Lipid biophysics and/or soft matter-inspired approach for controlling enveloped virus infectivity

Lamyaa Al-dalawi, Stephen P. Dunham and Cyril Rauch

School of Veterinary Medicine and Science, University of Nottingham, Sutton Bonington Campus, College Road, Sutton Bonington LE12 5RD, UK

Proven as a natural barrier against viral infection, pulmonary surfactant phospholipids have a biophysical and immunological role within the respiratory system, acting against microorganisms including viruses. Enveloped viruses have, in common, an outer bilayer membrane that forms the underlying structure for viral membrane proteins to function in an optimal way to ensure infectivity. Perturbing the membrane of viruses using exogenous lipids can be envisioned as a generic way to reduce their infectivity. In this context, the potential of exogenous lipids to be used against enveloped virus infectivity would be indicated by the resulting physical stress imposed to the viral membrane, and conical lipids, i.e. lyso-lipids, would be expected to generate stronger biophysical disturbances. We confirm that when treated with lyso-lipids the infectivity three strains of influenza virus (avian H2N3, equine H3N8 or pandemic human influenza H1N1) is reduced by up to 99% in a cell-based model. By contrast, lipids with a similar head group but two aliphatic chains were less effective (reducing infection by only 40–50%). This work opens a new path to merge concepts from different research fields, i.e. ‘soft matter physics’ and virology.

1. Background

Respiratory virus infections pose a significant threat to human health. Influenza viruses continue to cause recurrent epidemic and pandemic infections and the more recently emerged SARS-CoV-2 has threatened to overwhelm healthcare systems globally causing widespread mortality and disease. The human body has many natural or innate defences against viruses including pulmonary surfactant. Surfactant is composed of lipids (approx. 85% of phospholipids, approx. 10% neutral lipids) and proteins (approx. 5%) [1]. The initial physiological role and clinical importance discovered for pulmonary surfactant was its involvement in avoiding the collapse of pulmonary alveolus during end-exhalation [2]. More recently, pulmonary surfactant has been shown to participate in the innate host defence against respiratory pathogens [3–5]. Within the last decade, single lung surfactant phospholipids have been shown to protect, to some extent, cells from virus infectivity, including respiratory syncytial virus and influenza A [6–9]. Specific lipids found in the lung surfactant such as DPPG (di-palmitoyl-phosphatidylglycerol), POPG (palmitoyl-oleoyl-phosphatidylglycerol) or DPPC (di-palmitoyl-phosphatidylcholine) have been shown to antagonize virus infectivity [5,7]. However, a full understanding of their mechanism of action remains elusive.

The initial stage of replication requires the influenza virus to enter cells based on the interaction between viral surface hemagglutinin protein (HA) and cellular sialic acids. The stereochemistry and collective structure of viral HA, neuraminidase (NA) and matrix protein 2 (M2) are central for successful infectivity [10,11].
These proteins are embedded within a bilayer lipid envelope derived from cell membrane on budding.

Although it is often routine to think pharmacologically in terms of drug–target or surfactant component–virus chemical interaction, in the context of lipid surfactant the amphipathic nature of lipids, namely their property of being structurally composed of hydrophobic and hydrophilic parts, needs to be considered carefully.

Indeed, lipids have the unique property to self-assemble physically and it is perhaps not surprising then that in the presence of surfactant lipids, membrane blebs appear on the viral envelope [5]. Following Laplace’s Law, without any variation in the osmotic pressure difference between the inside and outside of the virus, the formation of nanometric membrane blebs suggests a direct change in the surface tension of the viral envelope linked to the inclusion of surfactant lipids. As a result, it is possible that lung surfactant lipids interact directly with the bilayer membrane of the viral envelope in turn destabilizing the collective structure of key viral envelope lipoproteins, i.e. HA, NA and M2, with a resulting impact on virus infectivity. Such an effect would not be unexpected, as in mammalian cells the mechanical perturbation of the bilayer membrane linked to the exogenous inclusion of amphipatic materials, e.g. phospholipids, has been shown to affect the function of ion channels [12], drug transporters [13], endocytosis [14] and exocytosis [15].

