SARS-CoV-2 NSP5 Antagonizes the MHC II Antigen Presentation Pathway by Hijacking Histone Deacetylase 2

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The clearance of SARS-CoV-2 requires a multi-faceted immune response that is initiated by innate immune cells, with infection ultimately resolved by adaptive immune mechanisms. Induction of adaptive immunity to SARS-CoV-2 is dependent on the presentation of viral antigens on MHC II by professional antigen presenting cells such as dendritic cells and macrophages, to induce robust activation of CD4+ T cells. SARS-CoV-2 interferes with antigen presentation by downregulating MHC II on the antigen presenting cells of COVID-19 patients, but the molecular mechanism mediating this process is unelucidated. In this study, analysis of protein and gene expression in human antigen presenting cells reveals that the expression of MHC II is inhibited by the SARS-CoV-2 main protease, NSP5. Suppression of MHC II expression occurs via downregulation of the transcription factor CIITA, which is required for MHC II expression. This downregulation of CIITA is independent of NSP5’s proteolytic activity, and rather, NSP5 delivers HDAC2 to the CIITA promoter via interactions with IRF3. Here, HDAC2 deacetylates and inactivates the CIITA promoter. This loss of CIITA expression prevents further expression of MHC II, with this suppression alleviated by ectopic expression of CIITA or knockdown of HDAC2. These results identify a novel mechanism by which SARS-CoV-2 can limit antigen presentation on MHC II, thereby delaying or weakening the subsequent adaptive immune response.

SARS-CoV-2 | NSP5 | Major Histocompatibility Complex II | CIITA | Antigen Presentation | IRF3

Introduction

First identified in December 2019, severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has rapidly become a leading global cause of morbidity and mortality. This high virulence is due, in part, to the presence of multiple mechanisms which allow SARS-CoV-2 to evade and alter host immune responses, thereby delaying viral clearance and prolonging the period in which patients remain infected [1,2]. Of note, humoral immunity against SARS-CoV-2 wanes twice as quickly compared to other human-infecting coronaviruses [3]. Even though vaccination has greatly reduced morbidity and mortality, breakthrough infections still occur and are becoming more common due to the emergence of new, more immunoevasive, SARS-CoV-2 variants [4,5]. Therefore, understanding the immunoevasion mechanisms of SARS-CoV-2 is important to better understand this deadly disease and to develop better targeted treatments and vaccines.

Although professional antigen presenting cells (pAPCs) such as macrophages and dendritic cells (DCs) do not express the SARS-CoV-2 receptor ACE2, they can be infected via Fc-receptor dependent phagocytosis of antibody-opsonized virions, and potentially through the efferocytosis of SARS-CoV-2 infected apoptotic cells [6–12]. While SARS-CoV-2 is unable to establish a productive infection in macrophages or DCs, viral early genes are expressed and drive a multi-pronged immunoevasion response. Firstly, the expression of pro-inflammatory cytokines is induced and contributes to the cytokine storm [12]. This cytokine response is typified by high
circulating levels of IL-2, IL-6, IL-7, IL-8, IL-10, G-CSF, MCP-1, MIP1-α, and TNF-α [13–15]. While this mechanism drives a potent inflammatory response, the cytokine profile is more typical of bacterial infections, and promotes both NK cell exhaustion and reduced NK cell cytotoxicity, thereby producing a non-productive innate immune response that can exacerbate tissue damage [16,17]. Secondly, SARS-CoV-2 suppresses the anti-viral interferon (IFN) pathway, reducing the production of type I and type II IFNs. This suppression is driven by ORF6, which sequesters inactive signal transducer and activator of transcription 1 (STAT1) and STAT2 in the cytosol, thereby preventing their nuclear translocation and blocking the primary signalling pathway that initiates IFN responses [18]. Moreover, membrane protein and non-structural protein 13 further inhibits IFN-I production by degrading TANK-binding kinase 1 [19,20]. Thirdly, infected macrophages and dendritic cells often die, resulting in long-term depletion of some subsets [21].Fourthly, SARS-CoV-2 directly suppresses antigen presentation on Major Histocompatibility Complex class I (MHC I) via ORF8, which directs the trafficking of MHC I to the lysosome where it is degraded [22]. MHC I is further suppressed by ORF6, which inactivates CIT/A/NLRC5—the master regulator of MHC I expression [23]. Finally, infection of alveolar DCs reduces their ability to migrate to draining lymph nodes and suppresses expression of MHC class II and the class II transcriptional activator (CIITA) required for MHC II expression [24]. This suppression of MHC II occurs across a range of pAPCs in SARS-CoV-2 infected patients and in cells infected in-vitro [25–27]. Moreover, some non-professional antigen presenting cells express MHC II in response to infection – including type II alveolar epithelial cells which are one of the primary targets of SARS-CoV-2 infection in the alveolus [28,29]. However, the cellular mechanism which downregulates MHC II in these cells has remained elusive.

MHC II expression is driven by type II interferon signaling [30]. Activation of the IFN-γ receptor complex leads to the structural rearrangements in the receptor complex and activation of Janus kinase 1 and 2 (JAK1/2) tyrosine kinases and phosphorylation and homodimerization of STAT1, which translocates to the nucleus to induce transcription of IFN-γ inducible genes [31]. Here, STAT1 activates interferon regulatory factor -1 and -3 (IRF-1/3), with STAT1 and IRF-1/3 then cooperatively inducing expression of CIITA [32,33]. CIITA, via its intrinsic acetyltransferase activity, can then acetylate histones at the MHC II promoter, which decondenses the chromatin to allow access for transcription factors that regulate MHC II expression [34]. Once the chromatin is opened, CIITA and regulatory factor X (RFX) form an enhanceosome complex on the MHC II promoter which recruits and activates additional transcription factors that induce the transcription of MHC II [35]. While inhibition of STAT1 nuclear import by SARS-CoV-2 ORF6 may account for some inhibition of MHC II expression [18], tissue-resident DCs natively express significant amounts of CIITA [36], whereas ORF6 expression is limited until 12-16 hours post-infection [37] – more than sufficient time to induce presentation of SARS-CoV-2 antigens following infection. Moreover, IFN-independent mechanisms can drive some MHC II expression in macrophages and DCs [38–40]. Therefore, a more direct form of MHC II suppression is likely invoked by SARS-CoV-2. A critical regulator of MHC II expression is histone deacetylase 2 (HDAC2), which can suppress the expression of both CIITA and MHC II through deacetylation of histones within their promoters [41]. Gordon et al. recently mapped the SARS-CoV-2 protein interactome and identified non-structural protein 5 (NSP5) as an HDAC2 interactor, and identified a putative NSP5 cleavage site near the nuclear localization signal (NLS) of HDAC2 [42]. NSP5 – also known as main protease and 3-chymotrypsin like protease – is translated as part of the polyprotein expressed early after viral entry, and cleaves this polyprotein into 11 individual proteins which form the complex that translates full viral RNA and allows for reproduction of the viral genome and the production of mature virions [43]. Through interactions with HDAC2, NSP5 may mediate the epigenetic reprogramming of infected cells, which in pAPCs may include suppression of MHC II expression. Indeed, epigenomic changes are required for SARS-CoV-2 reproduction [44,45], and similar epigenetic reprogramming is known to suppress MHC II expression in Middle East respiratory syndrome–related coronavirus (MERS-CoV) [46]. Therefore, in this study we tested the hypothesis that SARS-CoV-2 NSP5 inhibits MHC II expression through interactions with HDAC2.
Results

