reinitiation sites, and predicted their structural features. Several DI-RNAs possessed clusters of A-to-G or U-to-C transitions. Sequences flanking these mutation sites were characteristic of those favored by adenosine deaminase acting on RNA-1 (ADAR1), which catalyzes in double-stranded RNA the C-6 deamination of adenosine to produce inosine, which is recognized as guanosine, a process known as A-to-I RNA editing. In individual DI-RNAs the transitions were of the same type and occurred on both sides of the breakpoint. These patterns of mutations suggest that ADAR1 edits unencapsidated DI-RNAs that form doublestrand RNA structures. Encapsidated DI-RNAs were incorporated into virus particles, which reduced the infectivity of virus stocks. The C^{KO} phenotype was dominant: DI-RNAs derived from vac2 with a C^{KO} suppressed the replication of vac2, as shown by coinfections of interferon-incompetent lymphatic cells with viruses expressing different fluorescent reporter proteins. In contrast, coinfection with a C-protein-expressing virus did not counteract the suppressive phenotype of DI-RNAs.

IMPORTANCE

Recombinant measles viruses (MVs) are in clinical trials as cancer therapeutics and as vectored vaccines for HIV-AIDS and other infectious diseases. The efficacy of MV-based vectors depends on their replication proficiency and immune activation capacity. Here we document that copy-back defective interfering RNAs (DI-RNAs) are generated by recombinant vaccine and wild-type MVs immediately after rescue. The MV C protein interferes with DI-RNA generation and may enhance the processivity of the viral polymerase. We frequently detected clusters of A-to-G or U-to-C transitions and noted that sequences flanking individual mutations contain motifs favoring recognition by the adenosine deaminase acting on RNA-1 (ADAR1). The consistent type of transitions on the DI-RNAs indicates that these are direct substrates for editing by ADAR1. The ADAR1-mediated biased hyper-mutation events are consistent with the protein kinase R (PKR)-ADAR1 balancing model of innate immunity activation. We show by coinfection that the C-defective phenotype is dominant.

Measles virus (MV) is a negative-strand RNA virus of the *Paramyxoviridae* family with a genome comprising 15,894 nucleotides (nt). The MV genome encodes six genes from which eight proteins are expressed: nucleoprotein (N), phosphoprotein (P), matrix protein (M), fusion protein (F), hemagglutinin (H), and large protein (L; the catalytic subunit of the viral RNA-dependent RNA polymerase), as well as the host response-modulating V and C proteins (1–3).

MV replication occurs in the cytoplasm of infected cells and involves a helical ribonucleocapsid (RNP) formed by viral genomic RNA and N protein. The polymerase complex, consisting of the L and P proteins, recognizes the 3' end of the RNA and generates a complementary nascent RNA strand that is cotranscriptionally encapsidated (4). Both genomic RNA and antigenomic RNA can serve as a template for replication, whereas only the genomic RNA serves as a template for the transcription of capped and polyadenylated mRNAs (1).

The polymerase of negative-strand RNA viruses is error prone

and introduces nucleotide substitutions at fairly high frequencies (about 1 substitution in 10,000 nt) (5–8). Defective genomes are also generated, including abortive genomes and 5' copy-back defective interfering (DI) RNAs (DI-RNAs) (9). DI-RNAs are generated when the polymerase prematurely terminates genome synthesis and then—when it is still bound to the truncated ge-

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nome—binds back and reinitiates replication on this RNA, thereby generating a genome-antigenome hybrid with perfectly complementary ends. These complementary ends can form stable panhandle duplex RNA structures, as shown for MV and Sendai virus (SeV), a related paramyxovirus (9, 10).

These double-stranded RNA (dsRNA) structures are potent inducers of the cellular stress and innate immune response (11). They trigger the activation of innate receptors, including retinoic acid-inducible gene I (RIG-I [12]), melanoma differentiation-associated gene-5 (mda-5 [13]), and protein kinase R (PKR [14]). DI-RNAs contain the 3'-terminal promoter sequence and therefore can be replicated into their complementary counterpart, which also contains the promoter sequence at its 3' end. DI-RNAs, therefore, like the genome-antigenome pair, form a replication system, which, due to its reduced length and therefore increased replication frequency, can interfere with full-length RNA replication and temporarily outgrow full-length RNA (15).

We previously reported the generation of viral DI-RNAs in C-protein-deficient (C-protein-knockout [C^{KO}]) Moraten vaccine MV (MV-CKO) and showed that DI-RNAs induce the PKRmediated stress and interferon (IFN) responses (14). Here we provide mechanistic insights into the generation of DI-RNAs by a vaccine MV (vac2)-derived infectious cDNA (16) and a wild-type MV (IC323) infectious cDNA (17) immediately after virus rescue. We describe the molecular characteristics of DI-RNAs from Cprotein-expressing and C-defective viruses and find structural similarities between DI-RNAs derived from all viruses. Our analysis of multiple independent virus rescues indicated that deficiency in C-protein expression leads to a 10-fold increase in the abundance of DI-RNAs. Although initially demonstrated in serially undiluted virus infections performed at a high multiplicity (18-20), we detected DI-RNAs in stocks as early as passage 1 (P1) after virus rescue. We found clusters of A-to-G or U-to-C transitions in several DI-RNAs in a sequence context characteristic of the mutations introduced by cellular adenosine deaminase acting on RNA-1 (ADAR1) (21-23).

MATERIALS AND METHODS

Cells and viruses. Vero, Vero-hSLAM (a kind gift from Y. Yanagi [24]), and HeLa cells were cultivated in Dulbecco's high-glucose modified Eagle's medium (D-MEM; HyClone; GE Healthcare Life Sciences, Logan, UT) supplemented with 5% (vol/vol) fetal bovine serum (FBS; Gibco, Life Technologies, Carlsbad, CA). Vero-hSLAM cells were grown in the presence of Geneticin (G418; Corning, Tewksbury, MA) at a final concentration of 0.4 mg/ml. 293-3-46 cells (25), used for virus rescue, were cultivated in D-MEM supplemented with 10% (vol/vol) FBS and 1 mg/ml G418. The mantle cell lymphoma cell line Granta-519 (26) was cultivated in RPMI 1640 (HyClone) supplemented with 10% (vol/vol) FBS.

Recombinant MVs were IC323 (isogenic to the wild-type IC-B strain [17]), IC323-C^{KO}, vac2 (isogenic to the Moraten vaccine strain [16]), vac2-C^{KO}, vac2(GFP) (expressing enhanced green fluorescent protein [eGFP] from an additional transcription unit downstream of H [27]), vac2-C^{KO}(GFP), vac2(mCherryNLS), and vac2-C^{KO}(mCherryNLS). Viruses with the C^{KO} have specific mutations introducing early stop codons in the C open reading frame, as described previously (16).