Therefore, surfactant lipids may provide a biophysics-based generic immunity against a plethora of inhaled pathogens provided they have an outer bilayer membrane. If the physical properties of surfactant lipids are essential to set up this generic immunity, the next question to ask is: can we optimize them, i.e. what lipid surfactant would be the most efficient?

The basic geometry or topology of lipids is essential. Lyso-lipids, i.e. single aliphatic chain lipids, have a tendency to form small unilayer structures due to their conical shape whereas lipids with two aliphatic chains form larger bilayer liposomes due to their cylindrical shape [16]. The changes in the surface pressure generated upon inclusion into a bilayer membrane are expected to be different as lyso-lipids tend to promote sharper curvature.

Given the set of observations [5,17,18], the aim of this work was therefore to connect a physical intuition regarding the physical properties of lipids to virology. The current study is focused on biology/virology rather physics/modelling as its central purpose was to demonstrate, as a proof of concept, whether specific exogenous lipids reduce the infectivity of the influenza A viruses including H2N3, H1N1 and H3N8. Herein is shown that specific lipids are altering the virus morphology and inhibiting the binding of the virus to the host cell membrane with an impact on the expression of cellular cytokines (IL-8, TNF-α) and the expression of viral M-gene.

2. Results

2.1. Impact of exogenous lipids on H2N3 topology and structure

Hypothesizing that lipids have a physical impact on viruses, this should be visible using transmission electron microscopy (EM). Figure 1a shows a typical H2N3 virus where the surface glycoproteins and spherical shape of the virus are clearly visible. However, upon incubation with different classes of lipids including LPG, DPPG, LPC and DPPC for 30 min at 37°C, the virus morphology changed drastically as a function of concentrations used. EM pictures show that virions are surrounded by lipid structures visible as white foci next to the virus. Incubation of H2N3 with DPPC showed the least change in virion appearance, but at the other extreme, LPG treatment demonstrated rupture of the viral membrane suggesting an irreversible process (figure 1b). In order to quantify the impact of phospholipids on H2N3, the virus aspect ratio and surface area were measured (figure 1c,d, respectively) to document the observed change from a spherical shape toward a more elongated morphology. Uninfected H3N3 viruses showed a value for the aspect ratio of approximately 1 confirming that H2N3 are natively round (spherical). However, following treatment with amphipatic compounds, the mean aspect ratio increased demonstrating that viruses became ellipsoidal in shape. The change in aspect ratio was more pronounced following treatment with lysosilipids (LPG or LPC). Regarding H2N3 surface area, results demonstrated a larger spread of numerical values for all lipids used (figure 1d). A larger surface area would be expected by the incorporation of exogenous lipids into the virus membrane. However, virions with a reduced surface area, compared to untreated virus, were also observed, suggesting that upon incorporation of lipids, a virus may become unstable, undergoing fission to form smaller ‘daughter’ virions.

2.2. Minimal impact of exogenous lipids on mammalian cells

Given the observation that viruses pre-treated with lipids demonstrated an irreversible impact on their morphologies, the direct impact of lipids on cells needed to be assessed as free lipids could also interact directly with cells used in virus infection experiments. Lipid solutions were prepared and incubated directly with Madin Darby canine kidney (MDCK) cells. Direct observation by light microscopy identified minimal differences in cellular morphology (data not shown). Flow cytometry was, therefore, used to assess the impact on cell morphology. The addition of 500 µM LPG to cells for 10 min resulted in a small proportion of cells (approx. 1.4%) showing an increase in side-scatter, suggesting increased cell granularity suggesting a change in the membrane intracellular architecture (figure 2a).

As a result, a method was devised to remove free lipids following the treatment of virus to minimize any direct impact on cells. After several optimizations using flow cytometry and measuring cell viability using an MTT essay, the best method found was to centrifuge the solution containing lipids using ultrafiltration. Indeed, the flow cytometry data demonstrated that once the retentate was resuspended with the infection medium only a small fraction (less than 0.1%) of MDCK cells were affected (figure 2b) and that their viability remained unchanged for a period of up to 6 h (figure 2c).