SARS-CoV-2 NSP5 Downregulates MHC II in Professional Antigen Presenting Cells

SARS-CoV-2 NSP5 is the viral main protease that plays an essential role in viral infection and pathogenesis [47,48]. As main proteases are required for the processing of coronavirus polyproteins [49], deletion of NSP5 or its inactivation abrogates productive infection by SARS-CoV-2 [43,49,50]. To assess the effects of NSP5 on the MHC II antigen presentation system, primary human moDCs were transduced with either empty or NSP5-expressing lentiviral vectors bearing a zsGreen marker. Flow cytometry was used to quantify surface expression of total MHC II on transduced (zsGreen+) moDCs (Figure 1A-B, S1), with NSP5 expression reducing the cell surface expression of MHC II to an extent similar to that observed in SARS-CoV-2 patients (Figure 1C-D) [25–27]. This downregulation was not a general suppression of the MHC II presentation system, as expression of the co-stimulatory molecule CD86 was not affected by NSP5 expression (Figure 1E). Some pathogens reduce their immunogenicity by diverting intracellular trafficking such that MHC II molecules fail to reach the cell surface [51]. To test this possibility, we transduced J774.2 macrophages with NSP5-expressing or empty lentiviral vectors, labeled the plasma membrane wheat-germ agglutinin, followed by labeling for total cellular MHC II. and three-dimensional reconstructions of these cells used to differentiate between cytosolic/vesicular MHC II and cell-surface MHC II (Figure 1F). Quantitation of these micrographs revealed no changes in the portion of MHC II localized to the cell surface versus intracellular vacuoles (Figure 1G) but did identify the same decrease in MHC II expression that was observed with flow cytometry (Figure 1D,H), indicating that NSP5 does not affect the trafficking of MHC II to the cell surface, but rather decreases its overall expression.

![Figure 1: NSP5 Suppresses MHC II at the Protein Level in Primary Monocyte-Derived Human Dendritic Cells.](https://example.com/figure1)

moDCs were transduced with lentiviral vectors lacking a transgene (empty vector) or bearing NSP5, with both vectors containing an IRES-zsGreen marker. A-B) Flow cytometry dot plots showing cell surface MHC II expression and transduction (zsGreen+) of moDCs transduced with an empty vector control (A) or NSP5-expressing lentiviral vector (B). C) Histogram of cell surface MHC II expression levels on moDCs transduced with an empty vector (red) or NSP5-expressing vector (cyan). D-E) Quantification of cell surface MHC II (D) and CD86 (E) in moDCs transduced with either an empty (Empty) or NSP5-expressing (NSP5) lentiviral vector. MFI is normalized to the MFI of the untransduced cells in the Empty-vector sample. F) Z-slice through a macrophage stained for the plasma membrane and MHC II, showing the segmentation of vesicular/cytosolic versus surface (membrane) MHC II. Scale bar is 10 µm. G-H) Quantification of the fraction of MHC II on the plasma membrane (G) and total cellular MHC II (H) in macrophages. Data is representative of, or quantifies a minimum of 3 independent experiments, * = p < 0.05; n.s. = p > 0.05 compared to empty vector, Mann-Whitney test.
NSP5 Suppresses CIITA and MHC II Transcription

Next, the subcellular localization of NSP5 was determined to identify potential mechanisms accounting for the downregulation of MHC II. Quantitative microscopy of HeLa cells co-transfected with NSP5-FLAG, the ER marker KDEL-GFP, the Golgi marker GaIT-mCherry, and with the nuclei stained with Hoechst, determined that approximately half of the cellular NSP5 was localized to the nucleus, with the remainder associated with the ER (Figure 2A-B). The nuclear localization of NSP5 was confirmed by blocking importin-mediated nuclear transport (Figure 2C-D). While localization to the ER is consistent with the known role of NSP5 in forming the viral replication complex [52], the role for nuclear NSP5 remains unclear. Unexpectedly, bioinformatic analysis of NSP5 failed to identify either a classical or bipartite nuclear localization signal, or a nuclear export signal (data not shown). As such, NSP5 is likely carried into the nucleus through interactions with other cellular proteins and not via direct transport of NSP5 by the nuclear import machinery, similar to the hepatitis delta antigen [53].

Figure 2: NSP5 Suppresses CIITA and MHC II expression. A-B) Fluorescent z-projection (A) and Manders colocalization analysis (B) quantifying the portion of NSP5 colocalized with the nucleus of HeLa cells co-transfected with NSP5-FLAG (yellow), GaIT-mCherry (Golgi/magenta), KDEL-eGFP (ER/cyan), and stained with Hoechst (DNA/grey). Manders colocalization analysis compares the fraction of NSP5 colocalized with the nucleus, ER and Golgi (Manders) to the Manders ratio from the same image when the NSP5 image was randomized (Random). C-D) Fluorescent z-projections (C) and quantification (D) of the fraction of NSP5 in the nucleus of vehicle-treated (DMSO) versus ivermectin-treated HeLa cells expressing NSP5-FLAG (yellow) and stained for DNA with Hoechst (cyan). E) RT-qPCR
quantification of RFX5, CIITA, and MHC II mRNA levels in moDCs transduced with empty or NSP5-expressing lentiviral vectors. F) Quantification of the promoter activity of the CIITA pI, CIITA pIII/IV, and MHC II promoters using a dual-luciferase assay in RAW 264.7 macrophages co-transfected with empty or NSP5-expressing lentiviral vectors. G) Quantification of MHC II promoter activity in RAW 264.7 macrophages co-transfected with CIITA and NSP5. (-) indicates the sample was transfected with empty vector rather than CIITA or NSP5. Scale bars are 10 µm. Images are representative of a minimum of 30 cells captured across 3 independent experiments. n = minimum of 3. * = p < 0.05; n.s. = p > 0.05, compared to Random (B), DMSO (D), or Empty Vector (E-F), paired t-test (B) or Mann-Whitney test (D, E, F).

