Phosphoproteomics to Characterize Host Response During Influenza A Virus Infection of Human Macrophages*§

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Influenza A viruses cause infections in the human respiratory tract and give rise to annual seasonal outbreaks, as well as more rarely dreaded pandemics. Influenza A viruses become quickly resistant to the virus-directed antiviral treatments, which are the current main treatment options. A promising alternative approach is to target host cell factors that are exploited by influenza viruses. To this end, we characterized the phosphoproteome of influenza A virus infected primary human macrophages to elucidate the intracellular signaling pathways and critical host factors activated upon influenza infection. We identified 1675 phosphoproteins, 4004 phosphopeptides and 4146 nonredundant phosphosites. The phosphorylation of 1113 proteins (66%) was regulated upon infection, highlighting the importance of such global phosphoproteomic profiling in primary cells. Notably, 285 of the identified phosphorylation sites have not been previously described in publicly available phosphorylation databases, despite many published large-scale phosphoproteome studies using human and mouse cell lines. Systematic bioinformatics analysis of the phosphoproteome data indicated that the phosphorylation of proteins involved in the ubiquitin/proteasome pathway (such as TRIM22 and TRIM25) and antiviral responses (such as MAVS) changed in infected macrophages. Proteins known to play roles in small GTPase-, mitogen-activated protein kinase-, and cyclin-dependent kinase- signaling were also regulated by phosphorylation upon infection. In particular, the influenza infection had a major influence on the phosphorylation profiles of a large number of cyclin-dependent kinase substrates. Functional studies using cyclin-dependent kinase inhibitors showed that the cyclin-dependent kinase activity is required for efficient viral replication and for activation of the host antiviral responses. In addition, we show that cyclin-dependent kinase inhibitors protect IAV-infected mice from death. In conclusion, we provide the first comprehensive phosphoproteome characterization of influenza A virus infection in primary human macrophages, and provide evidence that cyclin-dependent kinases represent potential therapeutic targets for more effective treatment of influenza infections. Molecular & Cellular Proteomics 15: 10.1074/mcp.M116.057984, 3203–3219, 2016.

Influenza A viruses (IAVs)1 cause annual epidemics and occasionally worldwide pandemics, infecting millions of people. IAVs are negative-stranded RNA viruses belonging to Orthomyxoviridae family. The options to control the spread of human influenza are vaccination and antivirals. The current antiviral drugs used for treatment of IAV infections target the viral proteins. The influenza viruses mutate rapidly to escape host immune responses developed after previous infection or vaccinations (1) and these high mutation rates lower the effi-
cacy of antiviral treatment that target viral proteins (2, 3). To
tackle the resistance problem, it is crucial to find novel
cytoxic drugs targeting conserved virus-host interac-
tions, which are critical for IAV infection and replication. In-
hibiting such interactions provide improved opportunities to
reduce the occurrence of drug-resistant mutants and to min-
imize the morbidity and mortality caused by IAVs.

The host response to pathogens is complex and dynamic.
Proteomics can shed light on this response and provide the
system-level information of the immune response required for
detailed understanding of disease mechanisms (4, 5). Mass
spectrometry (MS)-based proteomics has become a powerful
tool for analyzing protein abundances, modifications and in-
teractions. The current MS technology allows identification,
quantification, and characterization of post-translational mod-
ifications of thousands of proteins in a single experiment. In
biomedicine, phosphoproteins are intriguing candidates for
therapeutic interventions because the upstream kinases are
relatively easy to target selectively with small molecule drugs
(6). Previous MS-based phosphoproteome studies have
shown major effects on host protein phosphorylation and reor-
ganization of signal-transduction pathways in viral infec-
tions such as HIV-1 (7), lytic γ herpesvirus infection (8), por-
cine reproductive and respiratory syndrome virus (9), rift valley
virus (10), Sendai virus (11), and human cytomegalovirus (12)
infections.

Macrophages are professional phagocytes and one of the
first cells to encounter pathogens in tissues. Macrophages
often reside in or are recruited to infection sites, and they are
important generators of inflammatory responses (13). Macro-
phages are also one of the most abundant phagocytes in the
respiratory tract tissue and lung, and they are therefore
thought to play a key role in the control and clearance of IAV
infections (14). We previously published a subcellular pro-
teome and secretome characterization of IAV infected primary
human macrophages, which demonstrated that viral infection
perturbs the expression and subcellular localization of more
than a thousand host proteins at early phases of infection (15).
Here, we have characterized the phosphoproteome of IAV
infected human macrophages in order to identify the host
kinases and their substrates that play a critical role in the early
stages of infection. We identified 1675 human phosphopro-
teins and in total 1113 proteins showed changes in their
phosphorylation status upon IAV infection, implying that the
IAV infection has a profound effect on host protein phosphor-
ylation. These host proteins are involved in all critical stages of
virus life cycle including viral entry, gene regulation, and
gress. The data also showed that IAV infection influences the
phosphorylation status of several cyclin-dependent kinases
(CDKs) and their substrates. Functional studies showed that
the CDK activity is required for efficient viral replication and for
the activation of antiviral and pro-inflammatory responses.

**EXPERIMENTAL PROCEDURES**

**Ethics Statement**—All mouse experiments were conducted in strict
accordance with national (Belgian Law 14/08/1986 and 22/12/2003,
Belgian Royal Decree 06/04/2010) and European (EU Directives 2010/
63/EU, 86/609/EEG) animal regulations. Animal protocols were
approved by the ethics committee of Ghent University (permit number
LA1400091, approval ID 2010/001). All human blood donors provided
written informed consent.

**Cells and Cell Stimulation**—Primary human macrophages were
derived from leukocyte-rich buffy coats from healthy blood donors
(Finnish Red Cross Blood Transfusion Service, Helsinki, Finland).
Monocytes were isolated and differentiated into macrophages as
described previously (16). In total 1.4 × 10⁶ monocytes were seeded
per well on 6-well plates or 5 × 10⁴ per well on 96-well plates.
The monocytes were cultured in serum free macrophage media (Gibco®,
Thermo Fisher Scientific, Waltham, MA) supplemented with 10 ng/ml
granulocyte-macrophage colony-stimulating factor (GM-CSF) (BIO-
SOURCE International, Thermo Fisher Scientific) and 50 U/ml peni-
cillin-streptomycin (Lonza, Basel, Switzerland) at 37 °C and 5% CO₂
for 7 days, polarizing the monocytes into macrophages of the acute
pro-inflammatory M1-phenotype. Before stimulation, the media was
replaced with fresh GM-CSF free macrophage media and macro-
phages were infected with influenza A virus. MDCK cells (Sigma-
Aldrich, St Louis, MO, catalog number B412190) were maintained at
37 °C and 5% CO₂ in DMEM (Sigma-Aldrich) supplemented with 10%
fetal bovine serum (Gibco®, Thermo Fisher Scientific), 2 mM L-gluta-
mine (Lonza), and 50 U/ml penicillin-streptomycin (Lonza).

