APOBEC3-mediated restriction of RNA virus replication

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APOBEC3 family members are cytidine deaminases with roles in intrinsic responses to infection by retroviruses and retrotransposons, and in the control of other DNA viruses, such as herpesviruses, paroviruses and hepatitis B virus. Although effects of APOBEC3 members on viral DNA have been demonstrated, it is not known whether they edit RNA genomes through cytidine deamination. Here, we investigated APOBEC3-mediated restriction of Coronaviridae. In experiments in vitro, three human APOBEC3 proteins (A3C, A3F and A3H) inhibited HCoV-NL63 infection and limited production of progeny virus, but did not cause hypermutation of the coronaviral genome. APOBEC3-mediated restriction was partially dependent on enzyme activity, and was reduced by the use of enzymatically inactive APOBEC3. Moreover, APOBEC3 proteins bound to the coronaviral nucleoprotein, and this interaction also affected viral replication. Although the precise molecular mechanism of deaminase-dependent inhibition of coronavirus replication remains elusive, our results further our understanding of APOBEC3-mediated restriction of RNA virus infections.

Apolipoprotein B mRNA-editing enzyme, catalytic polypeptide-like type 3 (APOBEC3) proteins are Zn2+-dependent cytidine deaminases that belong to the APOBEC superfamily. APOBEC3 proteins are considered to be part of the intrinsic defense system in eukaryotic cells, and can inhibit replication of some viruses and restrict mobile genetic elements. All APOBEC3 proteins share a cytidine-deaminase domain, which is identifiable by the primary amino acid motif His-Xaa-Glu-Xaa23–28-Pro-Cys-Xaa 2–4-Cys, in which Zn2+ is coordinated. Deamination of cytidine results in its conversion to uridine. During this process, the ammonium group on C4 of cytidine undergoes nucleophilic attack by an activated water molecule and the APOBEC3 glutamate residue.

A single copy of a gene encoding APOBEC3 is found in the genomes of rodents, cats, pigs and sheeps, with two genes in the genomes of cows, three the genomes of dogs and horses and seven genes in the human genome, encoding A3A, A3B, A3C, A3D, A3F, A3G and A3H1. The mechanisms and selective pressures that led to the APOBEC3 gene expansion in humans are unclear, but it is believed that all the genes originated from duplication of a single-copy primordial APOBEC3 gene. This multiplication may have provided protection from genomic instability caused by human endogenous retroviruses and retrotransposons2.

The best-studied protein within the human APOBEC3 family is A3G. Its function as an antiviral factor was discovered in 2002 through cDNA-transfer experiments that were designed to identify a cellular suppressor of an HIV-1 accessory protein, the virion infectivity factor (Vif). The primary antiretroviral activity of A3G requires its encapsidation into HIV-1 particles, and involves cytidine deamination of the reverse-transcribed first viral DNA strand, with mutations becoming fixed as G to A changes upon second-strand synthesis. Mutational frequencies can exceed 10% of all G residues (hypermutation), and may result in inhibition of viral replication. This activity is blocked by Vif protein, which specifically binds to and ubiquitinates A3G, directing it for degradation in the...
proteasome. Other APOBEC3 proteins have been found to affect a wide variety of viruses, including other retroviruses (e.g., simian immunodeficiency virus (SIV), equine infectious anemia virus (EIAV), murine leukemia virus (MLV), foamy viruses), and DNA viruses (e.g., hepatitis B virus, parovirus)13–15. As well as the inhibition of viral replication by hypermutation, APOBEC3-induced mutations also cause degradation of viral genomic material, mediated by cellular DNA-repair mechanisms16.

The RNA-editing capability of APOBEC3 proteins was identified in 200416, since when extensive mutational analysis of A3A mRNA transcripts in monocytes and macrophages has been demonstrated17. Our current understanding of the APOBEC3 proteins suggests that they may also constitute an innate defense system against RNA viruses. For example, A3G known to be involved in restriction of three RNA pathogens (the mumps, measles and respiratory syncytial viruses), but whether this effect is associated with cytidine deamination is not yet known18.

Members of the family Coronaviridae are positive-strand RNA viruses with large genomes, ranging in size from 27 to 32 kb. Six human coronaviruses (HCoVs) have been identified, four of which (HCoV-OC43, HCoV-229E, HCoV-NL63 and HCoV-HKU1) circulate continuously in human populations. These four viruses cause the common cold in otherwise healthy adults, and may cause more severe symptoms in young, elderly and immunocompromised individuals19–22. Other coronaviruses that infect humans can cause severe and life-threatening diseases. In 2002, severe acute respiratory syndrome coronavirus (SARS-CoV) emerged and affected ~8,000 individuals, causing ~800 deaths23,24. Middle East respiratory syndrome coronavirus (MERS-CoV) is the most recently identified coronavirus that infects humans. MERS-CoV was isolated in 2012 and is associated with severe pneumonia and renal failure, with a mortality rate of ~30%25,26.