Generation of mCherryNLS-expressing MVvac2 cDNA. A construct encoding mCherry with a nuclear localization signal (NLS), mCherryNLS, kindly provided by L. Enquist (28), was cloned into p(+)MV323(GFP)N to replace GFP, yielding p(+)MV323(mCherryNLS)N (Crystal Mendoza, unpublished data). mCherryNLS was amplified from this donor vector by PCR using the following primers (Integrated DNA Technologies [IDT], Coralville, IA), where the restriction sites are underlined: mCherryNLS EcoRI/MluI fwd (TATA<u>GAATTC</u>CCATCA<u>ACGCG</u> <u>T</u>TACGCCACCATG) and mCherryNLS NotI/AatII rev (ATAT<u>GCGGC</u> <u>CGC</u>GTTCGA<u>GACGTC</u>AGTTTATCTGGATCC). The PCR product was cloned into pCR3 (Life Technologies, Invitrogen, Carlsbad, CA) using the EcoRI/NotI restriction sites and then excised with MluI/AatII (New England BioLabs, Ipswich, MA). The GFP-encoding sequence in pB(+)MVvac2(GFP)H and pB(+)MVvac2-C^{KO}(GFP)H was then replaced with the mCherryNLS fragment to yield pB(+)MVvac2 (mCherryNLS)H and pB(+)MVvac2-C^{KO}(mCherryNLS)H, respectively.

Virus rescue and stock production. Recombinant viruses were generated as described previously (25). Briefly, 293-3-46 cells were transfected with the relevant plasmid carrying full-length viral cDNA and pEMC-La, expressing the viral polymerase. After a 42°C heat shock and 2 days of incubation, cells were transferred to 100-mm dishes with Vero or Vero-hSLAM cells and incubated until syncytium formation was observed. Single syncytia were picked and used to inoculate Vero or VerohSLAM cells in 6-well plates. Cells and supernatants were harvested when the maximum cytopathic effect (CPE) was achieved (2 to 4 days), yielding passage 0 stocks. These stocks were used to generate passage 1 stocks by infecting multiple 150-mm dishes of Vero or Vero-hSLAM cells. Virus stocks were grown at 32°C (vac2) or 37°C (IC323). Cells were harvested at the time of the maximum CPE by scraping them into 1 ml Opti-MEM medium (Gibco, Life Technologies, Carlsbad, CA) per 150-mm dish, followed by three freeze-thaw cycles (liquid nitrogen and 37°C). Cleared lysates were aliquoted and stored at -80°C. Viral titers on Vero-hSLAM cells were determined using the 50% tissue culture infectious dose (TCID₅₀) method. For further passages, Vero cells in 150-mm dishes were infected at a multiplicity of infection (MOI) of 0.03 and harvested, and titers were determined.

Virion purification. Vero cells (in 10 150-mm dishes) were infected with passage 1 vac2 or vac2-C^{KO} at an MOI of 0.05. Cells were harvested at 48 h postinfection and scraped into 1 ml Opti-MEM medium per dish, followed by three freeze-thaw cycles. Cleared lysates (total volume, about 15 ml) were mixed with Coomassie blue (Brilliant Blue G; final concentration, ~150 µg/ml; Bio-Rad, Hercules, CA), and the mixture was laid onto a 20%/60% (wt/vol) sucrose cushion (in TNE buffer [50 mM Tris-HCl {pH 7.4}, 100 mM NaCl, 0.1 mM EDTA]) and centrifuged in an SW32Ti rotor (Beckman Coulter, Indianapolis, IN) at 75,000 \times g at 4°C for 4 h. The visible blue band at the 20%-60% interface was collected with a 16.5-gauge needle and syringe in a total volume of about 6 ml, diluted with sterile phosphate-buffered saline (PBS; HyClone) supplemented with 0.5 mM MgCl₂ (PBS-MgCl₂) to a final volume of 15 ml, and then laid onto a 20 to 60% (wt/vol) discontinuous sucrose gradient (in TNE buffer, 5% intervals). Centrifugation was to equilibrium (75,000 × g at 4°C for 16 h) using the SW32Ti rotor. Virions that accumulated in the 30 to 40% region were collected in a total volume of 6 ml, diluted with sterile PBS-MgCl₂, and concentrated using Amicon Ultra filter tubes (EMD Millipore, Billerica, MA) with a molecular mass exclusion of 100 kDa (3,500 \times g, 4°C) in a tabletop centrifuge. The concentrate was washed once with 15 ml PBS-MgCl₂, concentrated to a final volume of 1 ml, and stored at -80° C. Titers were determined by the TCID₅₀ method. Prior to further analysis, purified stocks were subjected to treatment with Benzonase (EMD Millipore, Billerica, MA) at a final concentration of 12.5 U/µl for 2 h at 37°C in 10 mM Tris, pH 8.0, buffer containing 1 mM MgCl₂ and 10 mM NaCl.

RNA protection assay. One million TCID_{50} s of purified vac2 or vac2- C^{KO} virions were treated with combinations of nonionic detergent (Triton X-100), protease (trypsin), and nuclease (Benzonase). First, 1% (vol/ vol) Triton X-100 (Sigma-Aldrich, St. Louis, MO) was added to virions in reaction buffer (10 mM Tris, pH 8.0, 10 mM NaCl, 1 mM MgCl₂). Next, samples were heated to 80°C for 15 min and cooled on ice for 10 min. Then, 0.1 mg/ml trypsin (Worthington, Lakewood, NJ) was added and the samples were incubated at 37°C for 15 min. To inactivate the trypsin, 3 mg/ml soybean trypsin inhibitor (Worthington, Lakewood, NJ) was

added and the samples were incubated at 25°C for 30 min. Next, 2.5 U/ μ l Benzonase was added and the samples were incubated at 37°C for 2.5 h. Nuclease activity was inhibited by addition of 10 mM EDTA, before sample aliquots were subjected to protein analysis or RNA extraction.

RNA extraction, RT-PCR, and PCR analysis. Vero or Vero-hSLAM cells were infected with viruses at an MOI of 0.1 or left uninfected and harvested at 30 to 48 h postinfection. Total RNA was extracted using either an RNeasy minikit (Qiagen, Valencia, CA) or the TRIzol reagent (Ambion, Life Technologies, Carlsbad, CA) according to the suppliers' instructions and stored at -80°C. One microgram of total RNA was subjected to reverse transcription-PCR (RT-PCR) using SuperScript III reverse transcriptase (Life Technologies, Carlsbad, CA) with a mixture of MV-specific primers (IDT), where restriction sites are underlined: negative-polarity primers A1_{SnaBI} (GCGCGC<u>TACGTA</u>AAAGCTGGGAATA GAAACTTCG), A2_{vac2-NotI} (TAGAGAGAGAGCGGCCGCTCTGGTGTAA GTCTAGTATCAGA), A2_{323-Not}I (TAGAGAGA<u>GCGGCCGC</u>TCTGGT GTAAGTCTAGTACTAGA), A3_{NotI} (TAGAGAGAGAGCGGCCGCGATCT AGAATTGGCGGAAACC), and A4_{NotI} (TAGAGAGAGAGAGCGGCCGCTCC CTGATGGTCTATGATAATG) and positive-polarity primer B1 (ATGA CAGATCTCAAGGCTAAC). Briefly, RNA was mixed with primers and deoxynucleoside triphosphates and subjected to heat denaturation at 98°C for 10 min, followed by incubation on ice for 10 min. After addition of the enzyme mix, samples were incubated at 55°C for 60 min and then at 70°C for 15 min. cDNAs were stored at -20°C. An aliquot of the cDNA (1 µl) was further amplified using GoTaq polymerase (Promega, Madison, WI) and DI-RNA-specific primer combinations (primers A1/A2, A1/A3, and A1/A4) or a control primer pair for standard genomes (primers A1/ B1). Samples were analyzed by electrophoresis using 1% (wt/vol) agarose gels and a 100-bp or 1-kb DNA ladder as a size standard (Promega, Madison, WI).