The nuclear localization of NSP5 suggests that downregulation of MHC II occurs via a transcriptional mechanism. To test this hypothesis, we used RT-qPCR to measure the mRNA levels of MHC II, as well as RFX5 and CIITA—the two transcription factors which act as the master regulators of MHC II expression. Interestingly, while RFX5 expression was unchanged, NSP5 significantly downregulated the expression of CIITA and MHC II (Figure 2E). In humans, CIITA is transcribed from three separate promoters: pI—which drives expression in myeloid cells, pIII—which drives expression in lymphoid cells, and pIV—which drives IFN-γ-induced expression in non-immune cells such as epithelia [54,55]. To assay the activity of the MHC II and CIITA promoters, we constructed dual-luciferase reporters of the MHC II promoter, the CIITA pI promoter, and the CIITA pIII/pIV promoters, followed by quantification of NSP5’s effect on these promoters’ activity in macrophages. NSP5 expression strongly suppressed both the MHC II promoter and the CIITA pI promoter, whereas the CIITA pIII/pIV promoters were minimally active, producing insufficient signal to observe any effect of NSP5 (Figure 2F). MHC II transcription is dependent on CIITA, therefore this suppression of MHC II expression may be due to NSP5-dependent repression of CIITA expression, or alternatively, may be a product of NSP5 suppression of both the CIITA and MHC II promoters. To differentiate between these possibilities, we quantified MHC II promoter activity in cells ectopically expressing CIITA and NSP5 (Figure 2G). CIITA expression greatly increased MHC II promoter activity, with co-expression of NSP5 having no effect on MHC II promoter activity in the presence of ectopically expressed CIITA. These data indicate that NSP5 suppresses the transcription of CIITA, and the resulting absence of CIITA then limits MHC II expression.

NSP5-Mediated Suppression of MHC II Expression is Dependent on HDAC2

MHC II expression is regulated by CIITA through several mechanisms. Induction of MHC II expression begins with CIITA binding to distal enhancers located several kilobases 5' to the MHC II promoter [56], followed by acetylation of the core MHC II promoter by CIITA. This acetylation induces the formation of an enhancerome complex comprised of CIITA, RFX5, CREB and NF-Y, wherein CIITA activates TAF family transcription factors via its intrinsic protein acetylation and kinase activity, thus initiating MHC II transcription [57,58]. CIITA expression is induced by IFNγ through activation of the transcription factor IRF1, and via toll-like receptor (TLR) and IL-1 family cytokines via activation of the transcription factor IRF3 [32,59,60], and is negatively regulated by HDAC2-mediated promoter deacetylation [61]. Critically, SARS-CoV-2 NSP5 is known to interact with HDAC2, suggesting that NSP5 utilizes HDAC2 to silence the CIITA promoters [42]. Consistent with this model, HDAC2 co-precipitated with FLAG-tagged NSP5 in an anti-FLAG immunoprecipitation (Figure 3A). Knockdown of HDAC2 (Figure 3B) had a profound restorative effect on CIITA and MHC II mRNA levels, with HDAC2 knockdown not only reversing – but increasing above baseline – expression of both genes (Figure 3C). NSP5 is comprised of a proteolytic domain formed by the interface of the globular A and B domains, which positions two key catalytic residues (H41 and C145) in a binding cleft formed between the two domains. A C-terminal B’ chain folds over this cleft, coordinating both the substrate and three water molecules within the active site [62]. We inactivated the catalytic site by generating NSP5H41A and NSP5C145S point mutants, and also deleted the proteolytic (NSP5Δ1-192) and B’ domains (NSP5Δ199-306) (Figure 3D). The deletion mutants were unstable and could not be expressed at levels amenable for immunoprecipitation (data not shown), but immunoprecipitation of the point mutants revealed that both retained some binding capacity for HDAC2 (Figure 3E), and retained their ability to suppress CIITA and MHC II promoter activity (Figure 3F).
Figure 3: NSP5 Requires HDAC2 to Suppress MHC II Expression. A) Co-immunoprecipitation of HDAC2 with NSP5-FLAG from primary human moDCs transduced with empty or NSP5-FLAG-expressing lentiviral vectors. B) Expression of HDAC2, as quantified by immunoblotting, in moDCs that are untreated (UT) or treated with either scrambled (siScrm) or HDAC2-targeting (siHDAC2) siRNAs. HDAC2 protein level is quantified relative to GAPDH protein levels in the same lysates. C) RT-qPCR quantification of the impact of scrambled (siScrm) or HDAC2-targeting (siHDAC2) siRNA on CIITA and MHC II mRNA levels in primary human moDCs transduced with empty or NSP5-expressing lentiviral vectors. D) Protease-inactivating (H41A and C145S) point-mutants, and deletion of the catalytic (A/B, Δ1-192) and ligand-stabilizing (B’, Δ199-306) domains were generated to assay the roles of these sites in NSP5 activity. E) Immunoprecipitation of HDAC2 with NSP5<sub>H41A</sub> and NSP5<sub>C145S</sub> mutants. F) Dual-luciferase assay quantification of CIITA pI and MHC II promoter activity in RAW264.7 macrophages co-transfected with wild-type NSP5 (WT) or NSP5<sub>H41A</sub> and NSP5<sub>C145S</sub> mutants. n = 5, * = p < 0.05; n.s. = p > 0.05, Kruskal-Wallis test with Dunn correction (C) or Mann-Whitney test (F).

NSP5 Induces Deacetylation of the CIITA and MHC II Promoters
Given the dependence of NSP5 on HDAC2 for its suppressive effect on MHC II transcription, it is likely that NSP5 is modulating histone acetylation at the CIITA or MHC II promoters. FRET microscopy was used to quantify promoter acetylation, measuring the transfer of excitation energy from a Cy3-labeled anti-acetyl-lysine antibody to ATTO647-N labeled FISH probes specific to the MHC II promoter plus 3 kb of 5′ enhancer regions, the CIITA-pI promoter plus 1.5 kb of 5′ sequence, and the region containing the CIITA pIII and pIV promoters (CIITA pIII/IV). Primary human macrophages were transduced with empty or NSP5-expressing lentiviral vectors and treated with either scrambled or HDAC2-targeting siRNA. Acetylated-lysine staining was concentrated in the nucleus, with weaker staining present in the cytosol, with one or two FISH probes in-focus within each nucleus (Figure 4A). Neither NSP5 expression, nor HDAC2 depletion, altered the quantity or distribution of cellular acetyl-lysine staining (Figure 4B-C), indicating that neither NSP5 expression nor HDAC2 knockdown globally affected lysine acetylation. In untransduced cells, and in cells transduced with an empty vector, a significant FRET signal could be observed at the CIITA pI and MHC II promoters, while a much weaker FRET signal was detected.
at the CIITA pIII/IV promoter (Figure 4D-F), consistent with the pl promoter driving CIITA expression in myeloid cells. Critically, ectopic expression of NSP5 significantly reduced acetylation at all three promoters, with HDAC2 knockdown restoring promoter acetylation in NSP5-expressing cells (Figure 4D-F).