HCoV-NL63 was identified in 2004 and is distributed worldwide, with highest prevalence during winter and early spring in temperate climate27. HCoV-NL63 accounts for a considerable number of hospitalisations among children <18 years of age, the elderly and immunocompromised individuals19. A notable characteristics of coronaviruses is that their genomes are highly U/A-rich and C/G-poor. For example, HCoV-NL63 has 39% U and 27% A content, with only 14% C and 20%, G nucleotides28. The explanations for this bias is not yet known28,29. One possibility is that, over an evolutionary timescale, genomes of coronaviruses might have been shaped by cytidine deamination. Our aim, therefore, was to determine whether APOBEC3 proteins might be responsible for this modification. Our results demonstrate that three of the APOBEC3 proteins (A3C, A3F and A3H) can inhibit coronaviral infection in vitro.

Results

APOBEC3-family gene expression is upregulated in HCoV-NL63-infected cells. Modulation of human APOBEC3-family transcript levels in human airway epithelium (HAE) cultures during coronavirus infection was evaluated by quantitative RT-PCR (qRT-PCR) with primers and standards developed in our lab. The level of each APOBEC3 mRNA was analysed in fully differentiated cells from healthy adults, and a high level of expression was observed for six out of seven human APOBEC3 genes (Fig. 1a). The same six genes (APOBEC3A, APOBEC3C, APOBEC3D, APOBEC3F, APOBEC3G and APOBEC3H) were expressed during HCoV-NL63 infection, and these genes were analysed in subsequent experiments

HAE cultures were infected with HCoV-NL63 and cultured for 5 days at 32 °C. As a control, HAE cultures were infected with influenza A H1N1 virus, which induces considerable upregulation of APOBEC29. RT-qPCR analysis of total RNA extracted from infected cells revealed upregulation of APOBEC3A, APOBEC3C, APOBEC3D and APOBEC3F transcripts in HCoV-NL63-infected cells relative to mock-infected cells, with an almost 100-fold increase in APOBEC3A expression (Fig. 1b). As anticipated, APOBEC3G was upregulated in the influenza-infected cells, and APOBEC3H expression also increased (Fig. 1c).

Expression of APOBEC3 proteins in cultured cells. Plasmids encoding individual APOBEC3 proteins were prepared and transfected into 293T cells, which were cultured and analysed for protein expression by western blotting (Fig. 2a, Supplementary Figure 1). For expression in the naturally permissive LLC-Mk2 cell line, in vitro transcription. Transfection of LLC-Mk2 cells was efficient, reaching ~80% for the control GFP mRNA (Fig. 2b). Expression of HA-tagged APOBEC3 proteins in mRNA transfected LLC-Mk2 cells, measured by flow cytometry, was similar for all proteins (Fig. 2c).

Human APOBEC3 proteins A3C, A3F and A3H inhibit HCoV-NL63 replication. mRNA-transfected LLC-Mk2 cells were infected with HCoV-NL63 and cultured for 5 days at 32 °C. No cytopathic effect (CPE) was observed in cells expressing A3C, A3F and A3H proteins, but CPE was observed in untransfected or GFP-transfected cells, and in cells expressing A3A, A3DE and A3G (Fig. 3a). Analysis of HCoV-NL63 replication showed a 1.5–2 log reduction in virus yield in cells expressing A3C, A3F and A3H, relative to the control sample. By contrast, HCoV-NL63 replication was not inhibited in cells expressing GFP, A3A, A3D or A3G (Fig. 3b).

APOBEC3-mediated HCoV-NL63 restriction is associated with cytidine deamination. Because HCoV-NL63 infection resulted in upregulation of expression of A3A, A3C, A3D and A3F and A3C, A3F and A3H all inhibited HCoV-NL63 replication, we tested whether this inhibition was the result of the catalytic activity of the APOBECs. Plasmids encoding variants of A3C, A3F and A3H with Glu → Gln substitutions in the catalytic site were prepared and mRNAs encoding active or inactive proteins were transfected into LLC-Mk2 cells, which were infected with HCoV-NL63 and cultured prior to visualisation of viral proteins with specific antibodies. The percentages of HCoV-NL63-infected cells were measured by flow cytometry. Wild-type APOBEC3 proteins resulted in pronounced reductions in HCoV-NL63 infection rates, whereas catalytically inactive proteins produced moderate inhibition, with infection rates that were not significantly different from those in controls (Fig. 4a). Western blot analysis showed an even expression of all proteins, showing that the inhibitory effect
did not result from unequal mRNA translation (Fig. 4b, Supplementary Figure 2). This level of inhibition of HCoV-NL63 replication by catalytically inactive proteins required transfection of 3 µg of mRNA, whereas for active proteins, inhibition was achieved with transfection of 1 µg of mRNA.

Next, we tested the activity of APOBEC3 proteins in vitro. A DNA template containing several possible sites for cytidine deamination (GC-rich domains) was used for in vitro transcription, producing RNA that was incubated with lysates from 293T cells overexpressing A3C protein or GFP. The RNA was then isolated, reverse transcribed, amplified by PCR and sequenced, revealing several C → C/T changes following incubation with A3C (but not GFP) cell lysate (Fig. 4c). However, hypermutation did not occur, and no pattern was identified in the nucleotide changes.