**Virus Stocks and Infections**—Human pathogenic influenza
A/Urdu307/1972 (H3N2), provided by the National Institute for
Health and Welfare, was cultured in embryonated hen eggs and the
virus stock, with a titer of 256 hemagglutination U/ml, was stored at
−80 °C. A virus dose of 2.56 hemagglutination U/ml was used in the
infection experiments unless stated otherwise. The virus experiments
were carried out under BSL-2 conditions and in compliance with
regulations of the University of Helsinki (permit No 21/1/M/09). The mice
studies were conducted with a mouse-adapted influenza A virus,
H1N1 A/Puerto Rico/8/34 (PR8) (17), obtained as a kind gift from Dr.
Skehel. The PR8 virus was adapted to mice by serial lung passage
and propagated in MDCK cells in the presence of TPCK-treated
trypsin.

**Phosphoproteomics and Bioinformatics**—Macrophages grown
on six-well plates were used for the phosphoproteomics experiments.
The cell growth media was changed to GM-CSF free growth media (1
ml/well) one hour before infection with a virus dose of 2.56 hemag-
glutination U/ml. The macrophages were left untreated (control) or
infected with IAV for 6 h. The cell media was removed and the cells
were washed with PBS, and then incubated in PBS on ice for 5 min,
after which they were scraped off with a rubber policeman. The cell
solution was centrifuged 404 × g for 5 min. The supernatant was
removed and the cell pellet was re-suspended in 10 ml of ice-cold
PBS and the centrifugation step was repeated. The cells were lysed
with HEPES lysis buffer (50 mM HEPES, 150 mM NaCl, 1 mM EDTA,
1% Nonidet P-40, pH 7.4, including protease and phosphatase in-
hibitor cocktails from Sigma-Aldrich), the lysates were centrifuged,
and the protein content was measured with Bio-Rad DC™ (Bio-
Rad Laboratories, Hercules, CA). The digestion of the proteins in the
cell lysates (10 mg per sample), the clean-up, desalting steps, strong
cation exchange fractionation, and phosphopeptide enrichment with
IMAC were performed as described previously (18). The LC-MS/MS
analysis was done with a Q Exactive ESI-quadrupole-orbitrap mass
spectrometer coupled to an EASY-nLC 1000 nanoflow LC (Thermo
Fisher Scientific) as described previously (19). Two biological repli-
cates were analyzed (in total two control and two IAV infected sam-

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samples, both replicates including equal amounts of cells from four different donors.

Raw files were processed in the Proteome Discoverer interface (version 1.3, Thermo Fisher Scientific, SanJose, CA) and the MS/MS spectra were searched using the Mascot algorithm (Matrix Science, version 2.4.0) against a concatenated forward-reverse SwissProt database (release date April 2012, Homo sapiens) supplemented with common contaminants (total of 40,678 protein sequences) to identify human proteins. To identify IAV proteins, the LC-MS/MS data was searched against a SwissProt database with IAV and human protein sequences (version 02/2012, total of 21,545 sequences). Search parameters included trypsin specificity with allowance of maximum two missed cleavage sites. The peptide charge was specified as 2+ and 3+, the precursor mass tolerance as 5 ppm, and the MS/MS ion tolerance as 0.2 Da. Carbamidomethylation (C) was specified as fixed modification, and oxidation (M), phosphorylation (S, T, and Y) and acetylation (protein N-terminal) as variable modifications. The mass spectrometry proteomics data have been deposited at the ProteomeXchange Consortium (20) via the PRIDE partner repository (21).

To identify IAV proteins, the LC-MS/MS data was submitted to the PhosFox (22) analysis with the in-house developed phosphoproteomic data analysis tool. The results from two biological replicates were combined when performing the PhosFox analysis. In addition, PhosFox considered a phosphopeptide with different number of missed cleavages as single phosphopeptide. A phosphopeptide was considered “unique” to a sample if PhosFox detected it only in one sample (e.g. infected), but not in the other sample (e.g. control). The proteins with phosphopeptides that were unique to control and infected samples were chosen for bioinformatics analyses. Pathway analysis and classification of proteins based on gene ontology annotations were carried out with Ingenuity Pathway Analysis (IPA) (Qiagen, Hilden, Germany) and InnateDB database (23). The NetworKIN algorithm (24) in KinomeXplorer was used for predicting kinases associated with specified phosphosites and Kinase Enrichment Analysis (KEA) (25) for identifying kinase substrates in the phosphoproteomic data.

**Compound Efficacy Testing In Vitro and VirusTitration**—SNS-032, flavipoviridol, dinaciclib, rosvocinide, and palbociclib (all from Selleckchem, Houston, TX) were dissolved in DMSO (Sigma-Aldrich) to 10 mM stock solutions and stored at -80 °C. DMSO was used as control in the experiments. The compound efficacy testing was performed in 96-well plates with macrophages at 95% confluence. The compounds were added to the medium and one hour later the cells were infected with virus or noninfected (mock). The cell viability was analyzed with the Cell Titer Glo assay (CTG; Promega, Madison, WI) at 24 h post-infection. The luminescence was read with a PHERaster FS plate reader (BMG Labtech, Thermo Fisher Scientific).

The antiviral efficacies of the CDKIs were validated by the plaque assay as described previously (26). The macrophages were non- or compound-treated and infected with IAV (MOI of 0.01). The virus titers were determined by calculating the plaque forming units (PFU) (clear spots) for each sample and expressed as PFU/ml.

**Western Blotting**—Equal amounts of protein from total cell lysis samples were dissolved in Laemmli buffer and resolved by SDS-PAGE (Bio-Rad). The proteins were transferred onto Immobilon-P membranes (Millipore, Billerica, MA). The membranes were blocked with 5% nonfat milk or 5% BSA (Sigma-Aldrich) in TBST, stained with different primary antibodies overnight, followed by secondary antibody labeling and detection by the ImageQuant LAS 4000 mini Luminescent Image Analyzer (GE Healthcare, Little Chalfont, UK). The primary antibodies used in this study were anti-phospho(Ser396)-IRF3 (4D4G) (Cell Signaling, Danvers, MA, #4947), anti-IRF3 (FL-425) (Santa Cruz, Heidelberg, Germany, sc-9082), anti-iIκBα (4D4) (Cell Signaling, #4812), anti-Bcl-XL (Santa Cruz, sc-56021), anti-cleaved caspase 3 (Asp175) (Cell Signaling #9661). These antibodies were diluted 1:1000 in 5% BSA-TBST or in 5% milk-PBST (the IRF3 antibody). Anti-IAV (H3N2) NS1 and NP antibodies (1:5000 dilution in 5% BSA-TBST) have been previously described (16, 27). To confirm equal sample loading the membranes were stripped and labeled with anti-GAPDH (O141) (Santa Cruz, sc-47724), 1:1000 dilution in 5% milk-PBST.