Cloning and analysis of DI-RNAs. PCR products amplified from the cDNA of MV-infected cells were purified (PCR purification kit; Qiagen, Valencia, CA), digested with SnaBI and NotI-HF (New England BioLabs, Ipswich, MA), purified again, and then ligated with EcoRV/NotI-linearized pCR3 (Invitrogen, Life Technologies, Carlsbad, CA). Ligation products were transformed into OneShot TOP10 cells (Life Technologies, Carlsbad, CA), and the bacteria were plated on LB-ampicillin-agar plates and incubated at 37°C for 12 to 20 h. Single colonies were picked and resuspended in 10 µl H₂O, and 5 µl of the suspension was subjected to colony PCR using the primers (IDT) T7_{fwd} (TAATACGACTCACTATA GGG) and BGH_{rev} (TAGAAGGCACAGTCGAGG) and a 2× PCR mix (Promega, Madison, WI). Clones with individual inserts, as analyzed by 1% agarose gel electrophoresis, were selected and grown in 2 ml LBampicillin, and plasmids were extracted using a plasmid minikit (Qiagen, Valencia, CA). Sequences were determined by Sanger sequencing using the T7_{fwd} and BGH_{rev} primers and analyzed by alignment to MV reference sequences with GenBank accession numbers AF266287 for vac2 and AB016162 for IC323.

Protein gel electrophoresis and immunoblot analysis. Protein extracts of infected cells (MOI = 0.1, 30 to 48 h postinfection) were generated, and protein concentrations were determined as described previously (14). Protein (25 μ g) was fractionated using 12% polyacrylamide gels and transferred to Immobilon-P membranes (EMD Millipore, Billerica, MA) using a semidry protocol. Membranes were blocked with 5% (wt/vol) bovine serum albumin (Sigma-Aldrich, St. Louis, MO) in Tris-buffered saline (Bio-Rad, Hercules, CA) for 1 h and incubated with primary antibodies at 4°C for 12 to 16 h. Membranes were washed three times with PBS with 0.5% (vol/vol) Tween 20 (PBST) for 5 to 10 min, incubated with horseradish peroxidase (HRP)-conjugated secondary antibody at room temperature for 1 h, washed again three times with PBST, and incubated with SuperSignal West Pico chemiluminescent substrate (Pierce, Thermo Fisher Scientific, Rockford, IL). Membranes were exposed to HyBlot films (Denville, South Plainfield, NJ) for various amounts of time.

Antibodies. Rabbit peptide antisera recognizing MV N (N₅₀₅ antiserum), MV C (C_2 antiserum), and MV H (H_{cyt} antiserum) were described

previously (16, 29, 30) and were used at dilutions of 1:5,000 for primary antibody incubation. Mouse monoclonal anti-GFP antibody (eBioscience, San Diego, CA) was used at a 1:1,000 dilution. HRP-conjugated antiactin (Sigma-Aldrich, St. Louis, MO) was used at a 1:25,000 dilution. Rabbit and mouse secondary antibodies conjugated with HRP (Jackson ImmunoResearch, West Grove, PA, and EMD Millipore, Billerica, MA, respectively) were used at 1:10,000 dilutions. For immunofluorescence, mouse anti-dsRNA J2 (English and Scientific Consulting, Szirák, Hungary [31]), anti-MV N₅₀₅ antiserum, and secondary antibodies antimouse Alexa 488, antimouse Alexa 594, and antirabbit Alexa 405 (Molecular Probes, Life Technologies, Carlsbad, CA) were used as described previously (14).

Northern blot analysis. A 250-bp fragment of the L trailer region of MV (nt 15641 to 15890) was cloned into the pCR3 vector between the HindIII and XhoI restriction sites (amplification primers [IDT], MV15641-HindIII fwd [AGACCCAAGCTTTATCTTCGTTAAGAA TCT] and MV15890 XhoI rev [TGCATGCTCGAGGACAAAGCTGG GAATAGA], where the restriction sites are underlined). Positive and negative single-stranded RNA (ssRNA) probes were generated from HindIII- or XhoI-linearized plasmids using SP6 or T7 RNA polymerase of a digoxigenin (DIG) RNA labeling kit (Roche Diagnostics, Indianapolis, IN). Total RNA (3 µg) or an amount of RNA corresponding to 1×10^6 TCID₅₀s of purified virions was subjected to 1% agarose-2% formaldehyde gel electrophoresis, with DIG-labeled RNA molecular weight marker I (Roche Diagnostics, Indianapolis, IN) being included as described in the DIG Northern starter kit manual (Roche Diagnostics, Indianapolis, IN). RNA was transferred to nylon membranes (Roche Diagnostics, Indianapolis, IN) by bottom-up capillary transfer using 20× SSC (3.0 M NaCl, 0.3 M sodium citrate, pH 7.0) as the buffer. RNAs were cross-linked with 125 mJ of 254-nm UV light before incubation with RNA probes (100 mg/ml) at 68°C for 16 to 20 h under constant agitation. Probe detection was performed according to the instructions in the supplier's manual (Roche Diagnostics, Indianapolis, IN).

Confocal immunofluorescence analysis. Infection of HeLa cells and immunostaining for dsRNA and MV N protein were performed as described previously (14). Slides were analyzed using an LSM 780 inverted laser scanning confocal microscope and ZEN 2012 (Black Edition, v8.0) software (Carl Zeiss Microscopy, Jena, Germany).

Flow cytometry analysis of coinfections. Granta-519 cells (5×10^5) were infected with various combinations of GFP- and mCherryNLS-expressing vac2 and vac2-C^{KO} at an MOI of 1 for each virus and fixed with 4% paraformaldehyde (Santa Cruz Biotechnology, Santa Cruz, CA) at 48 h postinfection, and fluorescent reporter expression was analyzed using a FACSCalibur cytometer (BD Biosciences, San Jose, CA) and FlowJo software (Tree Star Inc., Ashland, OR).

RESULTS

DI-RNAs are generated at a high frequency immediately after virus rescue. Copy-back DI-RNAs have two characteristic features: their breakpoint, that is, the last nucleotide incorporated by the polymerase into the progeny genomic RNA before leaving the template antigenome strand (Fig. 1A, left end of black bar), and the reinitiation site, that is, the position of polymerase entry on the progeny RNA. Polymerase reentry results in the synthesis of a cRNA strand of opposite polarity (Fig. 1A, left end of bottom gray bar). Even if such RNA molecules have the potential of forming a dsRNA structure with a hairpin loop (Fig. 1B), they usually do not because the replicating RNAs are cotranscriptionally encapsidated. Copy-back DI-RNA can be specifically amplified by PCR from the cDNA of infected cells by the use of primer pairs in which one primer binds to the copy-back sequence (Fig. 1C, primer A1) and the second primer (primer A2, A3, or A4) binds to the DI sequence downstream of the breakpoint. Although only parts of