It seemed unusual that a virus whose closest known wild ancestor is found in bats (BANAL-20-236) would have such potent activity against the human MHC II and CIITA promoters [63,64]. This activity may suggest that NSP5 has undergone significant evolution following the zoonotic transmission of SARS-CoV-2 to humans, or alternatively, that HDAC2 is highly conserved between bats and humans. Phylogenetic and amino acid conservation analysis revealed that the latter is the case – HDAC2 is strikingly conserved across vertebrates, with most residues completely conserved between humans and a range of vertebrates known to be infected by SARS-CoV-2 or to frequently contact humans (Figure 4G). A similar degree of conservation is also observed across all major clades of vertebrates (Figure S2). In contrast, NSP5 is modestly conserved across the major coronavirus clades, but shows high conservation of its catalytic site, as well as complete conservation between SARS-CoV, SARS-CoV-2, and bat coronavirus BANAL-20-236 (Figure 4H).

Figure 4: NSP5 Modulates Acetylation of the CIITA and MHC II promoters. A) Representative micrograph of primary human macrophages transduced with empty or NSP5-expressing lentiviral vectors. Cells were stained for acetyl-lysine (Lys\(^{ac}\), cyan), ATTO647N-labeled MHC II FISH probes (yellow), with the FRET signal between the Lys\(^{ac}\) and FISH probes within the insert shown in the FRET panel. Scale bars are 10 \(\mu\)m. B-C) Quantification of the nuclear:cytosolic acetyl-lysine distribution (B) and total cellular acetyl-lysine content (C) in primary human macrophages transduced with empty or NSP5-expressing lentiviral vectors and treated with scrambled (siScrm) or HDAC2-targeting (siHDAC2) siRNAs. “Untransduced” quantifies macrophages which did not take up the empty or NSP5-expressing lentiviral vector. D-F) Quantification of the acetylation of the CIITA pl (D), CIITA pIII/pIV (E) and MHC II (F) promoters, as quantified by the FRET efficiency between Cy3-labeled anti-acetyl-lysine staining and ATTO647N-stained FISH probes in the same cells used for quantification of panels B and C. G-H) Domain structure, amino acid conservation (Shannon Entropy, H (X)), and phylogenetic trees for (G) HDAC2 across a range of vertebrate species which humans frequently contact or which are possible vectors of SARS-CoV-2, and (H) representative species of the four genera of coronaviruses. HDAC2 consists of four major domains, a homodimerization (HD) domain, deacetylase domain, an IAC (E/D)E motif, and a coil-coil domain which mediate interactions with other transcription factors. NSP5 contains a bipartite protease domain (A/B) with two critical catalytic residues (H41 and C145), and a B’ C-terminal domain which stabilizes ligands in the protease domain. Data are representative of (A) or quantifies (B-F) three independent experiments. * = p < 0.05; n.s. = p > 0.05, Kruskal-Wallis test with Dunn correction. Flat bars indicate the statistical significance for all groups beneath the bar; legged bars indicate the statistical significance between the groups below the legs.

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NSP5 Targets the CIITA Promoter via Interactions with IRF3

As NSP5 is a protease, and proteolytic cleavage of some HDACs are known to promote their activity [65], we next assessed whether NSP5 was capable of cleaving HDAC2. Immunoblot analyses were performed to compare differences in HDAC2 protein levels in cells transfected with either NSP5-expressing or empty lentiviral vectors. Despite a predicted NSP5 cleavage site [42], we observed no change in HDAC2 protein levels or mass in NSP5-expressing cells (Figure 5A). While this predicted cleavage site should produce an 11 kDa decrease in the mass of HDAC2, HDACs are often cleaved closer to the C-terminus, resulting in cleavage products barely distinguishable from the uncleaved protein. As such, we constructed an intramolecular FRET cleavage probe comprised of an mVenus-HDAC2-mTurquoise2 fusion protein. Intact, this probe has high FRET efficiency between the mVenus and mTurquoise2 fluorophores, with cleavage resulting in separation of the fluorophores and loss of the FRET signal (Figure S3). Consistent with the immunoblotting results, no change in FRET efficiency was observed in NSP5-transfected versus empty-vector transfected cells in either the cytosol or nucleus (Figure 5B), confirming that NSP5 does not modulate HDAC2 activity through proteolytic cleavage. We next quantified the subcellular distribution of HDAC2 and the NSP5 mutants generated in Figure 3 and found that inactivation of the NSP5 catalytic site had no impact on the distribution of either protein (Figure 5C-D). Although poorly expressed, the rare cells expressing NSP5 deletion mutants lacking the proteolytic (NSP5Δ1-192) or B′ (NSP5Δ199-306) domains displayed a hyper-nuclear localization compared to wild-type, but also had no impact on the distribution of HDAC2, indicating that HDAC2 unlikely to be the protein used by NSP5 to gain access to the nucleus (Figure 5C-D). Naik et al demonstrated that NSP5 can interact with IRF3, and IRF3 is known to bind to the CIITA PI promoter, suggesting that NSP5 may deliver HDAC2 to the CIITA promoter via interactions with IRF3 [66–68]. Using FISH-FRET analysis of A549 cells—a type II human alveolar lung epithelial cell line—we found that IFNγ stimulation increased CIITA PI, PII/IV and MHC II promoter acetylation, and as expected, NSP5 expression inhibited promoter acetylation (Figures 5E-G, S4). Critically, this suppression of promoter acetylation by NSP5 was largely reversed by an IRF3-targeting siRNA, consistent with a model in which NSP5 delivers HDAC2 to the CIITA promoter via interactions with IRF3 (Figures 5E-G, S5).
Discussion

In this study we demonstrate that SARS-CoV-2 NSP5 suppresses expression of CIITA across a range of professional and non-professional antigen presenting cells, thereby suppressing MHC II expression. This suppression of CIITA expression occurs via a novel pathway in which NSP5 delivers HDAC2 to the CIITA promoter via interactions between NSP5 and promoter-bound IRF3. HDAC2 then deacetylates histones at the CIITA promoter, decreasing CIITA expression, thereby blocking MHC II expression (Figure S5). These findings explain the decreased MHC II expression observed COVID-19 patients and in in-vitro infection models [25–27].

This suppressive mechanism occurred across multiple types of professional APCs (human moDCs, human macrophages, and a mouse macrophage cell line) and in human non-professional APCs (type II alveolar epithelial cells). As SARS-CoV-2 infects pAPCs including DCs, macrophages, and B cells—as well as non-professional APCs in the lung—this suppression of MHC II has the potential to dramatically decrease the activation of CD4+ T cells in COVID-19 patients [6–8]. Such a loss of CD4+ T cell activity likely contributes to the shorter-lived humoral immunity and propensity for re-infection of COVID-19 patients, and may contribute to the aberrant formation of memory and effector CD4+ T cell subsets in patients with severe disease [24,69–72].