**RT-PCR**—Total cellular RNA was isolated with the RNeasy Plus Mini Kit (Qiagen) and the RNA was reverse transcribed using the High Capacity cDNA Reverse Transcription kit (Applied Biosystems, Carlsbad, CA), according to the manufacturer’s instructions. RT-qPCR was run with an ABI PRISM 7500 Sequence Detection System, applying TaqMan chemistry and Predesigned TaqMan assay primers and probes (Applied Biosystems) and TaqMan® Fast Advanced Master Mix (Applied Biosystems). The following TaqMan probes were used: 18S: Hs01119599_g1_s1, β-actin: Hs01063069_g1, IL-6: Hs00600149_g1, and IL-10: Hs00985635_g1. The following primers and probes were used: IFN-α: Hs01119599_g1_s1, and IL-29 (IFN-α): Hs00601677_g1, CXCL10 (IP-10): Hs01124261_g1, CXCL11 (I-αC): Hs01417682_g1. Human GAPDH (forward primer: ggctggggctcatttg-cttaaa, reverse primer: ttgatctcccaacatgttagc-gg, probe: 5′-VIC-CCTGAGGAGAACATTCTTACCA-3′); human IκBα (forward primer: ccaactgcagagtagttgtg, reverse primer: gatctcagttctcctgctg, probe: 5′-FAM-TACACTGCTCAGGAGAGAATTTCGCT-3′). The human cytokine Luminex Bio-Plex Pro arrays were probed with IAV or mock. The cell growth media were collected after 18 h of stimulation and centrifuged. The human cytokine Luminex Bio-Plex Pro samples were analyzed routinely as negative controls.

**Cytokine Profiling, Luminex Assays and Caspase Activation Assay**—The cytokine arrays Proteome Profiler Human Cytokine Array Panel A kit (R&D Systems, Minneapolis, MN) were used for profiling cytokines according to the manufacturer’s instructions. For the human macrophage in vitro experiments, the concentration of SNS-032 and flavipoviridol was 0.3 μM and the cell culture supernatants were collected after 20 h of stimulation. The membranes were exposed to x-ray films and the films were scanned, or the chemiluminescence was measured with ImageQuant LAS 4000 mini (GE Healthcare, Uppsala, Sweden) and the fold change was calculated in comparison to the mock sample. The images were analyzed with ImageJ software (NIH).

For Luminex assay, primary human macrophages were pretreated with SNS-032 and flavipoviridol (0.3 μM) or left untreated and infected with IAV or mock. The cell growth media were collected after 18 h of stimulation and centrifuged. The human cytokine Luminex Bio-Plex Pro
Phosphoproteomic Analysis of Influenza Infection

Pro immunoassay (Bio-Rad Laboratories) designed to detect chemokines TNF, IL-1α, IL-1β, and IL-18 was performed according to the manufacturer’s instructions using the Bio-Plex 200 system hardware and version 4.1.1 of the Bio-Plex 200 software.

Caspase 3/7, 8 and 9 activities in infected or noninfected macrophages, nontreated or treated with SNS-032 or flavopiridol at 0.3 μM, were measured with the Caspase-Glo® 3/7. -8 and -9 assays (Promega) at 18 h. The Caspase-Glo® 1 Inflammasone assay (Promega) was used to measure the caspase 1 activity in infected or noninfected macrophages, nontreated or treated with SNS-032 at 0.3 μM at 18 h postinfected.

In Vivo Experiments—Pathogen-free 7-week-old female BALB/c mice were purchased from Charles Rivers Laboratories (Italy). The animals were housed in a SPF temperature-controlled environment with 12 h light/dark cycles and given water and food ad libitum. After 1-week adaptation in the animal room, six mice per group were treated with SNS-032 (15 mg/kg in 5% DMSO in 200 μl PBS) or vehicle (5% DMSO in 200 μl PBS) IP at 1 day before and 1, 3, 5, 7, and 9 days after infection. At day 0 the mice were sedated with isoflurane and challenged intranasally with 0.5 LD50 of mouse-adapted PR8. Body weight was monitored daily and mice that had lost more than 25% of their initial body weight were euthanized by cervical dislocation. The survival data of the mouse experiments were analyzed using SigmaStat 11.0 and the Log-rank (Mantel-Cox) test. The bodyweight changes were analyzed by two-way ANOVA test with Bonferroni’s multiple comparison adjustment.

In the follow-up experiment, four mice per group were treated with SNS-032 (15 mg/kg in 5% DMSO in 200 μl PBS) or vehicle (5% DMSO in 200 μl PBS) IP at 1 day prior infection and 1 and 3 days after infection. At day 0 the mice were sedated with isoflurane and challenged intranasally with 0.5 LD50 of mouse-adapted PR8. Mice were sacrificed at 5 days post infection and the lung and liver were collected and homogenized in 1.5 ml of PBS using a Heidolph RZR 2020 homogenizer. The homogenate was cleared using centrifugation at 1000 × g for 15 min at 4 °C. In total 200 μl of lung homogenate from four groups (nontreated/DMSO-treated, nontreated/SNS-032-treated, infected/DMSO-treated, and infected/SNS-032-treated) of four mice each were combined and assayed for cytokines with the Mouse cytokine array panel A (R&D Systems).

Mouse influenza virus infection experiments were carried out under BSL-2+ conditions and were authorised by the Institutional Ethics Committee on Experimental Animals of the Department for Medical Molecular Biology at Ghent University.

Experimental Design and Statistical Rationale—Data are shown as mean values and error bars represent standard deviation (S.D.) from the number of independent assays indicated in the figure legends. For two-group comparisons, a two-tailed unpaired t test was applied using Prism software (Graph-Pad Software, Inc.). The significance level is indicated as ***p < 0.001; **p < 0.01; *p < 0.05; p > 0.05 not significant (ns).

With phosphoproteomics, two independent biological replicate experiments were performed and the results from the replicates were combined when analyzing the data with the PhosFox software (22). See subsection Phosphoproteomics and Bioinformatics for more details.