To examine the progressive accumulation of mutations, we performed a series of virus passages on cells expressing A3C, A3F or A3H. Notably, after the first passage, the inhibitory effect of the three APOBECs diminished, and no decline in virus yield was seen (data not shown). Cell-culture supernatants were collected after four passages of HCoV-NL63, and several regions of the viral RNA genome were sequenced. Point mutations (G → A or C → T) were identified in the genes encoding the 1ab polyprotein and spike protein in virus passed

Figure 1. APOBEC3 gene expression in human airway epithelium (HAE) cell culture. APOBEC3 gene expression was analysed in 3D HAE cultures. mRNA levels were evaluated by quantitative RT-PCR. Expression was assessed (a) in uninfected HAE cultures, (b) in cultures infected with human coronavirus (HCoV)-NL63 and (c) in cultures infected with influenza A virus (IVA) strain H3N2. Results are presented as mRNA copy number per ml. Infection was performed at 400 TCID₅₀ per ml, for 4 days at 32 °C (for HCoV-NL63) or for 2 days at 37 °C (for IVA). Each experiment was performed at least three times with clinical material from different donors, and with two technical replicates. For comparisons by Student’s t-test, *Indicates P < 0.05; **Indicates P < 0.005.
in APOBEC3-expressing cells, but not in wild-type cells. These mutations were not specific to virus collected from cells expressing A3C, A3F or A3H, but were also identified in virus passaged in cells expressing A3A, ADE or A3G.

To identify whether APOBEC3 protein were capable of inducing hypermutation, we used a recombinant GC-rich HCoV-229E virus (ic229E-GC), into which we had introduced silent mutations (either A→G or T→C) in the A/T-rich region of nsp6, resulting in a potential ‘hot-spot’ for cytidine deamination. The virus was propagated and titrated to determine the 50% tissue-culture infective dose (TCID<sub>50</sub>) in naturally permissive Huh-7 cells. The effectiveness of mRNA transfection in Huh-7 cells was confirmed by detection of green fluorescence in cells transfected with GFP mRNA. Huh-7 cells were then transfected with mRNA encoding an APOBEC3 protein, infected with ic229E-GC at 400 TCID<sub>50</sub> per ml and cultured for 5 days. Pronounced inhibition of viral infection occurred in cells expressing A3C, A3F or A3H (data not shown). After four passages of ic229E-GC, viral RNA was sequenced. No G→A or C→T mutations were identified within the GC-rich region in any of the samples. However, ic229E-GC passaged in cells expressing A3F protein showed considerable sequence heterogeneity, compared with passaged in GFP-positive cells, indicating a possible A3F deaminase activity (Fig. 4d).

**APOBEC3 proteins interact with the HCoV-NL63 nucleoprotein.** Our results demonstrated that, although cytidine deamination was involved in inhibition of coronavirus by APOBEC3 proteins, deaminase mutants retained some anti-coronaviral activity, suggesting that an additional mechanism may be involved in APOBEC3-mediated coronavirus restriction. We therefore analysed the interactions between APOBEC3 proteins (A3C, A3F and A3H) and HCoV-NL63 nucleocapsid protein (HCoV-NL63-N), which is the most abundant viral structural protein. Human 293T cells were transiently transfected with plasmid DNA constructs for expression of individual HA-tagged APOBEC3 proteins (or their catalytically inactive derivatives) or GFP. Cell lysates containing the expressed proteins were incubated with purified HCoV-NL63-N, and protein complexes were co-immunoprecipitated with anti-HA-tag resin. Western-blot analysis demonstrated co-immunoprecipitation of HCoV-NL63-N with A3C, A3F and A3H, whereas no HCoV-NL63-N could be detected in samples incubated with GFP lysate. Notably, the catalytically inactive derivatives of A3C, A3F and A3H also bound to HCoV-NL63-N protein (Fig. 5a, Supplementary Figure 3). On the other hand, we could not observe the same effect with other APOBEC3 proteins (A3A, A3D and A3G).
Co-localisation of A3C with HCoV-NL63-N was investigated by co-transfection of LLC-Mk2 cells with mRNAs encoding A3C and HCoV-NL63-N, followed by immunofluorescence staining and confocal microscopy. The analysis showed several clearly visible foci of co-localisation (Fig. 5b). Calculation of Manders’ coefficient gave a value of 0.459, indicating the existence of an interaction between the two proteins.