2.3. Impact of lipid pre-treatment on the attachment, internalization and infectivity of H2N3 in MDCK cells

As the lipids used had a strong impact on virus morphology it was hypothesized that such an impact could impair their
ability to infect cells. To determine whether H2N3 virus attachment to cells was affected, virus (at a concentration to allow infection at a multiplicity of infection (MOI) of 5) was treated with lipids prior to ultrafiltration and the retentate was resuspended in cold infection medium (4°C). The cold infection solution was then incubated with cells for 10 min on ice. Cells were then washed with cold phosphate-buffered saline (PBS) and were either fixed with ice cold 1% PFA to determine the amount of viruses initially bound to cells, or reincubated with the infection medium at 37°C for different time points prior to fixation with PFA. HA was labelled using immunofluorescence without additional permeabilization of MDCK cells to allow determination of the amount of viruses remaining on the MDCK surface over time. Figure 3 illustrates that the initial attachment of viruses to cells is impaired upon virus pre-treatment with lipids in comparison with untreated virus. After 30 min, only MDCK cells infected with untreated virus showed a noticeable diffuse fluorescent signal for HA suggesting that virus had entered cells and was detected by immunohistochemistry despite no additional permeabilization having been performed (figure 3b). However when the viruses were pre-treated with lipids a reduction in the fluorescent intensity within cells was clearly visible. In order to assess quantitatively the amount of viruses bound to the cell membrane over time flow cytometry was used. Data in figure 3c derived from flow cytometry analysis of MDCK infected cells demonstrate that the initial attachment of viruses to the MDCK cell membrane is impaired and is a function of the concentration of lipids used to pre-treat the virus.

The impact of lipid treatment on virus infection of MDCK cells was also assessed by immunofluorescent staining of cells for viral nucleoprotein. MDCK cells were infected with H2N3 virus at an MOI of 1.0 for 2 h, with or without lipid pre-treatment of virus inoculum, prior to washing cells in PBS and incubation of cells for a further 4 h. Cells were then fixed and the viral nucleoprotein (NP) was labelled using immunohistochemistry to determine the proportion of infected cells (measured as focus forming units (ffu) per ml). Data presented in figure 4a–c demonstrate that H2N3 infectivity was significantly reduced when DPPG, LPG or LPC were used for the virus pre-treatment as opposed to DPPC which showed little impact. LPG used at 500 µM showed the greatest reduction in virus infectivity (to 3.5% compared with untreated virus; table 1).

However, with these data it was not possible to differentiate whether the drop in infectivity was simply linked to a reduction in the number of virions attached to cells initially or resulted from the impairment of the virus to multiply inside cells. To resolve this point, we studied...
the two lipids having the most significant impact on attachment and infectivity, i.e. LPG and DPPG. The infectivity experiment was then repeated with using virus at an MOI of 0.1 allowing infection to proceed to 48 h. Figure 4d demonstrates that when the virus was pre-treated with either LPG or DPPG the logarithmic increases of

![Figure 2](https://royalsocietypublishing.org/doi/10.1098/rsif.2021.0943)
infectivity over time were similar to the control, in turn suggesting that the ability of H2N3 to multiply inside cells was not affected.

Taken together, the results suggest that a small fraction of pre-treated viruses are still functional and that pre-treating H2N3 with lipids only reduces the initial level of virus infection.

**Figure 3.** Influence of phospholipids on H2N3 (MOI 5) virus binding to MDCK cells. (a) Detection of H2N3 via FITC-labelled anti-HA antibody after binding at 4°C for 10 min and (b) after a further 30 min incubation at 37°C. (c) Flow cytometry analysis showing the proportion of cells with H2N3 bound to the surface over time. Means and standard errors of at least three experiments are represented. These results demonstrate that pre-treating H2N3 with phospholipids impacts on its ability to bind to cells. (*p < 0.05; **p < 0.01; ***p < 0.001; one-way ANOVA.)
2.4. Transcription of IL-8, TNF-α and M-gene viral protein upon infection of MDCK cells with H2N3 pre-treated with lipids