RESULTS AND DISCUSSION

Phosphoproteome Analysis Shows Major Changes in Host Protein Phosphorylation After Influenza A Virus Infection of Human Macrophages—In order to get an integrated view of phosphorylation-regulated host proteins and signaling pathways by IAV infection in human macrophages, we used high resolution MS-based phosphoproteome analysis of control and infected samples combined with bioinformatics and functional assays. First, we studied the kinetics of viral replication and host response of human macrophages infected with influenza A/Udorn/307/1972 (H3N2) virus. The expression of viral mRNAs coding for matrix protein 2 (M2), nucleoprotein (NP), and RNA polymerase subunit 1 (PB1) was initiated at 4 h postinfection (supplemental Fig. S1). In accordance with the mRNA expression data, Western blot analysis demonstrated that viral proteins NP and nonstructural protein 1 (NS1) became detectable at 5−6 h after IAV infection (Fig 1A). Expression of antiviral cytokines IFN-β1 and IL-29 (IFN-λ1) mRNA started already 1−2 h after infection, reaching high levels of expression at 6 h postinfection (supplemental Fig. S1).

The IRF3 and NF-κB signaling pathways are activated during viral infection leading to the expression of inflammatory genes including type I (IFN-α/β) and III interferons (IL-28 and IL-29), proinflammatory cytokines, and chemokines (29, 30). The IRF3 signaling pathway is also involved in the activation of apoptosis during viral infections (31, 32). To study the activation of these transcription factors during IAV infection, we performed Western blot analyses to probe for phosphorylation of IRF3 and degradation of IκBα, an inhibitor of NF-κB. Phosphorylation of IRF3 was weakly initiated at 2 h postinfection and strong activation of IRF3 was seen after 6 h of infection (Fig 1A). No change in the total amount of IRF3 was detected during the infection progression. Degradation of IκBα was first seen at 2 h after IAV infection and at 6 h postinfection IκBα was barely detectable, indicating the activation of NF-κB. These results demonstrate that there is a weak activation of IRF3 and NF-κB pathways at early stages of infection, 1−2 h after viral entry. This is followed by a robust activation of these signaling pathways at 6 h after infection, which correlates with the initiation of viral RNA transcription and translation.

Based on these experiments, we chose an infection time point of 6 h for the high resolution MS-based phosphoproteomics analysis (Fig 1B). Control and IAV-infected macrophages were lysed and the protein samples were digested into peptides followed by peptide fractionation and phosphopeptide enrichment before LC-MS/MS analysis of the peptides. For phosphoprotein and -peptide identification, the LC-MS/MS data was first processed with the Mascot search engine, and the final list of phosphopeptides was generated using an in-house-developed data analysis tool, PhosFox (22). We analyzed two independent biological replicates, which led to the identification of 4004 phosphopeptides and 4146 nonredundant phosphorylation sites, originating from 1675 different human phosphoproteins (Fig 1C, supplemental Tables S1 and S2). In total, 1113 human proteins showed qualitative changes in their phosphorylation profiles upon infection (Fig 1C), meaning that specific Y/S/T sites were identified as phosphorylated only in control sample or after infection (supplemental Table S3). Approximately 30% of all the phosphopeptides in IAV-infected and control samples were
identified in both replicates (supplemental Fig. S2 and Table S4), which is comparable to phosphoproteomic data sets that we published previously (11, 18). More detailed comparison of the phosphopeptide and -protein identification results revealed that altogether 507 human proteins showed reproducible qualitative changes in their phosphorylation profiles upon IAV infection (supplemental Fig. S2 and Table S4). From all of the identifications, 285 phosphorylation sites in 222 different proteins have not previously been reported in the PhosphoSitePlus (33), Phosida (34), or UniProt (http://www.uniprot.org/) databases.
and were therefore considered as “novel” phosphosites. The proportion of phosphoamino acids in the data set unique to control was ~86% pSer, 14% pThr, and 1% pTyr, whereas in the data set unique to infection the proportion of phosphorylated threonines was higher (approx. 80% pSer, 20% pThr, and 1% pTyr) (Fig 1C).

The LC-MS/MS analysis identified IAV proteins exclusively in the infected samples, and only one virus phosphopeptide, belonging to the NS1 protein, was identified with high confidence (Fig. 1C). This amino acid residue (Thr215) in the NS1 protein (Udom IAV strain) was reported previously to be phosphorylated (35, 36), and it has been proposed that CDKs were the upstream kinases involved in NS1 phosphorylation (35). NS1 is the main viral protein that suppresses the innate anti-influenza immune responses in the host and it interferes with both IFN production and NLR family pyrin domain containing 3 (NLRC3) inflammasome activation (37–39). Furthermore, we recently reported that removing the NS1 C-terminal domain, which hosts Thr215, strongly reduces the capacity of IAV to antagonize antiviral responses in macrophages and in mice (40).

Transcription Regulators, Kinases and Phosphatases, and Transporters Are Differentially Phosphorylated During Influenza Virus Infection—To gain biological insight into the phosphoproteome data we performed in-depth bioinformatics characterization of the data sets. As the first step, the list of the 507 human proteins that were reproducible regulated in their phosphorylation status upon IAV infection was analyzed using the Ingenuity Pathway Analysis (IPA®) (supplemental Table S5). In addition, IPA analysis was done using the 1113 proteins identified as being regulated upon IAV in one of the biological replicates (supplemental Table S5). Classification based on known subcellular location revealed that most of the uniquely phosphorylated proteins had cytoplasmic or nuclear annotation, but several plasma membrane proteins were also identified. A specific annotated molecular type was found for 233 out of the 507 reproducibly identified phosphoproteins and for 502 out of the 1113 phosphoproteins identified in one of the biological experiments (approx. 45%). These phosphoproteins were mainly transcription regulators, kinases and phosphatases, other enzymes and transporters (Fig 1D). The transcription regulators included several distinct protein families such as forkhead box (FOX) proteins, nuclear receptor coactivators, tripartite motif family (TRIM) proteins and members of the mediator complex. This is in line with the previous studies that have linked FOX and TRIM proteins to host responses and control to viral infection (41–44). Our analysis also identified many enzymes involved in nucleic acid metabolism and energy production, such as DEAD box helicases. Kinases identified as uniquely phosphorylated during IAV infection included mitogen-activated protein kinases (MAPKs), as well as several members of CDK, Ca²⁺/calmodulin-dependent kinase (CaMK) and cAMP-dependent protein kinase (PKA) families, which are known to be activated by various viruses (45–49). Transporters are involved in the movement of ions, small molecules, or macromolecules across biological membranes. During IAV infection, there is a dynamic transport of viral components from the plasma membrane to the nucleus and finally back to the plasma membrane (50–52). Our analysis showed changes in the phosphorylation of many transporters, such as solute carrier family members, sorting nexins, syntaxins and nucleoporins. Sorting nexins and syntaxins are involved in endocytosis and vesicle transport, whereas nucleoporins regulate the movement of macromolecules between the cell nucleus and the cytoplasm. Moreover, transmembrane receptors such as adhesion receptors and IFN- and IL-receptors, ubiquitin-specific peptidases and eukaryotic translation initiation factors (EIFs) were uniquely phosphorylated upon infection.