**Discussion**

APOBEC3-mediated cytidine deamination has been described as an interim antiviral mechanism. Although APOBEC3 proteins were initially shown to primarily modify DNA templates, it has subsequently been demonstrated that they also use RNA as a substrate. The finding that APOBEC3 deaminases can mutate the HIV RNA genome first expanded their possible specificity. Thereafter, it was shown that A3G impairs the replication of mumps, measles and respiratory syncytial viruses. This antiviral activity is not an effect of APOBEC3...
Figure 4. Cytidine deamination is relevant for HCoV-NL63 inhibition. (a) LLC-Mk2 cells were transfected with capped and polyadenylated mRNA encoding active or catalytically inactive (m) APOBEC3 proteins. Following incubation for 24 h at 37 °C in an atmosphere containing 5% CO₂, cells were infected with human coronavirus (HCoV)-NL63 at 400 TCID₅₀ per ml. After 96 h incubation at 32 °C in an atmosphere containing 5% CO₂, cells were fixed and immunostained for HCoV-NL63 nucleoprotein (N) using specific antibodies. The rate of viral infection was analysed by flow cytometry and is presented as the percentage of cells that were infected. For comparisons by Student’s t-test, *Indicates $P < 0.05$. (b) LLC-Mk2 cells were transfected with capped and polyadenylated mRNA encoding active or catalytically inactive (m) APOBEC3 proteins. Following incubation for 24 h at 37 °C in an atmosphere containing 5% CO₂, cells were lysed in RIPA buffer. APOBEC3 proteins and control GAPDH protein were detected by western blotting using specific antibodies. (c) 293T cell lysates enriched in the APOBEC3 protein A3C (lower panel) or green fluorescent protein (GFP) (upper panel) were incubated with in vitro-prepared RNA containing a GC-rich sequence for 2 h at 37 °C. The RNA was then reverse transcribed, amplified by PCR and sequenced. Experiments were performed at least three times, with two technical replicates. (d) Huh-7 cells were transfected with capped and polyadenylated mRNA encoding active A3F (lower panel) or GFP (upper panel). Following 24 h incubation at 37 °C in an atmosphere containing 5% CO₂, cells were infected with the recombinant GC-rich HCoV-229E virus (ic229E-GC) at 400 TCID₅₀ per ml. After four cycles of virus passage in A3F-positive or GFP-positive cells, viral RNA was isolated, reverse transcribed and sequenced across the GC-rich domain.
enzymatic activity, but is thought to result from the interaction of A3G with viral transcripts. A3A has been shown to act as a mRNA-editing factor in human monocytes and monocyte-derived macrophages. However, APOBEC3-mediated deamination of viral RNA genomes has not previously been demonstrated for viruses that do not replicate through DNA intermediates. Here, we investigated the antiviral effects of APOBEC3 proteins on human coronaviruses. HCoV-NL63 was selected as the model pathogen because of its prominent bias in nucleotide composition (low GC content), suggesting the possible shaping of the genome by cytidine deamination.

We studied the interaction between HCoV and HAE cultures, as an in vitro model of the human airways, the primary entry site for human respiratory viruses. mRNA levels of genes encoding A3A, A3C and A3D were upregulated in HCoV-NL63-infected cultures. Upregulation of APOBEC3 protein expression was previously shown to be dependent on interferon alpha, and it is associated with non-cytolytic clearance of human hepatitis B virus in primary hepatocytes. Cytoplasmic DNA triggers production of type I interferon, leading to massive upregulation of APOBEC3A. Here, we investigated the effects of different APOBEC3 proteins, and found that expression of A3C, A3F or A3H limits coronavirus infection. Notably, some infectious progeny were produced in the presence of these proteins, and serial passaging of virus was possible. Surprisingly, if cells overexpressing A3C, A3F or A3H were infected with the progeny virions, no restriction of virus replication occurred. Possible explanations for this effect are that the first viral passage results in selection of viral subspecies that are not affected by APOBEC3 activities, or that viral passage activates a viral factor or mechanism that can counteract APOBEC3 activities.

Serial passage of HCoV in APOBEC3-expressing cells did not result in hypermutation in progeny viruses. It is possible that HCoV-NL63 has reached a maximum level of AT content, so that new C → T or G → A changes in the genome result in a drastic drop in viral infectivity. However, incubation of an RNA transcript containing potential deaminase hot-spots with A3C-enriched cell lysates did not result in hypermutation. We were not able to develop a HCoV-NL63 molecular clone containing GC-rich elements, most likely because of low stability of the reverse genetics system for this virus (data not shown). We therefore developed a HCoV-229E clone containing a GC-rich region. Replication of this virus was inhibited in cells expressing A3C, A3F or A3H, but no mutations were observed in the modified region. The lack of observed hypermutation may result from proofreading and repair of the coronavirus RNA genome, and suggests that inhibition of coronaviral replication by APOBEC3 proteins does not result from deamination of viral RNA.

Figure 5. APOBEC3 proteins interact with human coronavirus (HCoV)-NL63 nucleoprotein (N). (a) 293T cell lysates enriched in active APOBEC3 proteins (A3C, A3F or A3H) or their catalytically inactive derivatives (mA3C, mA3F or mA3H) were incubated with purified HCoV-NL63 N (produced in Escherichia coli) for 1 h at 37 °C. Subsequently, agarose spheres coated with anti-HA antibodies were added, and proteins were co-precipitated using spin columns. Samples were separated by electrophoresis and transferred onto PVDF membrane, and HCoV-NL63 N and HA-tagged APOBEC3 proteins were detected with specific antibodies. PC = positive control protein. (b) LLC-Mk2 cells were transfected with capped and polyadenylated mRNA encoding HCoV-NL63 N (Ctrl NL63 N), A3C (Ctrl A3C) or both transcripts (NL63 N + A3C), and incubated for 72 h at 37 °C. Cells were then fixed and immunostained for A3C (red), HCoV-NL63 N (green) and nuclei (blue). Co-localisation of A3C with HCoV-NL63-N was determined by confocal microscopy and marked with arrows. M, Manders’ coefficient (A3C overlapping with the virus). A representative image is shown.
It was previously shown that, for some RNA viruses, APOBEC-mediated inhibition is deamination-independent. In our system, catalytically inactive A3C, A3F and A3H proteins only had marginal inhibitory effect on virus replication. We suggest that APOBEC3 proteins may affect coronavirus replication by two mechanisms: first, by deamination of yet unidentified cellular targets or deamination of single sites in the coronavirus genome; and second, by direct, non-catalytic interaction with viral RNA and/or proteins. APOBEC proteins may interact with the coronaviral N protein, which is an essential structural protein for formation of the viral ribonucleocapsid, and which also has a role in virus replication. Our results demonstrated that all APOBEC3 proteins with inhibitory activity bound to the N protein. The direct, editing-independent mechanism by which APOBEC3 proteins inhibit HCoV-NL63 replication is still to be elucidated, but it is possible that interaction with the N protein interferes with virus replication and progeny production.