FIG 1 Schematic drawings of 5' copy-back DI-RNA generation, predicted secondary structure, and replication. (A) 5' copy-back DI-RNAs are hybrids of truncated genome (black) and antigenome (gray) sequences. During replication, viral polymerase initiates RNA synthesis at the 3' antigenome promoter (trailer; open triangle) and prematurely terminates synthesis at the breakpoint. The polymerase then reinitiates synthesis on the newly generated strand and finishes replication by recopying the antigenomic 3' end. The two ends of the DI-RNA are therefore perfectly complementary to each other. (B) Panhandle structure that may be formed through hybridization of the complementary ends of a copy-back DI-RNA in the absence of sufficient N protein. (C) 5^\prime copy-back DI-RNA replication. The genome DI-RNA [(-)DI-RNA] (top) contains the antigenome promoter in the copy-back sequence (open triangle). Therefore, it can be replicated by the viral polymerase into a complementary antigenome DI-RNA [(+)DI-RNA], which now contains the antigenome promoter at the opposite end. The polarity switch within one DI-RNA molecule allows specific amplification using combinations of primers (primer A1 with either primer A2, A3, or A4; arrows) that would bind in the same direction on a regular antigenome molecule.

the DI-RNA sequences are amplified with these primers (the 5' and 3' termini outside the primer binding sites are not amplified), the amplified sequence permits calculation of the length of the DI-RNA. Importantly, these primer combinations cannot yield a specific PCR product with standard genomes, since there they have the same polarity.

To gain insights into the frequency with which DI-RNAs are generated, we analyzed individual viral plaques obtained during the rescue of vaccine (vac2; Fig. 2A, top left) or wild-type (IC323; Fig. 2A, top right) recombinant MV. In addition, we also isolated plaques of mutant viruses unable to express the C protein (vac2- C^{KO} and IC323- C^{KO} ; Fig. 2A, bottom left and right, respectively). We generated passage 1 (P1) stocks from these plaques and used these to do infections at a low MOI of 0.1.

We designed primers binding to different positions in the L gene, allowing amplification of DI-RNA sequences from less than 0.5 kb to up to 5 kb. Using the A1/A2 primer combination, DI-RNAs were readily detected in at least 2 of 10 individual plaques of C-expressing vac2 (Fig. 2A, top left) and IC323 viruses (Fig. 2A, top right). DI-RNAs were prominent in nearly all plaques of both mutant C^{KO} viruses: multiple DI-RNAs were identified in each vac2- C^{KO} plaque (Fig. 2A, bottom left), and 7 of 10 IC323- C^{KO} plaques (Fig. 2A, bottom right) showed at least one DI-RNA. A unique DI-RNA pattern was detected in each plaque.

Molecular characterization of DI-RNAs from vaccine and wild-type MV strains. To further characterize the DI-RNAs, we cloned their cDNAs and obtained the sequences of 65 independent clones. Table 1 summarizes the data for vaccine-derived DI-



FIG 2 Molecular characterization of DI-RNAs generated by vaccine and wildtype MVs. (A) Analysis of DI-RNAs expressed by multiple clones of vaccine MV (vac2) and wild-type MV (IC323) able to express C protein or not (C^{KO}). Plaques were generated in different rescue experiments, amplified, and analyzed by DI-RNA-specific RT-PCR. Uninfected cells (UI) were included as a negative control. The first number above each lane is the experiment number, and the second number is the plaque number. cDNAs were amplified with the A1/A2 primer combination and analyzed by agarose gel electrophoresis. The numbers on the left of each panel indicate the sizes of the marker bands (in base pairs). (B) Nucleotide positions of the breakpoints and reinitiation sites mapped in DI-RNAs derived from vac2 and vac2-C^{KO} (Table 1). The combinations of primers used for amplification are indicated on the left of the three panels. (C) Breakpoint and reinitiation site analysis of IC323 and IC323-C^{KO} viruses (Table 2).

TABLE 1 Vaccine virus-derived DI-RNAs

			Nucleotide position ^a			Length (no. of nucleotides)				
Virus	Expt no. and plaque no.	DI clone	Breakpoint	Reinitiation site	Clone class ^b	Insertions/ deletions	dsRNA stem	ssRNA loop	DI-RNA length	Rule of six ^c
vac2	1.1	1/01	15560	15750	A		145	190	480	Y
	1.1	13/02	14475	15655		+2	240	1,180	1,662	Ŷ
	3.2	2/01	15286	15796			99	510	708	Y
	3.2	2/04	15412	15778	В		117	366	600	Ŷ
vac2-C ^{KO}	1.1	3/01	15575	15765			130	190	450	Y
	1.1	3/05	15560	15786	С		109	226	444	Y
	1.2	24/01	13306	15759			136	2,453	2,725	Ν
	1.2	24/03	12891	15659			236	2,768	3,240	Y
	1.2	4/01	15535	15787			108	252	468	Y
	1.2	4/06	15560	15792	D		103	232	438	Y
	1.2	4/07	15560	15750	A		145	190	480	Y
	1.3	17/02	14570	15786			109	1.216	1.434	Y
	1.3	17/04	14129	15765			130	1,636	1,896	v
	1.3	17/04	14624	15777		-3	118	1,050	1,386	v
	1.3	17/05	14548	15754		5	1/1	1,105	1,300	v
	1.3	25/07	13655	15741			154	2.086	2 304	v
	1.3	5/01	15494	15735		+3	154	2,000	2,394	I V
	1.5	5/01	15294	15750		15	145	441	754	I V
	1.5	18/05	13264	15750			143	400	1 410	I V
	2.1	18/05	14639	15741			154	1,102	1,410	Y V
	2.1	18/06	142/7	15/6/			128	1,490	1,746	Y V
	2.1	26/03	13/13	15/55			140	2,042	2,322	Y
	2.1	26/05	12612	15728			167	3,116	3,450	Y
	2.1	6/01	15344	15756			139	412	690	Y
	2.1	6/05	15412	15778	В		117	366	600	Y
	2.1	6/12	15402	15621		-107	167	219 + 107	660	Y
	2.2	19/03	13899	15791			104	1,892	2,100	Y
	2.2	7/01	15450	15794			101	344	546	Y
	2.2	7/12	15377	15777			118	400	636	Y
	2.3	27/10	13341	15677			218	2,336	2,772	Y
	2.3	8/01	15412	15778	В		117	366	600	Y
	2.3	8/03	15139	15799			96	660	852	Y
	2.3	8/07	14684	15792			103	1,108	1,314	Y
	3.1	20/05	14334	15800			95	1,466	1,656	Y
	3.1	9/02	15560	15792	D		103	232	438	Y
	3.1	9/05	15557	15759			136	202	474	Y
	3.1	9/11	15412	15778	В		117	366	600	Y
	3.2	10/05	15412	15778	В		117	366	600	Y
	3.2	10/09	15412	15802			93	390	576	Y
	3.3	11/01	15560	15786	С		109	226	444	Y
	3.3	21/03	14459	15735			160	1,276	1,596	Y
	3.4	12/02	15560	15750	А		145	190	480	Y
	3.4	12/05	15412	15778	В		117	366	600	Y
	3.4	22/01	13563	15749			146	2,186	2,478	Y
	3.4	22/04	14478	15655		+5	240	1,177	1,662	Y
vac-C ^{KO} (GFP)	UCSB (14)	220	15507	15797			98	290	486	Y

^a Nucleotide positions of breakpoints and reinitiation sites as aligned to the reference sequence for the Moraten vaccine virus (GenBank accession number AF266287).