The suppression of MHC I and MHC II expression, or removal of these proteins from the cell surface, is a commonly employed viral immunoevasion strategy. For MHC I, mis-directing or re-internalizing MHC I after its translation are the predominant mechanisms used to limit immunogenicity of infected cells [73]. For example, HIV-1 Nef mediates the AP-1-dependent endocytosis of MHC I and its sequestration in a Golgi-proximal compartment, thus limiting cytotoxic T cell activity against infected cells [74,75]. Similarly, SARS-CoV-2 utilizes ORF8 to direct MHC I into lysosomes for degradation, and uses ORF6 to suppress MHC I expression by downregulating expression of the transcription factor CITA/NLRC5 [18,22]. Influenza A and B also down regulate surface MHC I via proteasomal degradation or via endocytosis and sequestration in cytosolic vacuoles, respectively [76]. Downregulation of MHC II is rarer, as only viruses that infect APCs have the potential to exhibit this activity. HSV-1 reduces cell surface levels of MHC II by directing MHC II into multivesicular bodies [77,78]. HIV-1 Nef, through accumulation of αβII complexes in intracellular vesicles, suppresses trafficking of peptide-loaded MHC II to the cell surface [51]. Human cytomegalovirus protein US2 degrades HLA-DR-α and DM-α chains, while the US3 protein competes with the invariant chain for binding to MHC II α/β complexes, thereby restricting MHC II intracellular trafficking [79,80]. Transcriptional downregulation of MHC II is also observed, often through targeting of CIITA. Similar to what we have reported in this study, Kaposi sarcoma-associated herpesvirus also targets the IRF3 binding site in the CIITA promoter—but unlike SARS-CoV-2—this virus expresses a viral IRF3 which is inhibitory to CIITA expression [67]. Epstein-Barr virus impairs CIITA expression in B cells by suppressing the activity of E47 and PU.1, thereby preventing these transcription factors from binding the pIII promoter [81,82]. The coronavirus MERS-CoV was shown to transcriptionally downregulate both MHC I and MHC II through an undefined epigenetic mechanism [46]. Given the epigenetic basis of SARS-CoV-2’s suppression of CIITA and MHC II, it is possible that the downregulation of MHC II by MERS-CoV is mediated by the same NSP5-mediated mechanism that we identified in this study, and that this activity may be conserved across pAPC-infecting coronaviruses.

The proteolytic activity of coronaviral NSP5 is known to be important for its immunosuppressive activity. Porcine deltacoronavirus NSP5 suppresses host antiviral IFN signaling by proteolysis of the host antiviral proteins NF-kappa-B essential modulator [83], STAT2 [84], and mRNA decapping protein 1a [85]. Previous computational studies also predicted a SARS-CoV-2 NSP5 cleavage site (VQMQ|AIPE) in HDAC2, with cleavage removing an 11.6 kDa C-terminal fragment containing the HDAC2 NLS [42]. This cleavage was proposed to reduce HDAC2 localization to the nucleus and thus limit its ability to attenuate inflammatory responses. Surprisingly, we did not observe evidence that NSP5 cleaves or inactivates HDAC2, with no cleaved HDAC2 (~44 kDa) appearing in our immunoblotting experiments, nor with any cleavage detected in our intramolecular FRET assay of either cytosolic or nuclear-localized HDAC2. This is consistent with the work of Naik et al. who observed a similar non-proteolytic interaction of NSP5 with HDAC2 and IRF3 [66]. Further demonstrating that cleavage is not required for this interaction, the catalytically inactive NSP5H141A and NSP5C145S point mutants maintained their interaction with HDAC2 and had the same suppressive effect on CIITA pI and MHC II promoter activity as wild-type NSP5. Why
HDAC2 is not cleaved by NSP5, despite containing an accessible SARS-CoV-2 NSP5 consensus sequence, is unclear. However, non-proteolytic functions of SARS-CoV-2 NSP5 have been reported, including inducing SUMOylation of MAVS to promote inflammation [86] and suppressing activation of RIG-I by preventing formation of antiviral stress granules via interactions with G3BP1 [87]. Additionally, coronavirus-mediated epigenetic reprogramming has been reported previously, with both SARS-CoV and MERS-CoV altering inhibitory and activating histone modifications across a range of genes [88]. This regulation is targeted, with these viruses able to selectively activate or repress expression of specific genes. Consistent with this observation, we found that SARS-CoV-2 NSP5 selectively inhibited expression of MHC II and CIITA without affecting expression of CD86 and RFX5, or globally suppressing acetylation.

HDAC2 is a promising target for treatment of SARS-CoV-2 infection. Indeed, epigenetic changes consistent with HDAC2 activity drives the cytokine storm that is responsible for much of the pathology seen in SARS-CoV-2 patients, with the degree of epigenetic change correlated to disease severity [45,89–91]. NSP5 is thought to drive much of the cytokine response in the infected lung via HDAC2, and indeed, HDAC2 is required for the upregulation of many pro-inflammatory molecules in endothelial and myeloid cells undergoing SARS-CoV-2 infection [92–94]. HDAC inhibitors exhibit potent anti-inflammatory effects and therefore may antagonize many of the inflammatory pathways activated by SARS-CoV-2 [95,96]. Outside of inflammation, HDACs positively regulate ACE2 expression, and consequently, HDAC2 inhibitors decrease SARS-CoV-2 entry and replication through reduced ACE2 expression [95,97,98]. Moreover, HDAC inhibitors downregulate pro-inflammatory cytokines, reduce lung fibrosis, prevent viral entry into the central nervous system, and decrease neurological damage [96,99,100]. Thus, HDAC2 inhibition may improve patient outcomes through multiple mechanisms in addition to restoration of antigen presentation on MHC II. Critically, targeting a host protein would reduce the likelihood of SARS-CoV-2 evolving resistance to this treatment.

Understanding how SARS-CoV-2 modulates the MHC II antigen presentation pathway provides an important insight into the immunoevasion tactics used by this virus and may help to provide directions for the design of future COVID-19 vaccines or therapeutics. This study identifies one mechanism through which SARS-CoV-2 suppresses adaptive immunity by antagonizing the MHC II antigen presentation pathway. Furthermore, our data indicate that SARS-CoV-2 may utilize NSP5 to modulate a broad array of immune responses via targeting HDAC2, which in addition to its effects on MHC II expression identified herein, has also been identified as a positive regulator of the cytokine storm that underlies much of the pathology of COVID-19. Indeed, HDAC2 inhibition has been proposed as a therapeutic approach for COVID-19, with the findings from this study further validating HDAC2 inhibitors as potentially valuable treatments for this disease [99].