In conclusion, our results show that APOBEC3 proteins (A3C, A3F and A3H) inhibit coronavirus replication. The molecular mechanism of deaminase-dependent inhibition of coronavirus replication remains elusive and may depend on deamination of single sites in the coronaviral genome, or in a cellular target or targets. Passage of HCoV-NL63 in cells expressing APOBEC3 proteins results in virus that is resistant to APOBEC-mediated inhibition, possibly indicating the presence of an unidentified factor that counteracts APOBEC3 activity. The deaminase-independent component of viral inhibition may result from interaction between viral N protein and APOBEC3 proteins.

**Methods**

**Cell culture.** LLC-Mk2 cells (*Macaca mulatta* epithelial kidney cells; ATCC; CCL-7) were cultured in H/E MEM (Hanks’ MEM: Earle’s MEM, 2:1 ratio; Thermo Fisher Scientific, Poland) with 3% fetal bovine serum (heat-inactivated; Thermo Fisher Scientific, Poland) and antibiotics: penicillin (100 U/ml), streptomycin (100 μg/ml), and ciprofloxacin (5 μg/ml). Human hepatocellular carcinoma HuH-7 (kindly provided by dr Lia van der Hoek, UvA, The Netherlands) and 293T (ATCC: CRL-3216; kidney epithelial) cell lines were cultured in Dulbecco’s MEM (Thermo Fisher Scientific, Poland) with 10% fetal bovine serum (heat-inactivated; Thermo Fisher Scientific, Poland) and antibiotics: penicillin (100 U/ml), streptomycin (100 μg/ml), and ciprofloxacin (5 μg/ml). Cells were maintained at 37 °C under 5% CO₂.

**Human airway epithelium (HAE) cultures.** Human epithelial cells were isolated from conductive airways resected from transplant patients. The study was approved by the Bioethical Committee of the Medical University of Silesia in Katowice, Poland (approval no: KNW/0022/KB1/17/10 dated on 16.02.2010). A written informed consent was obtained from all patients. Tissues were procured at the Silesian Center for Heart Diseases, Zabrze, Poland. No organs/tissues were procured from prisoners. Cells were dislodged by protease treatment, and later mechanically detached from the connective tissue. Resulting primary cells were first cultured in selective media to proliferate. Further, cells were trypanospun and transferred onto permeable Transwell insert supports (ϕ = 6.5). Cell differentiation was stimulated by media additives and removal of media from the apical side. In such a manner cells were cultured for 6–8 weeks to form well-differentiated, pseudostratified mucociliary epithelium. All experiments were performed in accordance with relevant guidelines and regulations.

For the virus infection, HAE cultures were washed thrice with 100 μl of 1 × PBS, following inoculation with HCoV-NL63 (isolate Amsterdam 1) or influenza A virus (IVA, strain H1N1) or mock (cell lysate). After 2h incubation at 32 °C (for HCoV-NL63) or 37 °C (for IVA) unbound virions were removed by washing with 100 μl of 1 × PBS and HAE cultures were cultured at an air - liquid interphase until the end of the experiment.

**Virus preparation and titration.** The stock of HCoV-NL63 (isolate Amsterdam 1) was prepared using LLC-Mk2 cells. Six days post-infection (p.i.) infected cells were lysed by freeze-thawing and the resulting supernatant was stored at −80 °C. Mock sample was prepared in the same manner, from non-infected cells. Virus yield was determined according to the method described by Reed and Muench. Briefly, cells infected with serially diluted virus were incubated at 32 °C for 6 days and the appearance of cytopathic effect (CPE) was monitored.

**Isolation of nucleic acids and reverse transcription (RT).** Viral DNA/RNA Kit (A&A Biotechnology, Poland) was used for nucleic acid isolation from cell culture supernatants, according to the manufacturer’s instructions. Cellular RNA was isolated using Fenzol reagent (A&A Biotechnology, Poland), followed by DNase I treatment (Thermo Fisher Scientific, Poland). cDNA samples were prepared with a High Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific, Poland), according to the manufacturer’s instructions.