In order to determine whether the cell responses were influenced by the reduced level of virus infection following lipid treatment, H2N3 was pre-treated with either LPG or DPPG and incubated with cells for 2 h prior to resuspending cells in a new infection medium (using untreated virus at an MOI of 5 as performed in figure 3a). After 22 h, cellular RNA was extracted and qRT-PCR used to determine the expression level of key host proinflammatory cytokines (IL-8 and TNF-α). Figure 5a,b demonstrates that the pre-treatment of H2N3 with either LPG or DPPG significantly reduced the expression of these cytokine mRNAs in MDCK cells. Cells treated with 500 µM of DPPG or LPG only, i.e. without viruses, showed no significant change in cytokine expression compared with untreated cells, suggesting that lipid treatment per se is not proinflammatory. Finally, the ability of H2N3 to multiply inside cells was also assessed by measuring the amount of mRNA for the viral M-gene by qRT-PCR. As expected the amount of M-gene mRNA expression was also downregulated when H2N3 viruses were pre-treated with lipids (figure 5c).

Altogether, these results confirm that pre-treating H2N3 with lipid reduces the level of virus infection, delays virus replication kinetics and consequently reduces the proinflammatory cell response.

2.5. Impact of lipid pre-treatment of H2N3 using A549 cells: attachment, infectivity and cell response

Our previous results demonstrated that H2N3 pre-treated with lipids impacts its ability to bind to and infect MDCK cells. However, such results could be specific to the cell line used. In order to assess this further, experiments were repeated using human adenocarcinomic alveolar basal epithelial (A549) lung cells. As previously determined with MDCK cells, the ultrafiltration method efficiently removed LPG and DPPG, with no impact on A549 cell viability noted using the MTT assay over a period of 6 h (figure 6a). Using the same method as for MDCK cells, H2N3 attachment to A549 cells was visibly impaired when H2N3 (sufficient to give an MOI of 5) was pre-treated with lipids (figure 6b,c). As expected, this resulted in a drop in virus infectivity, assessed as before, using immunohistochemistry for viral NP (at MOI of 1; figure 6d,e) that was more pronounced when LPG was used (infectivity reduced to 0.4% with 500 µM LPG or 5.5% with 50 µM LPG; table 1). IL-8 and TNF-α expression was similarly impacted by pre-treating H2N3 with either DPPG or LPG (figure 6f). Finally, it was found that DPPG and LPG significantly reduced H2N3 M-gene copy number.
reflecting a reduction in cell infection (figure 6f). Those results confirmed that the virus ability to infect cells is altered upon treatment with lipids, irrespective of the cell line used.

2.6. Impact of lipid treatment on filamentous H3N8 virus

Although H2N3 is natively spherical in shape, viruses can differ in their morphology and this is particularly the case for influenza viruses where filamentous viruses appear to be more frequently associated with clinical infections [19,20]. In order to assess the effect of lipid treatment on the infectivity of filamentous virus we used equine H3N8 which exhibits a largely filamentous morphology [21] to perform similar studies as for H2N3, using LPG or DPPG. Using EM, results demonstrated that treatment of H3N8 virus with either LPG or DPPG impacts its morphology (figure 7a,b), affecting the distribution of envelope glycoprotein. However, the reduction in infection elicited upon DPPG incubation was moderate when compared to greater reductions when H2N3 was used. Only when LPG was used were multilayered viral structures observed suggesting rupture of the viral membrane (compare figure 7a,b for LPG at 500 µM). Given the nature and three-dimensional shape of the filamentous virus it was not possible to quantify those changes through the measurements of H3N8 aspect ratio or its surface area using EM.

Regarding H3N8 attachment to cells, it was found that the interaction of the filamentous virus with MDCK cells was lower for untreated virus when compared to H2N3 in similar conditions (compare figure 7c with figure 3a), despite using virus at a similar MOI of 5. This may reflect a greater proportion of defective interfering particles in H2N3 virus preparation which are non-infectious but bind to MDCK cells. However, treatment with either LPG or DPPG still had a clear effect as the fluorescence intensity upon labelling H3N8 HA was much lower than the control (figure 7d). After a further 30 min incubation, infected cells were clearly visible for the control through the presence of clusters of fluorescence (figure 7d). Those clusters were not visible when H3N8 was pre-treated with either LPG or DPPG.