### Materials & Methods

#### Materials

The TGN46-GFP and KDEL-mRFP plasmids were gifts from Dr. Sergio Grinstein (Hospital for Sick Children, Toronto, Canada), plVX-zsGreen lentiviral vector and pMD2.G and pDR8.2 packaging vectors were a gift from Dr. Jimmy Dikeakos (University of Western Ontario, London, Canada). pcDNA3-myc-CIITA (plasmid 14650) were purchased from AddGene. All DNA primers and synthesized genes were from IDT (Coralville, Iowa), and the sequences for all primers used in this study can be found in Supplemental Table S1. Tissue culture medium, fetal bovine serum (FBS), and trypsin were from Wisent (St. Bruno, Canada). Recombinant cytokines were from Peprotech (Cranbury, NJ). All cell lines and Lympholyte-poly were from Cedarlane labs (Burlington, Canada). Polybrene, ivermectin, and 100K Amicon centrifugal filters were purchased from EMD Millipore Corp (USA). CD14 Positive Cell Selection Kit, FcBlock, and anti-DYKDDDDK Tag (L5) were from BioLegend (San Diego, California). The #1.5 thickness coverslips and 16% paraformaldehyde (PFA) were from Electron Microscopy Sciences (Hatfield, PA). FDA-traceable PLA filament was purchased from Filaments.ca (Mississauga, Canada). RNeasy Mini Kit was from Qiagen (Germantown, MD), Permaflor, WGA-Alexa Fluor 647, DAPI, Hoescht, HALT protease/phosphatase inhibitors, Dithiobi[succinimidyl propionate] and dithio-bismaleimidoethane, were purchased from ThermoFisher Canada (Mississauga, Canada). The suppliers and all antibodies and labeling reagents used in this study can be found in Supplemental Table S2. Atto647N NT Labeling Kit was from Jena Biotech (Jena, Germany). Accell cell-penetrating SMARTpool scrambled and HDAC2-targeting siRNAs were from Horizon Discovery (Cambridge, UK). Instagene, iScript Select cDNA Synthesis Kit, 4%-20% SDS-PAGE gels, SsoFast EvaGreen Supermix, and all
protein blotting reagents/gels were from BioRad Canada (Mississauga, Canada). Renilla luciferase internal control vector pRL-TK and the dual luciferase reporter assay kit was from Promega (Madison, WI). Phusion PCR enzyme, all restriction enzymes, HiFi Gibson Assembly Kit, and T4 DNA ligase were from NEB Canada (Whitby, Canada). Retro-X Universal Packaging System was purchased from Takara Bio (San Jose, California). All lab plasticware, PolyJet and GenJet transfection reagent, and DNA isolation kits were from FroggaBio (Concord, Canada), and all laboratory chemicals were from BioShop Canada (Burlington, Canada).

**NSP5 Cloning and Retroviral Packaging**

The RNA sequence NSP5 from the founder SARS-CoV-2 strain [101] was synthesized such that a start and stop codon were added to the 5’ and 3’ end of the NSP5 sequence, along with 20 bp of homology to the pLVX-zsGreen vector at the EcoRI restriction site. The resulting NSP5 sequence was cloned into EcoRI-digested pLVX-zsGreen by Gibson assembly. Point mutants were generated by amplifying the entirety of this original vector with phosphorylated primers that incorporate the point mutation in the first base pair of the forward primer, while deletion mutants were generated by amplifying the vector from either side of the desired deletion with phosphorylated primers. After amplification, the parental plasmid was removed by DpnI digestion and the amplicons circularized with T4 DNA ligase. All primers used for RT-qPCR can be found in Supplemental Table S1. To produce pseudotyped lentivirus containing empty vector or NSP5, 3x10^6 HEK293T cells were grown in 75 cm^2 tissue culture flasks, then transfected with PolyJet transfection reagent (500 μL complex containing 4 μg pMD2.G and 10 μg pDR8.2 packaging vectors, and 10 μg of pLVX expression vector). Following transfection, cells were incubated at 37°C/5% CO_2 for 18 hr at which point the media was exchanged for 8 mL of DMEM supplemented with 10% FBS and returned to the incubator for 48 hr. The media was transferred to a sterile 50 mL conical centrifuge tube and topped up to 20% FBS, then centrifuged at 4000×g for 5 minutes, and the supernatant filtered with a 0.2 μm syringe filter into a new 50 mL conical tube. The pseudotyped lentivirus was then concentrated using a 100 kDa centrifugal filter unit at 4000×g at 4°C for 45 min per 15 mL of filtrate. Concentrated pseudotyped lentivirus was aliquoted and stored at -80°C and thawed at room temperature prior to use.

**Human Macrophage and Dendritic Cell Culture, Transduction, and siRNA Treatment**

The collection of blood and cells from healthy donors was approved by the Health Science Research Ethics Board of the University of Western Ontario and was performed in accordance with the guidelines of the Tri-Council policy statement on human research. Blood was drawn into heparinized vacuum collection tubes, layered on an equal volume of Lympholyte-poly and centrifuged at 300×g for 35 min at 20°C. The top band of peripheral blood mononuclear cells was collected and washed once (300×g, 6 min, 20°C) with phosphate-buffered saline. For dendritic cell differentiation, a CD14 selection kit was used to isolate monocytes according to manufacturer’s instruction. The selected CD14+ cells were cultured in RPMI-1640+10% FBS and 1% antibiotic-antimycotic solution with GM-CSF (100 ng/ml) and IL-4 (100 ng/ml) for 4 days yielding immature monocyte-derived DCs (moDCs). To produce macrophages, selected CD14+ cells were cultured in the presence of M-CSF (10 ng/ml) for 6 days. ~1 × 10^6 moDCs or macrophages were centrifuged with 20 transducing units of lentiviral vectors per cell at 800×g at 32°C for 90 min with Polybrene (10 μg/ml). After centrifugation, the cells were incubated at 37°C with 5% CO_2, and 8 hr later fresh media plus cytokines were added and the cells incubated for 72 hr. For siRNA knockdown, 1 μM of Accell cell-permeant siRNA was added to the cells and incubated at 37°C with 5% CO_2 for 72-96 hr.

**Cell Line Culture and Transfection**

HeLa, RAW264.7, and J774.2 were cultured in DMEM supplemented with 10% FBS, A549 cells were cultured in Ham’s F12 supplemented with 10% FBS and were grown at 37°C/5% CO_2 incubator. Cells were split 1:10 upon reaching >80% confluency by either scraping cells into suspension (RAW and J774) or by trypsinization, diluting in fresh medium, and replating in a new tissue culture flask. GenJet DNA transfection reagent was used to transfect plasmids into HeLa cells, as per the manufacturer’s instructions. Briefly, for each well in a 6-well plate, 1 μg of DNA was diluted into 50 μl of serum-free DMEM, followed by 3 μl of GenJet reagent. The resulting mixture was incubated for 10 min at room temperature, and then added dropwise to the HeLa cells. Cells were incubated for at least 18 hr at 37°C in a 5% CO_2 incubator. A549 cells were transfected using Lipofectamine 3000, transfecting 1 μg of DNA per well of a 12-well plate, using 2 μL of P2000 and 3 μL Lipofectamine per transfection. A Neon transfection system (Thermo Fisher Scientific AG) was used to transfect plasmids into RAW264.7 cells according to the manufacturer’s instructions. Briefly, 1×10^6 cells were resuspended in 10 μL of buffer R containing 5 μg of plasmid DNA and electroporated using a single 20 ms pulse at 1680V. Lentiviral transductions were used for transducing plasmids into J774.2 cells as described above.