**Preparation of GC-rich infectious clone 299E virus mutant.** Recombinant HCoV-229E containing a GC-rich region was generated using the vaccinia virus-based reverse genetic system as described previously. In short, vaccinia virus containing the full-length HCoV-229E cDNA in which an Escherichia coli guanine phosphoribosyltransferase (GPT) was inserted between HCoV-229E 10098 and 10930 nucleotides was used to recombine with a plasmid containing HCoV-229E 9398–11580 nucleotides with a number of silent mutations (GC-rich region). The resulting vaccinia virus was used to rescue the mutant HCoV-229E, as described previously. Plasmid DNA, recombinant vaccinia virus and recombinant HCoV-229E identities were confirmed by sequencing.

**quantitative PCR (qPCR).** RNA yields were assessed using real-time PCR (7500 Fast Real-Time PCR; Life Technologies, Poland). HCoV-NL63 cDNA was amplified in a reaction mixture containing 1 × TaqMan Universal PCR Master Mix (Thermo Fisher Scientific, Poland), in the presence of FAM/TAMRA (6-carboxyfluorescein/6-carboxytetramethylrhodamine) probe (100 nM specific) and primers (450 nM each). Reaction was carried out according to the scheme: 2 min at 50 °C and 10 min at 92 °C, followed by 40 cycles of 15 sec at 92 °C and 1 min at 60 °C. Cellular cDNA was amplified in a reaction mixture containing 1 × SYBRGreen Ready PCR Master Mix (Sigma-Aldrich, Poland) in the presence of primers (500 nM each). TATA-box binding protein gene (TBP) was
used as a household gene control and Rox was used as the reference dye. Primer sequences are provided in Table 1. Reaction was carried out according to the scheme: 10 min at 50 °C and 5 min at 95 °C, followed by 40 cycles of 30 sec at 95 °C, 30 sec at 56 °C and 45 sec at 75 °C. In order to assess the copy number for each gene, DNA standards were prepared, as described before47.

Flow cytometry. LLC-Mk2 cells cultured in 6-well plate format (TPP) were washed with sterile 1× PBS, trypsinised, fixed with 4% formaldehyde, permeabilised with 0.1% Triton X-100 in 1× PBS and incubated for 1 h with blocking buffer (10% BSA in 1× PBS with 0.5% Tween 20).

To visualise HA-tagged APOBEC3 proteins after mRNA transfection, cells were incubated for 2 h at room temperature with mouse anti-HA-tag antibody (1 μg/ml, Antibodies-online, Germany) in 1× PBS with 3% BSA and 0.5% Tween 20, followed by 1 h incubation with Alexa Fluor 488-labeled goat anti-mouse antibody (2.5 μg/ml, Molecular Probes, Poland).

To evaluate HCoV-NL63 infection using flow cytometry, cells were incubated for 2 h at room temperature with mouse anti-HCoV-NL63-N antibody (1 μg/ml, Ingenansa, Spain) in 1× PBS with 3% BSA and 0.5% Tween 20, followed by 1 h incubation with Alexa Fluor 488-labeled goat anti-mouse antibody (2.5 μg/ml, Molecular Probes, Poland). Cells were then washed, re-suspended in 1× PBS and analysed with FACS Calibur (Becton Dickinson, USA) using Cell Quest software.

APOBEC3 expression constructs and site-directed mutagenesis. Sequence verified, C-terminal HA-tagged pTracer CMV2 plasmids (Thermo Fisher Scientific, Poland) encoding human A3A (GeneBank accession number (GB) NM_145699), A3C (GB: NM_014508), A3DE (GB: NM_152426), A3F (GB: NM_145298), APOBEC3G (GB: NM_021822) and A3H (GB: NM_181773) were prepared. Catalytic mutants of A3C (A3C_E68Q = mA3C), A3F (A3F_E251Q = mA3F) and A3H (A3H_E56Q = mA3H) were prepared using Quick Change mutagenesis with Phusion polymerase (Thermo Fisher Scientific, Poland) 48. Sequences of oligonucleotides used are listed in Table 2.

Design of “hot-spot” RNA. Plasmid construct containing several GC-rich domains was ordered at GeneArt Gene Synthesis (Thermo Fisher Scientific, Poland). The region was sequenced with primers 5′-CTG GCC TCG TAG GCC TAA TAC GAC TCA CTA TAG GGA GAC TGA GAG AAC CCA CTG CTT AC-3′ and 5′-CTG TGG AAA ACC TTT GGC ATC-3′. The plasmid was amplified in E. coli TOP10 strain and further used as a template in in vitro transcription reaction using T7 promoter, as described below.

Preparation of mRNA molecules for transfection. APOBEC3 plasmids described above were used as templates for amplification of APOBEC3 genes with forward primer containing the T7 promoter region (5′-TGG GCC TCG TAG GCC TAA TAC GAC TCA CTA TAG GGA GAC TGA GAG AAC CCA CTG CTT AC-3′) and specific reverse primer (listed in Table 2, 500 nM each). Reaction was carried out with Phusion polymerase (Thermo Fisher Scientific, Poland), according to the manufacturer’s instructions. DNA templates were purified using GeneJET PCR Purification Kit (Thermo Fisher Scientific, Poland) and further used for in vitro transcription with T7 RiboMAX Express Large Scale RNA Production System and Ribo m7G Cap Analog (Promega, Poland). Resulting RNA was precipitated with LiCl, purified with 70% ethanol and polyadenylated using Poly(A)