^b The letters A to D indicate clone classes when identical DI-RNAs were isolated from independent plaques.

^c The rule of six indicates whether the calculated total length of the DI-RNA is a multiple of six (yes [Y]) or not (no [N]).

RNAs: 44 sequences were obtained, and 4 of these were derived from parental vac2 (top four lines) and 40 were derived from mutant vac2-C^{KO}. Table 2 summarizes the data for IC323 virusderived DI-RNAs: 21 sequences were obtained, and 3 of these were derived from parental IC323 (top three lines) and 18 were derived from mutant IC323-C^{KO}. The breakpoints and reinitiation sites for all DI sequences are summarized in Fig. 2B for vac2 and in Fig. 2C for IC323 viruses. Breakpoints were distributed broadly (Fig. 2B and C). Their frequency of detection depended on the location of the primer used for amplification (primer A2, A3, or A4) and was inversely proportional to the distance from the 5' end of the genome (Fig. 2B and C; compare the top, middle, and bottom panels). The average breakpoint position for vac2-derived DI-RNAs was nucleotide 14,843, with a standard deviation of 831 nucleotides. For IC323-derived DI-RNAs, the average breakpoint was nt 15,122 \pm 681. In contrast, deduced reinitiation sites for

TABLE 2	Wild-type	virus-derived	DI-RNAs
INDLL 2	whu-type	vii us-ucrivcu	D1-1(11/10

		DI clone	Nucleotide position ^a			Length (no. of nucleotides)				
Virus	Expt no. and plaque no.		Breakpoint	Reinitiation site	Clone class ^b	Insertions/ deletions	dsRNA stem	ssRNA loop	DI-RNA length	Rule of six ^c
IC323	2.1	29/03	15595	15793	Е		102	198	402	Y
	2.3	55/04	15282	15788			107	506	720	Y
	3.2	30/06	15600	15794			101	194	396	Y
IC323-C ^{KO}	1.1	57/03	15595	15793	Е		102	198	402	Y
	1.1	64/01	14271	15659			236	1,388	1,860	Y
	1.2	33/04	15595	15793	Е		102	198	402	Y
	1.2	58/01	15545	15759			136	214	486	Y
	1.2	$58/04^{d}$	15545	15759			136	214	486	Y
	1.3	34/09	15562	15688			207	126	540	Y
	1.3	59/05	14960	15773			122	813	1,057	Ν
	1.3	68/01	13677	15779			116	2,102	2,334	Y
	2.2	60/01	15608	15798			97	190	384	Y
	2.2	60/06 ^e	15608	15798			97	190	384	Y
	2.2	70/01	13664	15624			271	1,960	2,502	Y
	2.3	61/01	15190	15742			153	552	858	Y
	2.3	61/06	15385	15703			192	318	702	Y
	2.3	65/03	13850	15792			103	1,942	2,148	Y
	3.1	66/10	14624	15784		+4	111	1,160	1,386	Y
	3.2	63/03	15244	15730			165	486	816	Y
	3.3	40/01	15560	15786			109	226	444	Y
	3.4	41/02	15595	15793	Е		102	198	402	Y

^{*a*} Nucleotide positions of breakpoints and reinitiation sites as aligned to the reference sequence for the IC-B wild-type virus from which IC323 is derived (GenBank accession number AB016162).

^b The letter E indicates the clone class when identical DI-RNAs were isolated from independent plaques.

^c The rule of six indicates whether the calculated total length of the DI-RNA is a multiple of six (yes [Y]) or not (no [N]).

^d This clone differed from 58/01 of the same rescue by a biased hypermutation event.

 e This clone differed from 60/01 of the same rescue by a biased hypermutation event.

most DI sequences clustered in a narrow region within the last 200 nucleotides of the MV genome (for vac2, nt 15,758 \pm 43; for IC323, nt 15,758 \pm 50) close to the region containing the sequence of primer A1 used for amplification. The total length of the DI-RNAs deduced from the positions of breakpoints and reinitiation sites (Tables 1 and 2, DI-RNA length columns) ranged from about 400 nt to about 3.5 kb. Interestingly, a few identical DI sequences appeared in virus plaques generated from independent rescue experiments. Identical DI-RNAs were assigned to the same class, labeled with the letters A to E (Tables 1 and 2, Clone class columns).

We also assessed whether the DI-RNAs conformed to the rule of six, which stipulates that only paramyxovirus genomes with hexameric lengths are efficiently replicated (10, 32). Indeed, the calculated total DI-RNA length was a multiple of six for 63 out of 65 clones. However, DI-RNA clones 24/01 (vac2- C^{KO}) and 59/05 (IC323- C^{KO}) did not comply with this rule. Interestingly, 6 out of 65 DI-RNAs possessed internal deletions or insertions (Tables 1 and 2, Insertions/deletions columns) that made them compliant with the rule of six (Tables 1 and 2, Rule of six columns). Assuming that DI-RNA generation is a purely random process, about 83% (5 out of 6) of newly generated DI-RNAs are predicted to be noncompliant with the rule of six, but because we found that only about 3% of the DI-RNAs were noncompliant, we conclude that DI-RNAs of hexameric length had a replicative advantage.

Clusters of A-to-G and U-to-C transitions in DI-RNA sequences. DI-RNAs of negative-strand RNA viruses (19, 33) and even full-length MV genomes from diseased human brains (34) can accumulate clusters of A-to-G or U-to-C transitions, characteristic of the action of the cellular ADAR1 deaminase (35, 36). We detected clusters of A-to-G or U-to-C transitions (biased hypermutation events) in 9 DI-RNA sequences (Fig. 3A and Table 3). The percentage of nucleotides affected by these transitions varied from about 4 to 42% (average 21%). Notably, all identified clones exhibited only one type of transition, and we did not observe a switch in the type of transition at the breakpoint. Figure 3A illustrates one biased hypermutation event: 27 transitions to C among 70 U residues (39%) within the sequence prior to the breakpoint and then an additional 9 transitions to C among 41 U residues (22%) after the reinitiation site. Interestingly, the DI-RNA carried an internal deletion in the copy-back sequence that spanned 107 nt between positions 15,653 and 15,759 (Fig. 3A, dashes). The internal deletion generated a DI-RNA compliant with the rule of six.

ADAR1 is a cellular enzyme that catalyzes the deamination of adenosines in dsRNA to produce inosine, which is recognized as guanosine (35, 36). The A-to-I editing activity of ADAR1 is affected by the neighboring nucleotides of the A that is deaminated (Fig. 3C) (21). We assessed whether the nucleotides flanking the modification sites in the MV DI-RNAs conformed to the established signature for ADAR1. The neighboring nucleotide frequencies (positions -4 to +4) of all observed hypermutations (Fig. 3B) were in good agreement with the nearest neighbor signature reported by Eggington et al. (21) (Fig. 3C is reconstructed from data available in the supplemental material of the original publication [21]). Most significantly, our data set and that of Eggington et al. (21) show an underrepresentation of C and G residues at the -1



FIG 3 ADAR1-like hypermutations in DI-RNA clones. (A) Alignment of the sequence of DI-RNA clone 06/12 to the vac2 genome (nt 15639 to 15402) and antigenome (nt 15621 to 15887) sequences (GenBank accession number AF266287). The binding sites of the A1 and A2 primers are underlined (top line and bottom two lines, respectively). Identical nucleotides are shown as dots. The breakpoint and reinitiation site nucleotides are indicated. Deleted nucleotides are shown as dashes. (B) Frequency analysis of nucleotides flanking hypermutated sites in DI-RNA clones (all 9 events detected here; Table 3). (C) Frequency of neighboring nucleotides flanking ADAR1 hypermutation sites defined *in vitro* (21).

position and the overrepresentation of U at the -2 position. This suggests that ADAR1 may have acted on the DI-RNAs generated during MV infection.