To determine which kinases are activated upon viral infection we used the Kinase Enrichment Analysis (KEA) (25). KEA is a standard tool for linking mammalian proteins with the kinases that phosphorylate them. To get a complete overview of activated kinases we used for this analysis all the uniquely phosphorylated proteins identified in the control and the infected samples (771 and 654 proteins, respectively). We identified ~20% of the phosphoproteins from the two phosphoproteome data sets as known kinase substrates (Table I and supplemental Table S6). In both samples, there was a significant overrepresentation of substrates for glycogen synthase kinase beta (GSK3B), MAPK14 (p38α), cyclin-dependent kinases CDK1 (CDC2), and CDK2. GSK3B has been shown before to be an important player in IAV entry (53). Recently, it was shown that GSK3B activation is needed for the expression of IRF3-regulated IFN-stimulated genes (54). The KEA analysis results included substrates of several members of the MAPK family (p38α, JNK2, ERK1, JNK1, ERK2, MAPKAPK5, MAPK10, MAPK11, and MAPK13) as highly enriched after infection. The MAPKs ERK1/2, p38, and JNK1/2 have been reported to be activated upon IAV infection (55–57). Especially p38 is considered the most important MAPK in the IAV infection-related immune response (58). Our results support these findings, demonstrating that many substrates of MAPKs are differently phosphorylated upon infection.

Other kinases whose known substrates were enriched in the phosphoproteins that were identified from IAV-infected macrophages included Serine/threonine-protein kinase PAK1 and Rho-associated coiled-coil containing protein kinase 2 (ROCK2). These two kinases are involved in the actin cytoskeleton regulation. PAK1, a target of Ras GTPases, is located upstream of interferon signaling molecules IKKε and TBK-1, which in turn phosphorylates IRF3 to induce IFN-β expression. ROCK2 is regulated by RhoA and RhoC, which belong to the Ras superfamily. ROCK2 plays a role in apoptosis, cell survival, and proliferation. In a siRNA screen where specific host membrane-trafficking genes were silenced, ROCK2 was identified as a cofactor in the hepatitis C virus replication (59). In our analysis, Ribosomal protein S6 kinase
<table>
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alpha-5 (RPS6KA5), also known as mitogen- and stress-related kinase 1 (MSK-1), was enriched in the infected sample compared with control. RPS6KA5 has been described to play a role in innate inflammatory response, for instance in responses to human respiratory syncytial virus infection (60). Taken together, our results indicate that kinases with documented roles in cytoskeleton regulation, viral infection and replication, cell survival, and interferon signaling are activated upon IAV infection in primary macrophages.

**Fig. 2.** The phosphorylation profiles change upon influenza A virus infection for host proteins known to be involved in different stages of viral infection. In total 180 proteins of the 1113 uniquely phosphorylated proteins were directly associated with the biological function “viral infection” according to IPA®, and 177 of these had annotated subcellular locations. These 177 proteins are shown and their molecular type is indicated by specific shapes. The purple nodes are proteins, which have unique phosphorylation sites in both uninfected (control) and infected conditions. The red nodes are proteins, which have unique phosphorylation sites only after infection. The green nodes are proteins that have unique phosphorylation sites only in the control sample. Proteins associated with small GTPase signaling (connected with magenta colored edges), replication of influenza A virus (connected with orange colored edges), and endocytosis (connected with blue colored edges) are highlighted with edges connected to the respective biological function. Phosphoproteins with novel phosphorylation sites are highlighted in bold text coding and yellow colored outlining.

Influenza A Virus Infection Changes the Phosphorylation of Many Host Proteins and Signaling Pathways Related to Viral Infection—One of the top biological functions in the pathway analysis was “infectious disease,” and within this category, the most significant annotated function was “Viral infection” (supplemental Table S5). From the 507 reproducibly regulated proteins upon IAV infection 99 were directly linked to viral infection (supplemental Table S5). Of all the 1113 proteins regulated by IAV infection, 180 proteins were directly associated with viral infection (Fig. 2). All the phosphorylated proteins regulated by IAV infection were further analyzed with the InnateDB database (23), which was developed explicitly for systems biology research on the innate immune system (supplemental Table S7). The KEGG pathway-based analysis was very similar to the IPA® results, showing that endocytosis, mTOR signaling, spliceosome, and RNA transport, all of which are important in the IAV life cycle, are enriched in the infected macrophage samples.

Viruses use host signaling pathways for promoting their life cycle, and consistently, our analysis showed here that several host factors which appear to be important for the replication of IAV were uniquely phosphorylated during infection. For example, twelve of the virus infection-associated phosho-
proteins that we identified have previously been linked to endocytosis, an early step in the IAV cellular replication cycle (Fig. 2). These include FCH domain only 2 (FCHO2), which has been described to regulate the early steps of clathrin-mediated endocytosis of IAV, and its silencing inhibited the replication of A/WSN/33 (H1N1) influenza virus (61). Small GTPase signaling was also activated during infection: Cdc42, Rac1 and Rho canonical signaling pathways were top pathways and several proteins associated with IAV infection were linked to small GTPase signaling (Fig. 2 and supplemental Table S5). G-protein-coupled receptors on the surface of phagocytes such as macrophages sense microbial presence and indirectly activate Rac, Rho, and Cdc42. These proteins transduce intracellular signaling leading to the production of reactive oxygen species, which increases the antimicrobial potency of macrophages. Rho signaling pathway has been linked to IAV uptake and internalization, as RhoA is thought to influence the Ca$^{2+}$ response in IAV infected cells (51). Thus, both the IPA® results and the InnateDB database REACTOME pathway results (supplemental Table S7), with “Signaling by Rho GTPases” as the top scored pathway, strongly suggest an important role for Rho family of GTPases in regulating IAV replication and/or antiviral response.