| Target   | Primer | Primer sequence (5′-3′) |
|----------|--------|------------------------|
| HCoV-NL63 | Sense   | CTG TGG AAA ACC TTT GGC ATC |
|          | Antisense | CTG TGG AAA ACC TTT GGC ATC |
|          | Probe   | ATG TTA TTC AGT GCT TTT GTG TCT CTC GTG AT |
| A3A      | Sense   | TGG TTC CTT CTT TGC AGT TGG ACC |
|          | Antisense | GCA GCA TTT GCA GTG CCT CTT TAT |
| A3B      | Sense   | AGC ACA TGG GCT TCC TAT GCA ACG |
|          | Antisense | ACG AGA TGA ACC AAG TGA CCC TGT |
| A3C      | Sense   | AAC GAA ACT TGG CTG TGC TTC ACC |
|          | Antisense | AGA CAG TAT GTC GTC GCA GAA CCA |
| A3D      | Sense   | ACG CAG GAG GTG TAT TCC CGG TT |
|          | Antisense | GGT GCT CAG CCA AGA ATT TGG TCA |
| A3F      | Sense   | TCC GTG GAG ATC ATG GGC TAC AAA |
|          | Antisense | TGC AGC TGG TGT TCC AGG AAT AGA |
| A3G      | Sense   | GCT GTG CCC AGG AAA TGG CTA AAT |
|          | Antisense | ACA AAG GTG TTC CAG CAG TGC TTA |
| A3H      | Sense   | TGA CTT CAT CAA GGC TCA CGA CCA |
|          | Antisense | AGC AGC AAA CTG GGA GCC CAT |
| TBP      | Sense   | CCC ATG ACT CCC ATG ACC |
|          | Antisense | TTT ACA ACC AAG ATT CAC TGT GG |

Table 1. Primers and probes used for a RT-qPCR.
CO₂ and transfected with 30 mM 4-(2-hydroxyethyl)-1-piperazine-ethane-sulfonic acid (HEPES; pH 7.5), 100 mM KCl, 25 mM 3 days post transfection in deaminase activity buffer, containing 0.2% Surfact-Amps NP-40 (Sigma-Aldrich, Canada). A rabbit anti-HA Taq antibody (1:500 dilution; Cell Signalling Technology, Poland) and anti-HA goat IgG (65 ng/ml; Dako, Denmark) were used to detect the HCoV-NL63 nucleocapsid protein. A rabbit anti-GAPDH antibody (1:5000 dilution; Cell Signalling Technology, Poland) and horseradish peroxidase-labeled goat anti-rabbit IgG (0.35 μg/ml; Sigma-Aldrich, Poland) were used to detect the GAPDH in cell lysates. All antibodies were diluted in 1% skimmed milk/TBS-Tween (0.1%). The signal was developed using the Immobilon Western Chemiluminescent HRP Substrate (Millipore, USA) and visualised by exposing the membrane to an X-ray film (Kodak, Poland).

**Table 2.** Primers used for generation of expression constructs and deaminase mutants.

| Target  | Primer  | Primer sequence (5’–3’) |
|---------|---------|------------------------|
| mA3C    | Sense   | GTC ATG CAC AAA GGT GCT TCC TCT CTT G |
|         | Antisense | GCA CCT TTT GTC ATG ACA ATG GGT CTC |
| mA3F    | Sense   | GTC ATG CAC AAA GGT GCT TCC TCT CTT G |
|         | Antisense | GCA CCT TTT GTC ATG ACA ATG GGT CTC |
| mA3H    | Sense   | GCC ATG CAC AAA TTT GCT TTA TTA ACG |
|         | Antisense | GCA AAT TTT GTC ATG GCA CTT TTT CTT G |

**Transfection of plasmid DNA and mRNA.** 293T cells were seeded on 10 cm² dishes, cultured for 24 h at 37°C with 5% CO₂ and transfected with 8 μg of plasmid per dish using polyethylenimine (Sigma-Aldrich, Poland). LLC-Mk2 and Huh-7 cells were seeded on 6-wells plates (TPP, Switzerland), cultured for 24 h at 37°C with 5% CO₂ and transfected with 3 μg mRNA per well (transcripts encoding APOBEC3 proteins or control GFP) using TransIT mRNA Transfection Reagent (Mirus, USA). Efficiency of transfection was assessed 24 h post-transfection in GFP-transfected cells using EVOS fluorescent microscope (Thermo Fisher Scientific, Poland).