DI-RNAs are encapsidated and protected by lipid envelopes. To assess whether DI-RNAs are incorporated into viral particles,

TABLE 3 ADAR1-mediated hypermutations in DI-RNA clones

	Expt no. and			No. of hypermutations observed/no. of total	%
Virus	plaque no.	DI clone	Polarity ^a	nucleotides ^b	editing ^c
vac2-CKO	2.1	06/12	$U \rightarrow C$	36/111	31.5
	2.3	08/03	$U \rightarrow C$	6/141	4.3
	3.1	20/05	$U \rightarrow C$	16/126	12.7
IC323-С ^{ко}	1.2	58/04	$U \rightarrow C$	5/52	9.6
	1.3	$59/03^{d}$	$A \rightarrow G$	17/67	25.4
	2.2	60/06	$U \rightarrow C$	8/21	38.1
	3.1	$62/04^{d}$	$U \rightarrow C$	19/54	35.2
	3.1	$62/06^{d}$	$A \rightarrow G$	28/67	41.8
	3.1	$62/11^{d}$	$U \rightarrow C$	11/54	20.4

 a Polarity indicates whether the hypermutations were observed as A \rightarrow G or U \rightarrow C in the genomic orientation.

^b The number of mutations observed/number of total nucleotides of the respective type (A or U) in a cluster.

^c The percentage is the calculated frequency of transitions observed over possible events.

^d These sequences were derived through semispecific hybridization of primer A2 at position – 15670 of the MV genome.

we generated two virus stocks: one without detectable DI-RNA (vac2; we started with plaque 1.2; Fig. 2A) and the other with high DI-RNA levels (vac2- C^{KO} ; we started with plaque 2.1; Fig. 2A). Since >90% of infectious MV remains cell associated, intracellular infectivity was purified using a two-step density gradient protocol (Fig. 4A). The visible bands for vac2 and vac2- C^{KO} particles had similar intensities in both steps, suggesting that the amount of physical particles was similar for the two preparations. However, the infectious titers of the purified virus preparations differed between vac2 and vac2- C^{KO} by about 10-fold (Fig. 4B, right columns), similar to the ratio of the original stocks (Fig. 4B, left columns).

To assess whether the viral RNAs were encapsidated and protected by a lipid envelope, purified virions were treated with different combinations of nuclease (Benzonase), protease (trypsin), and nonionic detergent (Triton X-100). To control for enzymatic activities, the viral envelope glycoprotein H and nucleocapsid protein N were analyzed after treatment (Fig. 4C, top two panels). For both parental and C^{KO} virion preparations, trypsin treatment in the absence of detergent led to the proteolytic cleavage of most H protein (Fig. 4C, lanes 4 and 9), whereas N protein was affected only to a small extent, if at all. However, when virions were treated with Triton X-100 prior to trypsinization, both H and N were completely degraded by trypsin (Fig. 4C, lanes 5 and 10).

We also extracted RNA from the samples, reverse transcribed it, and subjected the cDNA to PCR analysis using DI-RNA-specific primers (Fig. 4C, panel A1/A2) and full-length RNA-specific primers (Fig. 4C, panel A1/B1). Benzonase treatment in the ab-



FIG 4 Analysis of nucleic acids in vac2- and vac2- C^{KO} -infected cells and in purified vac2 and vac2- C^{KO} particles. (A) Two-step gradient purification of MV particles. Virus bands are indicated by brackets. (B) Titers (numbers of TCID₅₀s per milliliter) of virus stocks prior to and after purification. (C) RNA protection assay with purified virions. Samples were treated with the indicated combinations of trypsin, Triton X-100, and Benzonase; immunoblots were against MV H and N proteins; RT-PCR/PCR analysis was performed for DI-RNAs (A1/A2 primers) and full-length RNA (A1/B1 primers). (D) Northern blot analysis of total RNA from about 3×10^5 unified (UI; lane 1) or infected (lanes 2 and 3) Vero cells harvested at 32 h postinfection, as well as RNA extracted from 1×10^6 TCID₅₀s of purified (pur.) and Benzonase-treated vac2 (lane 4) and vac2- C^{KO} (lane 5) particles. The positions of the antigenome and L, H-L, and F-H-L mRNAs (+probe) are indicated on the right side of the left panel. The identities of the genomic RNA and the DI-RNAs (-probe) mapped by PCR amplification are indicated on the right side of the left panel. Lane M, molecular size marker. The numbers on the left side of the left panel indicate the sizes of the marker bands (in bases).

sence of Triton X-100 and trypsin affected both DI-RNA- and full-length RNA levels only to a small extent (Fig. 4C, lanes 2 and 7). Treatment of virions with trypsin only prior to nuclease treatment did not affect the strength of the PCR signals (Fig. 4C, lanes 4 and 9). However, when virions were treated with Triton X-100, the viral RNA became accessible to Benzonase, leading to the complete loss of the PCR signals (Fig. 4C, lanes 3, 5, 8, and 10). We conclude that both DI-RNA and full-length RNA are encapsidated by N and that the majority of these ribonucleocapsids (RNPs) are protected by a lipid envelope. Notably, treatment of the RNPs with trypsin was not required when virions were treated with detergent (Fig. 4C, lanes 3 and 8), indicating that RNPs became nuclease sensitive after Triton X-100 treatment.

High levels of DI-RNA genomes are found in particles. To further characterize the viral RNP, after Benzonase treatment we extracted RNA from 1×10^6 TCID₅₀s of purified virions. As a control, we also extracted total RNA from Vero cells infected with parental and C^{KO} viruses and from uninfected cells. These samples were analyzed by Northern blotting using single-strand RNA probes complementary to the L/trailer region of the antigenome (Fig. 4D, +probe) or genome (Fig. 4D, -probe).

Figure 4D shows an analysis of the antigenomic (left) and genomic (right) RNA from uninfected cells, infected cells, and

purified virus particles. As expected, the 16-kb full-length antigenomic and genomic RNAs were detected in all four virus samples. However, whereas the antigenome bands for vac2 and vac2-C^{KO} in infected cells had similar intensities (Fig. 4D, left; compare lanes 2 and 3), vac2-C^{KO} showed less genomic RNA than vac2 (Fig. 4D, right; compare lanes 2 and 3), correlating with the reduced infectivity recovered from vac2-CKO-infected cells compared to the infectivity recovered from vac2-infected cells. The difference in intensity of the genome band was not detected in purified virion samples (Fig. 4D, right; compare lanes 4 and 5), since we loaded RNA from equal numbers of infectious units of both viruses. However, antigenome ratios were altered in purified virions (Fig. 4D, left; compare lanes 4 and 5), indicating that vac2-C^{KO} had incorporated higher levels of the antigenome into virions than vac2. For total RNA samples from infected cells, the antigenome probe also detected bands for L mRNA (about 7 kb) and two bands that corresponded to polycistronic H-L (9 kb) and F-H-L mRNAs (11 kb) (37). As expected, the L, H-L, and F-H-L mRNAs were not detected in the purified viruses.