Ubiquitination through the Lys63 residue of ubiquitin is associated with signal transduction, and the cross-talk between phosphorylation and ubiquitination is an emerging topic in the regulation of many signaling pathways. Furthermore, ubiquitination and deubiquitination critically regulate virus infection-induced type I IFN production. Two interesting proteins that were specifically phosphorylated in response to IAV infection were TRIM22 and TRIM25, which are ubiquitin E3 ligases. TRIM22 has been shown to restrict the replication of a number of viruses including encephalomyocarditis, hepatitis B, and human immunodeficiency viruses (62). Interestingly, TRIM22 inhibits IAV infection by targeting the viral NP for degradation (42). TRIM25 has been shown to be essential for RIG-I mediated antiviral activity (63). It interacts with RIG-I and delivers the Lys63-linked ubiquitin moiety to the second CARD domain on the N terminus of RIG-I, resulting in its binding to MAVS and a marked increase in RIG-I downstream signaling and type I interferon induction. Influenza A virus NS1 was shown to specifically counteract this TRIM25-mediated RIG-I CARD ubiquitination (43). In addition to ubiquitin conjugating enzymes, IAV infection induced phosphorylation of several ubiquitin carboxyl terminal hydrolases, which reverse the function of ubiquitin conjugating enzymes by hydrolyzing ubiquitin adducts. Of these, ubiquitin carboxyl terminal hydrolases 15 and 25 are of special interest. These two proteins have been shown to negatively regulate virus-induced type I IFN response (64, 65). Our data shows that carboxyl terminal hydrolases 15 and 25 are phosphorylated on Ser225 and Thr149, respectively, during IAV infection. Taken together, our data show that both ubiquitination and deubiquitination events are regulated by protein phosphorylation during IAV infection of human macrophages.

Mitochondria act as critical platforms for the assembly of signaling complexes that operate in innate immunity (66). It was recently shown that MAVS is phosphorylated in response to stimulation on two different C-terminal serine clusters, including Ser426/Ser430/Ser433 and Ser442/Ser444/Ser445/Ser446 residues (67). Of these, Ser442 phosphorylation was essential for MAVS interaction with IRF3, for IRF3 activation and type I IFN production. Our data shows that MAVS is phosphorylated on serine residues Ser165 and Ser222 in response to IAV infection. The Ser165 amino acid residue is part of a target sequence motif (pS/pTXR/K) of protein kinases PKA and PKC. The Ser222 amino acid residue, on the other hand, is situated in kinase substrate motifs of ERK1, ERK2, CDK5, GSK3, G protein coupled receptor kinase, and casein kinase I. Future studies are needed to elucidate the role of these phosphorylations in MAVS-induced innate immune response.

**Cyclin-dependent Kinases are Activated Upon IAV Infection and Their Specific Inhibitors Rescue Primary Macrophages From Virus-induced Cell Death and Suppress IAV Replication**—Next, the NetworKIN algorithm in KinomeXplorer (24) was used for predicting kinases associated with the unique phosphorylation sites. Of the identified 842 phosphosites unique to infection and 1181 unique to control (Fig 1c), kinases were predicted for 240 and 372 phosphosites, respectively (supplemental Table S8). Approximately 10% of the unique phosphorylation sites were predicted to be phosphorylated by cyclin-dependent kinases (see the separate sheet “CDK_sites” in supplemental Table S8 for details). Proteins which were differently phosphorylated upon IAV infection and identified by KEA or NetworKIN as substrates of CDK1 or CDK2 are shown in Fig 3A. In total 122 phosphoproteins were identified as CDK1 or/and CDK2 substrates; 34 were common for both samples, 55 were unique to the control and 33 unique to the IAV sample. The reproducibly identified phosphoproteins as CDK 1 and/or CDK2 substrates are shown in supplemental Table S9. In particular, several CDK1/2 substrates with unique phosphorylation sites identified after IAV infection were linked to virus host interaction. These included proteins known to be involved in transcriptional regulation, mRNA processing, clathrin-mediated endocytosis, and apoptosis. Of these serine/arginine-rich (SR) protein SRRM2 which is involved in messenger RNA splicing is especially interesting. It has been previously shown that SRRM2 is phosphorylated in HIV-1 infection specific manner. It was shown that HIV-1 modulates host cell alternative splicing machinery through SRRM2 during virus entry in order to facilitate virus replication and release (7). Our findings suggest a similar role for SRRM2 during IAV infection. Utilizing the PhosphoNetworks database (68), we further identified substrates for CDK7 and CDK9 in our phosphoproteome analysis (Fig 3A). All these analyses suggest that several CDKs are activated upon IAV infection.
FIG. 3. Cyclin-dependent kinases are activated upon influenza A virus infection, and their specific inhibitors rescue primary macrophages from virus-induced cell death. A, The uniquely phosphorylated substrate proteins of CDK1, CDK2, CDK7, and CDK9 are shown with their gene names. The substrates for CDK1 and CDK2 are classified according to their biological process (note that one protein can belong to more than one class). Black colored coded proteins have unique phosphorylations in control and IAV samples, red colored coded proteins have unique phosphorylations only after IAV infection, and blue colored proteins have unique phosphorylations only in the untreated, control sample. The bold text coded proteins were identified to have novel phosphosites. B, Mock- or influenza A virus (MOI 3) infected primary macrophages were treated with increasing concentrations of various CDK inhibitors (SNS-032, flavopiridol, dinaciclib, roscovitine, and pablociclib). Cell viability was measured with the CTG assay at 24 h post-infection.
Interestingly, also a previous study had identified that CDK-related signaling events were overrepresented in the virus infection-associated phosphoproteome and that CDKs facilitate lytic gammaherpesvirus replication (8).

In addition to a large number of substrates for different CDKs we identified phosphopeptides from CDK11, CDK12, CDK13 in the phosphoproteomics data, suggesting that these could play a role in the IAV life cycle, in particular in primary macrophages, which do not undergo cell division. A proviral function has been attributed to CDK13 in lung epithelial cells (69), and a previously published siRNA screen also concluded that CDK13 is vital for IAV replication (70).