**Western blot analysis.** Cells were trypsinised, centrifuged and resuspended in RIPA buffer (50 mM Tris, 150 mM NaCl, 1% Nonidet P-40, 0.5% sodium deoxycholate, 0.1% SDS, pH 7.5) followed by lysis in RIPA buffer for 30 min on ice. Subsequently, samples were centrifuged (10 min at 12,000 × g) and the pelletted cell debris was discarded. Resulting supernatants were mixed with sample buffer (0.5 M Tris pH 6.8, 10% SDS, 50 mg/ml DTT), boiled for 5 min, cooled on ice, and separated on 10% polyacrylamide gels alongside dual color Page Ruler pre-stained Protein size markers (Thermo Fisher Scientific, Poland). The separated proteins were then transferred onto a Westran S PVDF membrane (GE Healthcare, Poland) by wet blotting (Bio-Rad, Poland) for 1 h, 100 Volts in transfer buffer: 25 mM Tris, 192 mM glycine, 20% methanol at 4°C. The membranes were then blocked by overnight incubation at 4°C in TBS-Tween (0.1%) buffer supplemented with 5% skimmed milk (BioShop, Canada). A rabbit anti-HA Taq antibody (1 μg/ml; Antibodies-online, USA) and horseradish peroxidase-labeled goat anti-rabbit IgG (0.35 μg/ml; Sigma-Aldrich, Poland) were used to detect the HA-tagged APOBEC3 proteins in cell lysates. Mouse anti-HCoV-NL63-N protein antibody (500 ng/ml; Ingenansa, Spain) and horseradish peroxidase-labeled rabbit anti-mouse IgG (65 ng/ml; Dako, Denmark) were used to detect the HCoV-NL63 nucleocapsid protein. A rabbit anti-GAPDH antibody (1:5000 dilution; Cell Signalling Technology, Poland) and horseradish peroxidase-labeled goat anti-rabbit IgG (0.35 μg/ml; Sigma-Aldrich, Poland) were used to detect the GAPDH in cell lysates. All antibodies were diluted in 1% skimmed milk/TBS-Tween (0.1%). The signal was developed using the Immobilon Western Chemiluminescent HRP Substrate (Millipore, USA) and visualised by exposing the membrane to an X-ray film (Kodak, Poland).

**Nucleocapsid protein binding analysis.** APOBEC3-expressing and control 293T cells were harvested 3 days post transfection in deaminase activity buffer, containing 0.2% Surfact-Amps NP-40 (Sigma-Aldrich, Poland), 30 mM 4-(2-hydroxyethyl)-1-piperazine-ethane-sulfonic acid (HEPES; pH 7.5), 100 mM KCl, 25 mM NaCl, 1.5 mM MgCl₂, 1 mM DTT, 1 × Halt protease and phosphatase inhibitor cocktail, and 10% glycerol. Samples were centrifuged (10 min at 12,000 × g), pelleted cell debris was discarded and resulting supernatants were stored at −80°C. HCoV-NL63 N protein was prepared as described previously. Cell lysates were incubated
for 1 h at 37 °C with the purified N protein in 1 × activity buffer, containing 100 mM KCl, 10 mM HEPES (pH 7.4), 100 µM ZnCl₂, 1 mM DTT and 1 mM EDTA. Subsequently, 15 µl of agarose spheres coated with anti-HA antibodies (Pierce, Poland) was added and samples were loaded onto co-immunoprecipitation columns. After over-night incubation at 4 °C on a rotary shaker, proteins were eluted, according to the manufacturer’s instructions.

Analysis of HCoV-NL63 co-localisation with APOBEC3C. LLC-Mk2 cells were seeded on 6-wells plates and cultured for 24 h at 37 °C with 5% CO₂. 1.5 µg of HCoV-NL63 N mRNA and 1.5 µg of A3C mRNA was co-transfected using the mRNA-In reagent (AMS Biotechnology, United Kingdom). After 72 h incubation at 37 °C with 5% CO₂, cells were fixed with 4% formaldehyde, permeabilised with 0.1% Triton X-100 in 1 × PBS and incubated overnight at 4 °C with 10% BSA in 1 × PBS with 0.5% Tween 20. To visualise HCoV-NL63 N molecules, cells were incubated for 2 h at room temperature with rabbit anti-HCoV-NL63-N serum (1:100 dilution, kindly provided by dr Lia van der Hoek, UvA, The Netherlands) in 1 × PBS with 3% BSA and 0.5% Tween 20, followed by 1 h incubation with Atto 488-labeled goat anti-rabbit IgG (5 µg/ml; Sigma-Aldrich, Poland). To detect HA-tagged A3C protein cells were incubated for 2 h at room temperature with mouse Alexa Fluor 647-labeled anti-HA-tag antibody (10 µg/ml; Thermo Fisher Scientific; Poland) in 1 × PBS with 3% BSA and 0.5% Tween 20. Nuclear DNA was stained with DAPI (0.1 µg/ml; Sigma-Aldrich, Poland). Immunostained cultures were mounted on glass slides with ProLong Gold antifade medium (Thermo Fisher Scientific; Poland). Fluorescent images were acquired under a Zeiss LSM 710 confocal microscope (Carl Zeiss Microscopy GmbH). Images were acquired using ZEN 2012 SP1 software (Carl Zeiss Microscopy GmbH) and processed using ImageJ 1.47v (National Institutes of Health, Bethesda, Maryland, USA).

Statistical analysis. All the experiments were performed in triplicate and the results are presented as mean ± SD. Student’s t-test was used to determine significance of the obtained results. P values < 0.05 were considered significant.

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Author Contributions

A.M. and P.V. conducted the experiments. E.K., S.Z., M.O. and V.T. provided materials for the study. Z.R. participated in confocal imaging. A.M. and K.P. designed the experiments, analysed the data and wrote the manuscript. All authors reviewed the manuscript.

Additional Information

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