Using either the antigenome probe or the genome probe, we observed multiple RNA species with a smaller molecular size that were present only in vac2- C^{KO} samples. The sizes of these bands ranged from 0.1 kb to about 7 kb, with most being in the 0.3- to



FIG 5 DI-RNA and dsRNA generation by C^{KO} viruses expressing either GFP or mCherryNLS. (A) Western blot and PCR analysis of P1 and P2 of fluorescent reporter-expressing vac2 and vac2-C^{KO}. The antisera used for detection are indicated on the left of the top four panels. Primer pairs used for amplification are indicated on the left of the bottom two panels. UI, uninfected cells. (B) Confocal microscope analysis of HeLa cells infected with fluorescente reporter-expressing vac2 and vac2-C^{KO}. (Top) Autofluorescence of GFP (green); (bottom) autofluorescence of mCherryNLS (red); (all panels) immunofluorescence staining for dsRNA (yellow) and MV N protein (blue). Magnifications, ×40.

2-kb range. The sizes of some RNAs were similar to those of specific DI-RNAs of vac2-C^{KO} plaque 1 from experiment 2 (referred to as 2.1), as identified by the PCR cloning approach (Table 1; compare the DI-RNA clone identifiers with those marked in Fig. 4D). Similarly sized RNAs were also detected, but in greater abundance, in purified vac2-C^{KO} virions (Fig. 4D, right, lane 5). We detected more RNA species with the genome probe than with the antigenome probe, consistent with the notion that these RNAs may be truncated genomes. We conclude that DI-RNAs are incorporated into intracellular particles since they are protected by a lipid envelope. The 10-fold reduced infectivity of vac2-C^{KO} correlates with the enhanced incorporation of DI-RNA genomes into intracellular particles.

 C^{KO} viruses generate dsRNA very early. Since DI-RNAs are most efficiently amplified at a high MOI, we sought to limit cellcell fusion, a process resulting in the bulk transfer of genomes. We therefore carried out infections on Vero cells, rather than on the more fusogenic Vero-hSLAM cells. Because we needed viruses expressing an alternative fluorescent reporter protein for subsequent competition experiments, we generated vac2-based variants expressing mCherry (mCherryNLS [28]). Figure 5A shows protein expression and DI-RNA levels after two passages of the vac2based viruses expressing either GFP or mCherryNLS. As expected, passage 1 DI-RNA amplification in Vero cells was minimally detectable, if it was detectable at all (lanes P1, A1/A2 panel). However, significant amounts of DI-RNA-specific PCR products were detected for both mutant C^{KO} viruses at passage 2 but not for parental vac2 (lanes P2, A1/A2 panel).

Having previously documented the kinetics of dsRNA production of standard and C^{KO} viruses in HeLa cells (14), we infected these cells and performed immunofluorescence staining for viral N protein and dsRNA (Fig. 5B). We found strong dsRNA staining (Fig. 5B, yellow, seen as white when superimposed onto blue) colocalizing with viral replication sites (revealed by N protein; Fig. 5B, blue signal) in passages 1 and 2 of either C^{KO} virus. In contrast, the C-expressing parental vac2 of either passage did not elicit detectable dsRNA staining at viral replication sites. Thus, immunofluorescence staining detects dsRNA before DI-RNA can be detected by PCR amplification.

The C^{KO} phenotype is dominant. To assess whether C-protein expression in *trans* can limit DI-RNA amplification, we coinfected cells with virus stocks possessing either high or undetectable DI-RNA levels. These viruses expressed different fluorescent proteins (either GFP or mCherryNLS) for easy replication tracking. We infected interferon response-defective Granta-519 cells with each virus at an equivalent MOI and quantified the GFP and mCherry expression at 48 h postinoculation by fluorescence-activated cell sorter analysis (Fig. 6).

Figure 6A shows the levels of GFP expression in cells left uninfected (panel 1) or infected only with either virus expressing GFP [vac2(GFP) in panel 2; vac2- C^{KO} (GFP) in panel 3] or only with either virus expressing mCherryNLS [vac2 (mCherryNLS) in panel 4; vac2- C^{KO} (mCherryNLS) in panel 7]. Panels 5, 6, 8, and 9 document the levels of GFP expression in cells coinfected with two viruses, as indicated on the top of each column and on the left of each row. The median fluorescence intensity (MFI) of vac2(GFP) (panel 2) was about 4,500, or 110 times over the background MFI of 41 (panel 1); the MFI of vac2- C^{KO} (GFP) (panel 3) was about 70 times over the background MFI. Infection with mCherryNLS-expressing viruses was not detected in the GFP channel (panels 4 and 7).

We then measured GFP expression levels in cells coinfected with 1:1 mixtures of viruses that were either DI-RNA poor (vac2derived stocks) or DI-RNA rich (vac2-C^{KO}-derived stocks). We



FIG 6 Dominance of the C^{KO} phenotype. (A) GFP fluorescence intensities of cells coinfected with viruses indicated on the top and on the left. UI, uninfected cells. Vertical axis, cell counts (linear scale); horizontal axis, GFP fluorescence intensity (FI; \log_{10} scale). (B) Overlay of graphs from panel A. (Left) Summary of coinfections with vac2(GFP); (right) summary of coinfections with vac2(GFP); Gray area, uninfected cells (panel 1); light green area, single GFP-virus infection (panel 2 or 3); dotted green line, GFP-virus coinfected with vac2(mCherryNLS) (panel 5 or 6); solid green line, GFP-virus coinfected with vac2-C^{KO}(mCherryNLS) (panel 8 or 9).

undertook these mixed infections to assess whether DI-RNA-rich stocks interfere with the replication of DI-RNA-poor stocks (dominant negative effects). Alternatively, this experiment might reveal a positive effect of DI-RNA-poor (C-protein-expressing) stocks on the replication of DI-RNA-rich stocks, presumably through *trans*-complementation by the C protein.

Figure 6B illustrates the results of mixed infection analyses. **Figure 6B**, left, shows that addition of a DI-RNA-poor stock [vac2(mCherryNLS)line] to another DI-RNA-poor stock [vac2(GFP)] reduced the GFP MFI about 1.5-fold (compare the light green area with the dark green dotted line). Addition of a DI-RNA-rich stock [vac2-C^{KO}(mCherryNLS)] reduced the GFP MFI of the DI-RNA-poor stock more than 8-fold (compare the light green area with the dark green solid line). In contrast, as shown in **Fig. 6B**, right, addition of a DI-RNA-poor stock [vac2-(mCherryNLS)]ine] to a DI-RNA-rich stock [vac2-C^{KO}(GFP)] reduced the GFP MFP 1.2-fold (compare the light green area with the dark green dotted line), while addition of a DI-RNA-rich stock [vac2-C^{KO}(mCherryNLS)] to another DI-RNA-rich stock had a more pronounced negative effect (a 3.5-fold reduction of the GFP MFI; compare the light green area with the dark green solid line). Because DI-RNA-rich virus stocks in any combination had a negative effect on the growth of the DI-RNA-poor stocks and because DI-RNA-poor stocks which expressed C protein had no positive effect on DI-RNA-rich stocks, we conclude that the C^{KO} effect is dominant.