Cyclin-dependent kinases regulate several cellular processes, including the cell cycle, transcription, RNA processing, and cell survival. To study the role of CDKs in IAV-induced cell death and virus replication we used small molecule (molecular mass < 900 Da) kinase inhibitors of CDKs (CDKIs). Pharmacological CDKIs have been originally developed to inhibit CDK signaling in Alzheimer’s and Parkinson’s disease, ischemia, and cancer. More recently, it was shown that CDKIs have antiviral activity against clinically important viruses including hepatitis C virus (71), human cytomegalovirus (72–74), human immunodeficiency virus type-1 (75–77), Epstein-Barr virus (78), and herpes simplex virus type 1 and 2 (79). These small molecule inhibitors have high specificity, good pharmacological properties and little side-effects (80, 81). Furthermore, safety data from numerous clinical trials are available for these drugs. First, we tested the effects of SNS-032, flavopiridol, dinaciclib, roscovitine, and pablociclib on the viability of noninfected (mock) and IAV-infected macrophages (MOI 3, Fig 3B). SNS-032, flavopiridol, and dinaciclib rescued macrophages from virus-mediated death at noncytotoxic concentrations at 24 h post-infection, whereas roscovitine and pablociclib did not have similar potency at the selected range of concentrations. Importantly, the protective effect of SNS-032 was lost when the drug was added to the IAV-infected cells later than 3 h post-infection (supplemental Fig. S3A). We propagated the viruses at MOI 0.01 in macrophages pre-treated at different compound concentrations and calculated the half maximum effective concentrations (EC$_{50}$), half maximum cytotoxic concentrations (CC$_{50}$), and selectivity indexes (SI) for both SNS-032 and flavopiridol (supplemental Fig. S3B). With an EC$_{50}$ of 40 nm and a SI > 500, the SNS-032 proved to be a potent small molecule inhibitor of IAV-associated cellular cytotoxicity in primary human macrophages.

Based on these experiments, we decided to characterize the effects of SNS-032 and flavopiridol (EC$_{50}$ = 200 nm and SI > 100) on IAV-infected human macrophages in more detail. RT-qPCR analysis showed that both SNS-032 (0.3 µM) and flavopiridol (0.3 µM) inhibit expression of viral M2 and NP mRNAs in macrophages that were infected with IAV for 6 h (Fig 4A). In line with this result, these CDKIs also blocked the expression of viral NS1 and NP proteins in macrophages that had been infected with IAV for 18 h (Fig 4B). Thus, it is highly likely that these CDKIs inhibit IAV replication.

Selected Cyclin-dependent Kinase Inhibitors Suppress Antiviral and Chemokine Response in IAV-infected Human Macrophages—Human macrophages respond to IAV infection by producing antiviral cytokines, including type I and III IFNs and chemokines that can attract other immune cells to the site of infection. To study the effect of CDKIs on IAV-induced antiviral cytokine and chemokine response, macrophages were infected with IAV for 6 h in the presence and absence of CDKIs. After this, total cellular RNA was isolated, cDNA was prepared and the expression of IFNs and chemokines was analyzed by RT-PCR. The inhibitors had modest, but significant, effect on IAV infection-induced IFN-β mRNA levels (Fig 4C). In contrast, both flavopiridol and SNS-032 treatment completely abolished IAV infection-induced expression of IL-29 (IFN-λ1), CXCL10, and CXCL11 genes (Fig 4C). Flavopiridol or SNS-032 administration did not affect GAPDH or 18S ribosomal RNA expression, indicating that these inhibitors do not interfere with the general cellular transcription (supplemental Fig. S4).

Next, we analyzed with the cytokine array proteome profiler the production of pro-inflammatory cytokines and chemokines from cell culture supernatants of IAV-infected human macrophages in the presence or absence of CDKIs. The secretion of chemokines CCL2, CCL3, CCL4, RANTES (CCL5), and CXCL10 was completely abolished by flavopiridol or SNS-032 in IAV-infected human macrophages (SSA Fig). Similarly, the secretion of cytokines IL-6 and IL-16 as well as complement component C5/C5a was totally inhibited by flavopiridol or SNS-032 in IAV-infected human macrophages (Fig S4B, S4C). Furthermore, IAV-infected macrophages—

IRF3 and NF-κB signaling pathways are involved in the activation of antiviral and pro-inflammatory response during viral infections, and we studied the effect of SNS-032 on these signaling pathways during IAV infection. Human macrophages were infected with IAV for 6 h in the absence and presence of SNS-032 after which cell lysates were prepared and Western blot analysis was performed with anti-phospho IRF3 and anti-IκBα antibodies (Fig 4D). SNS-032-treatment clearly but not completely inhibited the IAV-induced phosphorylation and activation of IRF3. The IFN-β promoter has been shown to have a very high affinity for activated IRF3 compared with other antiviral genes (82). Therefore, the expression of IFN-β can be induced by IRF3 that is only modestly phosphorylated which was seen in SNS-032 treated and IAV-infected macrophages (Figs. 4C and 4D). Interestingly, SNS-032 alone induced degradation of IκBα demonstrating activation of NF-κB. Furthermore, IκBα was almost completely lost in SNS-032-treated and IAV-infected human.
macrophages indicating robust activation of NF-κB under these conditions. In conclusion, these results show that SNS-032 differentially regulates IRF3 and NF-κB signaling pathways.

CDKIs Abrogate Apoptosis and Inflammasome Activation in IAV-infected macrophages—IAV infection of human macrophages is associated with apoptosis of the infected cells (83). We therefore decided to study the effect of SNS-032 and flavopiridol on IAV-induced apoptosis. There are two main apoptotic pathways, the extrinsic pathway, regulated by activation of extracellular death receptors that functions through caspase 8, and the intrinsic pathway that functions through caspase 9 and is initiated by the disruption of the mitochondrial membrane potential. Both caspase 8 and 9 activate caspase 3 by proteolytic cleavage. Caspase 7 is also a downstream effector protease of the apoptotic pathway. The activity of caspases 3/7, 8, and 9 was measured with the Caspase-Glo assay in human macrophages that were infected with IAV in the presence and absence of SNS-032. IAV infection of human macrophages clearly enhanced caspase 3/7 activity at 18 h post-infection and this activation was completely inhibited by SNS-032 (Fig 5A).

In accordance with this result, both CDKIs abrogated the up-regulation of...
Bcl-xL. Bcl-xL is an anti-apoptotic protein whose expression can be induced in response to IAV infection (84). Bcl-xL is a component of a cell signaling network that governs cell survival and cell death by interacting with pro-apoptotic proteins such as Bad and Bax (85). The expression of Bcl-xL is regulated by NF-κB and other transcription factors, e.g., CREB and STATs. JAKs and Src kinases phosphorylate and activate STAT3, which is known to induce the expression of Bcl-xL (86). It is possible that in the absence of IFNs and other cytokines in infected but SNS-032 treated cells, the JAK and Src signaling pathways are not activated and the expression of Bcl-xL is not induced.