**Flow Cytometry**

HLA-DR and CD86 expression on the surface of moDCs was measured following 72 hrs transduction with NSP5-ZsGreen or empty-ZsGreen pseudotyped lentivirus and subsequent 24 hr stimulation with 100 ng/μL IFN-γ and incubated at 37°C/5% CO_2. After stimulation 3×10^5 cells per condition were washed with PBS and blocked for 30 min on ice with FcBlock. Cells were stained on ice for 30 min using eFluor780-FVD and conjugated primary antibodies as indicated in Supplemental Table S2. Cells were fixed with 4% PFA in PBS for 15 min then washed with PBS. Expression levels were measured using a FACScanTo (BD), live moDCs were identified based on FVD-eFluor780 viability dye staining and forward scatter and side scatter profiles. Singlets were gated on the forward area scatter and forward height scatter profiles (Figure S1). For cell sorting, singlets were gated on the forward area scatter and forward height scatter profiles, then transduced cells were identified by a positive zsGreen signal and this population sorted into the receiving tube. Flow cytometry data were analyzed using FlowJo (v10.8). All antibodies, dyes, and dilutions used for flow cytometry can be found in Supplemental Table S2.
**Immunoprecipitation and Immunoblotting**

Prior to lysis, cells were washed 3x with cold PBS. For immunoprecipitations, proteins were reversibly cross-linked using the ReCLIP method [102]. Briefly, cells were incubated for 1 hr at room temperature in PBS + 0.5 mM dithiobis[succinimidy] propionate and 0.5 mM dithio-bismaleimidoethane). This medium was aspirated, and crosslinking quenched by the addition of 5 mM L-cysteine in 20 mM Tris-Cl, pH 7.4 for 10 min at room temperature. Cells were suspended in 300 µL of RIPA lysis buffer, and 50 µL pre-washed anti-DYKDDDDK Tag (L5) beads, rotating for 1 hr. Beads were washed with PBS and immunoprecipitated protein eluted using 0.1 M glycine at pH 2.8. For immunoblotting, cells were lysed with 300 µL RIPA buffer supplemented with 1 mM PMSF and Halt protease and phosphatase inhibitor cocktail at the manufacturer’s recommended concentration. Proteins were loaded on a 4–15% gradient SDS-PAGE gels and transferred onto PVDF membrane. The membrane was blocked for 5 minutes with EveryBlot Blocking Buffer (BioRad), incubated overnight at 4°C with the desired primary antibodies (Supplemental Table 2), washed 3 x 5 min with TBS-T, incubated with appropriate IR700 or IR800 secondary antibodies, 1:2,500 dilution, for 1 hr at room temperature in TBS-T. The blots were washed 3 x 15 min washes in TBS-T and visualized with an Odyssey CLx (LI-COR Biosciences, Lincoln, Nebraska). Densitometry was performed in ImageJ/FIJI [103,104].

**RT-qPCR**

Total RNA was isolated from FACS sorted cells transduced with either empty or NSP5-expressing lentiviral vectors using RNeasy Mini Kit as per manufacturer’s instructions. Samples were eluted in 30-50 µL of RNase-free water. RNA concentration and quality were measured using a NanoDrop 1000 Spectrophotometer. CDNA was obtained from total RNA using the iScript Select cDNA Synthesis Kit according to manufacturer’s instructions using an equal amount of starting RNA and equal mix of the oligo (dT)20 primer mixes. RT-qPCR was performed using SsoFast EvaGreen Supermix with an equal amount of starting cDNA. Reactions were run on a QuantStudio 3 Real-Time PCR System for 40 amplification cycles. Relative expression of genes of interest was calculated using the ΔΔCt method, with GAPDH serving as the reference gene.

**Dual-Luciferase Promoter Activity Assay**

RAW264.7 cells were seeded at 1 x 10⁶ cells/well in a 12-well plate and transfected as indicated with Luc-HLA-DRA, Luc-CIITA pI/, Luc-CIITA pII/IV, Renilla luciferase internal control vector pRL-TK, and NSP5-FLAG or pLVX-IRES-ZsGreen using the Neon electroporation system according to the manufacturer’s protocol. 72 hr post-transfection, cells were lysed with 1x Passive lysis buffer supplemented with EDTA-free HALT protease inhibitor at the manufacturer’s recommended concentration. Dual luciferase assays were performed using a Dual-Luciferase Reporter Assay Kit according to the manufacturer’s instructions, with measurements performed on a Cytation 5 luminescence microplate reader. Firefly luciferase readings were relative to Renilla luciferase readings to account for differences in transfection efficiencies and cell count between samples.

**Immunofluorescence Microscopy**

Cells of interest were seeded at a density of 1000 cells/mm² into either 18 mm circular coverslips placed into the wells of a 12-well plate, or into the wells of a custom-printed 15-well imaging chamber [105]. Cells were fixed in 4% paraformaldehyde in PBS for 20 min at room temperature. For plasma membrane staining, cells were stained with 5 µg/mL Alexa Fluor 647-conjugated wheat germ agglutinin for 10 min at 10 °C, then fixed in 4% paraformaldehyde in PEM buffer (80 mM PIPES, 1 mM EGTA, 1 mM MgCl₂) for 10 min at 37 °C. If permeabilization was required, fixed cells were treated with permeabilization buffer (PBS + 0.1% Triton X-100 + 2.5% BSA); otherwise, cells were blocked with antibody buffer (2.5% BSA in PBS). Anti-FLAG, -MHC II, or -acetyl-lysine were diluted to the concentration indicated in Supplemental Table S1 in antibody buffer and incubated with the cells for 1 hr. Cells were then washed 3 x 15 min with PBS, and then an appropriate secondary antibody added at a 1:1,000 to 1:2,500 dilution in antibody buffer for 1 hr, followed by washing 3 x 15 min with PBS. Samples were either imaged immediately or mounted on a slide using Permafluor before imaging. All incubations and washes were performed at room temperature.