DISCUSSION

MV C-protein function. Viruses of the *Morbillivirus*, *Henipavirus*, and *Respirovirus* genera of the *Paramyxoviridae* family express one or more C proteins (2). However, different studies have revealed alternative functions of C proteins. In the case of MV and Nipah virus, minigenome assays have suggested that C protein down-regulates genome replication (38, 39). Inhibition of transcription but not of replication was described for Sendai virus (SeV) in *in vitro* assays (40), although SeV C protein also is reported to be essential for the maintenance of negative-strand RNA genomes in virus particles (41) and an SeV-C^{KO} mutant did not undergo a shift from early (antigenome-dominated) to late (genome-dominated) replication (42). The generation and accumulation of DI-RNAs in C-defective paramyxoviruses would account for many of these observations.

In the absence of C protein, production of full-length genomes would be reduced and defective genomes would be incorporated into virions instead. In addition, the dsRNA structures formed by DI-RNAs are known to activate PKR (14), which induces an arrest in protein synthesis via the cellular stress response, thereby delaying viral replication. While in a preceding study we detected DI-RNA in one MV-CKO stock that had been passaged several times at a low multiplicity of infection (14), here we provide evidence for the generation of DI-RNAs as early as passage 1 following virus rescue. Moreover, we observed about 10 times more DI-RNA in $C^{\rm KO}$ virus infection than in parental virus infection, implying a genetic link between the $C^{\rm KO}$ phenotype and DI-RNA generation. Notably, C^{KO} mutants of SeV and human parainfluenza virus type 1 have also been reported to generate double-stranded RNA and activate PKR, but these studies did not address the origin or the structure of these RNAs (43, 44). Although CKO mutants of several paramyxoviruses can be rescued and are only moderately attenuated in cultured cells (16), they are strongly attenuated in vivo (45, 46). Thus, C proteins of diverse paramyxoviruses may share a similar function, to enhance RNA replication processivity.

The frequency but not the quality of DI-RNA generation is determined by the C protein. Most breakpoints of the DI-RNAs are widely distributed along the MV genome, with the frequency of the appearance of a breakpoint at a specific position being inversely proportional to the distance from the 5' end of the genome. In contrast, reinitiation sites are clustered in a narrow region close to the 5' end of the genome. We note, however, that repositioning of the copy-back primer (Fig. 1C, primer A1) to a more internal binding site resulted in the detection of DI-RNAs with more internal reinitiation sites (data not shown).

Our Northern blot analysis showed several RNAs hybridizing with the genome probe (Fig. 4C, right) that did not hybridize with the antigenome probe (Fig. 4C, left) and whose sizes did not correspond to the calculated sizes of any cloned DI-RNA. These species may possibly reflect truncated genomes lacking the copy-back sequence. Indeed, several kinds of defective replication products have been observed for vesicular stomatitis virus (19, 47) and SeV (48, 49).

The C^{KO} phenotype of MV is dominant: a C-protein-defective virus inhibited the replication efficiency of a C-protein-expressing virus in *trans*. On the other hand, a C-protein-expressing virus did not prevent DI-RNA replication once DI-RNA had been generated. Conceivably, the C protein might function together with the polymerase complex in separating the nascent and template RNA hybrid, or, alternatively, C protein might support the rapid cotranscriptional encapsidation of the nascent RNA with N protein. Future studies will discriminate between these possibilities.

ADAR1-mediated editing and the PKR-ADAR balancing model. DI-RNAs of negative-strand RNA viruses are potent triggers of innate immune receptors, including PKR, mda-5, and RIG-I (12-14), thereby inducing the expression of type I IFNs in vitro (50) and in vivo (51). The MV DI-RNAs characterized here, if not cotranscriptionally encapsidated, form perfect hairpins throughout the double-stranded structure. These hairpins are ADAR1 substrates in vitro (23). Destabilization of these hairpins through A-to-I editing could account for the proviral function of ADAR1, which was documented in several viral infections and attributed to suppression of activation of innate immune responses (52, 53). For example, in cell lines deficient in ADAR1 expression, MV strongly induces IFN (54), PKR activation (55), and the cellular stress response (56, 57). Furthermore, higher levels of dsRNA accumulate in ADAR1-deficient cells infected with MV than in cells sufficient in ADAR1 expression (14).

Herein we provide evidence for the enzymatic activity of ADAR1 on MV DI-RNAs. We identified several DI-RNA sequences bearing clusters of A-to-G or U-to-C mutations and show through a computational approach that the sequences flanking the edited nucleotides, as well as the average editing frequency, were characteristic of ADAR1, but not ADAR2 (21). Of the three known ADARs, only the IFN-inducible ADAR1 isoform (also called p150) is cytoplasmic (58, 59) and localizes to stress granules (56). These granules are induced during MV-C^{KO} infection and form in close proximity to viral replication sites (14). In contrast, the constitutively expressed isoforms of ADAR1 (p110) and ADAR2 are exclusively found in the cell nuclei (53). Thus, the subcellular cytoplasmic localization of only ADAR1-p150 is correct for editing of MV RNAs, which are exclusively replicated in the cytoplasm (2).

Notably, in individual DI-RNAs the transitions were of the same type (either A to G or U to C) and occurred on both sides of the breakpoint and reinitiation site. This pattern indicates that the generated DI-RNAs were direct substrates for editing by ADAR1 but not the full-length MV genomes. If the latter were the case, the polarity switch between breakpoint and reinitiation site would have also caused the ADAR1 transitions to change within the DI-RNA molecule. Thus, our findings suggest that ADAR1 editing destabilizes unencapsidated DI-RNAs may be the double-stranded RNA that elicits innate immune responses in negative-strand RNA virus infections. ADAR1 A-to-I editing has also been documented in full-length RNA virus genomes (33, 34, 60, 61) and was recently inferred by deep sequencing analyses of several paramyxoviruses (62, 63).

DI-RNAs as a frequent by-product of wild-type and vaccinelineage infections. DI-RNAs were originally detected in laboratory strains propagated at a high multiplicity (9) and later also in commercial vaccines (64, 65). Here we establish that DI-RNAs can accumulate in the earliest passage available for analysis following the initial virus rescue. Moreover, we documented DI-RNA generation in a wild-type MV strain, thereby providing evidence that DI-RNAs are not restricted to vaccine-lineage viruses and have the potential to influence virulence.

DI-RNAs may alter the characteristics of MV live-attenuated vaccines and modulate the efficacy of recombinant MV engineered for vaccination against heterologous pathogens or used as oncolytic vectors (27, 66–68). While vaccine producers seek to avoid DI-RNAs in order to standardize efficacy, oncolytic potency may not rely uniquely on efficient viral replication and cell lysis but may also rely on immunostimulatory properties (69, 70). The balance between replication efficiency and immune activation of an oncolytic vector may be an important key to a successful therapy, and immunostimulatory DI-RNAs might be a desirable feature of certain oncolytic vectors. Further studies of the MV C protein and its function may lead to the design of viruses that strike an optimal balance between replication efficiency and immune activation.

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