IAV infection of human macrophages is associated with caspase 1 activation and the subsequent secretion of pro-inflammatory cytokines IL-18 and IL-1β (16). Caspase 1 is activated in a molecular platform called NLRP3 inflammasome, comprising the protein NLRP3, the adaptor apoptosis-associated speck-like protein (ASC) and pro-caspase 1. The NLRP3 inflammasome has been shown to be essential for the activation of host response during IAV infection (87, 88). To study the effect of CDKIs on NLRP3 inflammasome activation we infected human macrophages in the presence and absence of SNS-032 and flavopiridol and analyzed IL-18 secretion and caspase 1 activation. Our results show that these CDKIs completely block IAV-induced secretion of IL-18 (Fig 5C). The activity of caspase 1 was increased in human primary macrophages after 18 h of IAV infection and this activity was significantly decreased when cells were treated with SNS-032 before infection (Fig 5D). These data demonstrate that the CDKIs prevent NLRP3 inflammasome activation. It is likely that CDKIs, and especially SNS-032, block IAV life cycle before the NLRP3 inflammasome can be assembled and activated.

CDKIs Protect Against IAV Challenge in Mice—Finally, we decided to validate the protective potential of the observed antiviral effects of SNS-032 in vivo in a standard mouse model for influenza infection. As a challenge virus, we used A/Puerto Rico/8/34 (PR8) virus because this virus is very well adapted to mice meaning that it causes disease in e.g. BALB/c mice after inoculation with as little as 10 plaque forming units.

Fig. 5. Cyclin-dependent kinase inhibitors block influenza A virus-induced apoptosis and inflammasome activation. A, Macrophages were pretreated with SNS-032 (0.3 μM) or flavopiridol (0.3 μM) for 1 h after which they were left uninfected or infected with IAV for 18 h. Caspase 3/7, caspase 8, and caspase 9 activity was measured as described under Experimental Procedures. The changes in activity were calculated and compared with the mock/untreated sample (= 1). Data is presented as mean and S.D. of n = 3 biological replicates. The results were compared with the infected and untreated sample. B, Macrophages were pretreated with SNS-032 (0.3 μM) or flavopiridol (0.3 μM) for 1 h after which they were left uninfected or infected with IAV for 18 h. After this, total cell lysates were prepared and expression of cleaved caspase 3 (Asp175) and Bcl-xL expression was analyzed with Western blotting. C, Macrophages from three different donors were pretreated with SNS-032 (0.3 μM) or flavopiridol (0.3 μM) after which they were infected with IAV for 18 h. After this, the cell culture supernatants were collected and IL-18 secretion was analyzed with a Luminex assay. Data is presented as mean and S.D. of n = 3 independent measurements. The results were compared with the infected and untreated sample. D, The activity of caspase 1 was measured in 18 h infected or noninfected macrophages, nontreated or treated with SNS-032 at 0.3 μM at 18 h post-infected. Data is presented as mean and S.D. of n = 3 biological replicates. The results were compared with the infected and untreated sample.
addition, this virus productively infects murine macrophages (89). We focused on SNS-032 because it showed better antiviral activity in vitro compared with other CDKIs. In a first experiment female BALB/c mice (6 mice/group) were treated with SNS-032 (15 mg/kg in 5% DMSO) or vehicle (5% DMSO) administered intraperitoneally every other day, starting on day 1 before and until day 9 after infection. For infection we used 0.5 LD50 (corresponding to 20 plaque forming units per mouse) of mouse adapted PR8 strain. Kaplan-Meier survival curves of the infected mice show perfect protection gained by SNS-032 treatment (log-rank test, p < 0.01). B. Relative body weight curves for the two different groups of infected mice. SNS-032 treated mice lost significantly less body weight than the DMSO treated mice at days 4–7 (two-way ANOVA, Bonferroni’s multiple comparison adjustment, p < 0.0001). At 9 days post infection more than half of the DMSO treated mice had died and hence from day 9 on, the body weight of the surviving DMSO treated mice was not included in the figure. C. Cytokine levels in mouse lung homogenates at day 5 were determined using mouse cytokine array panel A kit. Lung homogenates from four mice per group were combined before measuring the relative intensity of the luminescence for the different cytokines. Representative results from two experiments are shown.

Fig. 6. SNS-032 treatment rescues influenza A virus-infected mice. Mortality and morbidity of SNS-032 treated and DMSO treated mice after lethal influenza infection. At day 0 the mice were infected with 0.5 LD50 of mouse-adapted PR8 or mock infected. Six mice per group were repeatedly treated intraperitoneally with SNS-032 at 1 day prior and 1, 3, 5, 7, and 9 days post infection. A. Kaplan-Meier survival curves of the infected mice show perfect protection gained by SNS-032 treatment (log-rank test, p < 0.01). B. Relative body weight curves for the two different groups of infected mice. SNS-032 treated mice lost significantly less body weight than the DMSO treated mice at days 4–7 (two-way ANOVA, Bonferroni’s multiple comparison adjustment, p < 0.0001). At 9 days post infection more than half of the DMSO treated mice had died and hence from day 9 on, the body weight of the surviving DMSO treated mice was not included in the figure. C. Cytokine levels in mouse lung homogenates at day 5 were determined using mouse cytokine array panel A kit. Lung homogenates from four mice per group were combined before measuring the relative intensity of the luminescence for the different cytokines. Representative results from two experiments are shown.

Concluding Remarks—Like all other viruses, influenza A viruses rely on host factors for their life cycle; they manipulate and hijack cellular proteins for promoting their uptake, replication, and egress. Knowing the molecular mechanisms behind these events, for instance, how the viruses impact host cellular processes through protein phosphorylation, is important for understanding how viral diseases progress and for developing new antiviral treatments. Here, we took a mass spectrometry-based phosphoproteomics approach com-
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bined with bioinformatics and functional studies to elucidate the host-response to influenza A virus infection in primary human macrophages. The focus was on the early phase of infection, corresponding to the time frame in which innate immune responses are typically initiated. Our data shows that there are phosphorylation changes in the host proteins involved in all critical stages of viral replication including entry, gene regulation and egress. Our results also highlight the importance of using primary cells in proteome-level studies to obtain novel and biologically most meaningful data for further functional studies. An important finding of our work is the remarkably altered phosphorylation status of several cyclin-dependent kinases and their substrates following IAV infection. Functional studies based on well-characterized small molecule CDK inhibitors showed that the CDK activity is required for efficient viral replication and for the activation of host response in vitro. Further, in vivo experiments showed that SNS-032, one of the selected CDKIs, could rescue IAV-infected mice.

In conclusion, we provide the first comprehensive phosphoproteome characterization of IAV infection in primary human macrophages, and provide extensive evidence that specific CDK inhibitors could be excellent candidates to treat severe influenza virus infections.

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