All samples were imaged on a Leica DMi6000B equipped with a Hamamatsu ORCA-Flash4 CMOS camera, fast filter wheels equipped with a Chroma Sedat Quad and custom Fluorescent Resonance Energy Transfer (FRET) filter wheels, operated using Leica LAS-X software. Unless otherwise noted, all cells were imaged using a 100x/1.40 NA objective lens, with Z-stacks acquired with 0.4 µm between slices. Z-stacks were deconvolved in LAS-X using a 10-iteration blinded deconvolution. Images were exported to ImageJ/FIJI for analysis [103,104]. For co-localization studies, the JaCoP plugin was used to calculate the Manders Ratio of NSP5-FLAG and nuclear DAPI or Hoechst staining, or transgene-delineated ER or Golgi markers [106]. To calculate the fraction of NSP5 in the nucleus, a manual region of interest (ROI) was drawn around the nucleus and whole cell and the integrated intensity of each was measured. The fraction of NSP5 in the nucleus was calculated as Nuc/Total NSP5 FRET and FISH-FRET were quantified as described below. To calculate the mean fluorescence intensity of MHC-II in transduced macrophages, the background subtracted channel was thresholded to create a binary mask using the default setting in ImageJ, then a sum slices Z-projection was created. A manual ROI was drawn around the whole cell and the integrated intensity was measured. To determine the surface to cytosol ratio of MHC II in transduced macrophages, the background subtracted channels for wheat germ agglutinin and MHC II were thresholded as described above. Then, the image calculator was used to display all overlapping and non-overlapping pixels, representing MHC II on the cell surface and cytosol, respectively. A summed Z-projection was created, and a manual ROI was drawn around the whole cell to measure the integrated density for both surface- and cytosolic-MHC II. The surface-to-cytosol ratio was calculated as Surface MHC II and the fraction of total MHC II on the membrane was calculated as Cytosolic MHC II.
**HDAC2 Intramolecular FRET**

DNA comprised of the human HDAC2 gene with flanking BglII and BamHI restriction sites was synthesized and cloned into pmVenus (L68V)-mTurquoise2 (AddGene #60493) such that a fusion protein of mVenus-HDAC2-mTurquoise2 was produced. HeLa cells were then transfected with this construct with or without NSP5, with mTurquoise2 alone (donor-only sample), with mVenus alone (acceptor-only sample), or with the pmVenus (L68V)-mTurquoise2 vector (positive control). Tiled images of the each well were collected, acquiring the donor (mTurquoise2), acceptor (mVenus) and FRET channels at 40× magnification, using the same excitation and camera settings across all samples. FRET efficiency was then calculated using an implementation of the approach of van Rohenen et al. [107] using a custom-written script in FIJI. In each repeat, the correction values for donor cross-talk (β, donor-only Ida/Idd), donor cross-excitation (α, acceptor-only Idd/laa), acceptor cross-excitation (γ, acceptor-only Ida/laa), and FRET cross-talk (δ, acceptor-only Idd/Ida), were calculated using donor-only or acceptor-only images and custom-written scripts in FIJI. FRET efficiency (E) was then calculated in background subtracted images using the formula:

\[
E(i) = \frac{I_{dd} - \beta I_{dd} - (I_{aa} \times (\gamma - \alpha \beta))}{I_{Idd}(1 - \beta \delta)}
\]

In Image/FIJI the acceptor-only image was then thresholded, and the “Analyze particles” feature used to generate separate ROIs for each cell in each image, and these ROIs were used to quantify the FRET signal of each cell.

**FISH-FRET**

To measure levels of acetyl-lysine at the MHC II and CIITA promoters, human DNA was purified from a cheek swab using Instagene as per the manufacturer’s instructions. 3500 bp amplicons starting before the promoter and ending at the end of the first exon were amplified with Phusion DNA polymerase as per the manufacturer’s instructions using the primers from Supplemental Table S1. Amplicons were gel purified and cloned into EcoRV-digested pBluescript II using a HiFi Assembly Kit as per the manufacturer’s instructions. The resulting plasmids were labeled with ATTO647N and fragmented using a ATTO647N NT Labeling Kit, producing fragments averaging 200 nucleotides. Primary human macrophages or A549 cells were plated into the 7.5 mm wells of a customized imaging chamber [105], transduced with lentiviral vectors (macrophages) or transfected (A549 cells) with NSP5 or an empty vector, and treated with siRNA as per the protocol of Ye et al. [108], including wells which were left unstained, or stained only with the donor (acetyl-lysine-Cy3) or acceptor (FISH probes). Briefly, cells were fixed, permeabilized and immunostained for acetyl-lysine as described above. After labeling a secondary fixation was performed for 10 min with 2% PFA. FISH probes diluted 1:2,500 in hybridization solution (50% formamide, 10% dextran sulfate, 0.3 M NaCl, 30 mM sodium citrate) and denatured at 75°C for 10 min and then cooled to 37°C. Simultaneously, the cells were incubated at 70°C for 2 min in 70% formamide, 0.3 M NaCl, and 30 mM sodium citrate. The cells were dehydrated by immersing in 75%, 90% and 100% ethanol, 2 min/immersion, then air-dried. The cells were incubated with the denatured FISH probes overnight at 37°C, washed 3 × 5 min with 50% formamide, 0.3 M NaCl, and 30 mM sodium citrate at 42°C, then washed 3 × 5 min with 0.05% Tween 20 in 0.6 M NaCl, and 60 mM sodium citrate. The cells were then washed 3 × 5 min in PBS and immediately imaged.

Tiled images of each well were collected, acquiring the zsGreen, donor, acceptor, and FRET channels at 40× magnification, using the same excitation and camera settings across all samples. FRET efficiency was then calculated as described above. A trained algorithm in ilastik [109] was used to identify cells based on the acetyl-lysine staining and to classify each cell as zsGreen+ (transduced) or zsGreen- (untransduced). The resulting classifications were exported to FIJI where they were used to assign each FISH probe in the image and the corresponding FRET signal to transduced or untransduced groups. The FRET signal in each sample was then normalized to that observed in the scrambled siRNA-treated, zsGreen+ nuclei.

**NSP5 NLS Analysis**

The protein sequence of NSP5 was analyzed for the presence of monotonic and bipartite nuclear localization signals using the default settings on four different prediction algorithms: cNLS Mapper (https://nls-mapper.iab.keio.ac.jp/cgi-bin/NLS_Mapper_form.cgi), 4 state HMM on NLSstradamus (http://www.moseslab.csb.utoronto.ca/NLSstradamus/), seqNLS (http://mleg.cse.sc.edu/seqNLS/), and NLStradamus in InterProScan (https://www.ebi.ac.uk/interpro/search/sequence/).

**Phylogenetic Analysis**

Using the protein sequence of human HDAC2 and the NCBI BLASTp tool (https://blast.ncbi.nlm.nih.gov), the protein sequences of HDAC2 from a range of species representing the major vertebrate clades were identified. The same approach, using the protein sequence of NSP5 from SARS-CoV-2 was used to identify NSP5 protein sequences across the four coronavirus genera. These sequences were imported into MEGA XI, and a MUSCLE alignment of the protein sequences was generated [110]. Pairwise distances were then calculated using a Poisson model assuming uniform rates across sites, and maximum likelihood trees were generated using a 500-iteration bootstrapping approach. Per-residue conservation was quantified using the Shannon Entropy calculator on the Protein Residue Conservation Prediction server (https://compbio.cs.princeton.edu/conservation/) [111].

**Statistical Analysis**

Using GraphPad Prism, a Shapiro-Wilk test was used to determine whether data was parametrically or non-parametrically distributed, and data was then analyzed using an appropriate 2-tailed statistical test, as indicated in the figure legends. Parametric data is presented as mean ± SEM, while non-parametric data is presented as box-and-whisker or violin plots with median and quartiles.
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Competing Interest Statement

The authors have no competing interests.

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