Measurement of infectious virus, by immunostaining for virus nucleoprotein, following incubation of virus for 30 min with LPG or DPPG (5, 50 or 500 µM) showed that virus pre-treatment with LPG had a more pronounced impact on H3N8 infectivity (MOI 1) when compared with DPPG (figure 7c,f). Treatment with 500 µM or 50 µM lipid reduced H3N8 infectivity to 7.4% or 14.7% for LPG and 52.0% or 54.4% for DPPG, respectively (table 2).

Interestingly, it was found that only IL-8 transcription was significantly affected by pre-treatment of H3N8 (MOI 5) with either LPG or DPPG after 24 h infection, producing a modest reduction in the increased IL-8 expression observed following infection with untreated virus (figure 7g), whereas no significant changes were noted for TNF-α.

Those results suggest that the initial morphology of the virus, i.e. spherical or filamentous, has a significant impact on virus sensitivity to lipids. The impact of DPPG on virus infection was less marked when H3N8 was used when compared with H2N3. However, LPG had a greater impact on virus infection than DPPG for both viruses. Those results suggest that LPG is highly efficient whatever the initial morphology of the virus, but that the impact of DPPG is probably a function of the excess of viral membrane surrounding the virus, i.e. that the effect of such lipid is ‘diluted’ by the amount of lipid membrane forming the envelope of the filamentous virion. To confirm this, another spherical virus, H1N1, was used.

2.7. Impact of lipid pre-treatment on H1N1 infectivity and the transcription of IL-8 and TNF-α using MDCK cells

H1N1 (MOI 1) was pre-treated with DPPG and LPG using the same conditions as for H2N3 to measure its infectivity. Both LPG and DPPG altered H1N1 infectivity (figure 8a,b). The drop in infectivity for H1N1 treated with 500 µM DPPG to 4.6% was more pronounced than for H2N3 (32.1%; table 2). In line with the previous observations, LPG produced greater inhibition of virus infection compared with DPPG, resulting in a decrease to 1.3% or 5.6% infectivity when treated with 500 µM or 50 µM, respectively (table 2). Finally, the transcriptions of IL-8 and TNF-α were significantly and similarly reduced upon lipid pre-treatments of H1N1 (MOI 5) (figure 8c). This is consistent with the changes in IL-8 and TNF-α expression measured for H2N3 but differs from the results obtained using H3N8.
occasions, the viral envelope is ruptured. A less quantifiable
transformed into a giant/swollen structure and on some
outer surface of the virus. In the presence of lipids, H2N3 is
inserted into bilayer membrane, lipid surfactant analogues
have been successfully applied to a wide range of complex
systems, very little has been done in the field of virology.
More precisely, based on the physical principle that single
barrier against host pathogen infection, focusing on IAV.
In normal conditions H2N3 has a spherical shape show-
ing outer glycoprotein spikes evenly distributed around the
outer surface of the virus. In the presence of lipids, H2N3 is
transformed into a giant/swollen structure and on some
occasions, the viral envelope is ruptured. A less quantifiable
but similar impact was observed on H3N8, a filamentous
virus, suggesting that lipid interaction with the viral mem-
brane is independent of the initial virus morphology.

Influenza virus has a serious, worldwide, effect on human
and animal health with a yearly seasonal death toll of
many thousands, which increases during deadly pandemic
episodes. Inactivated and attenuated vaccines against influ-
enza A and B viruses (IAV and IBV) have been used to
control influenza virus in addition to prophylactic and treat-
ment attempts by using antiviral drugs but, still, significant
research efforts are needed to control influenza viral infec-
tions. More recently the emergence of SARS-CoV 2 has
resulted in an unprecedented virus pandemic that has had
a devastating effect worldwide; mortality rates remain high
and new treatments are urgently required.

The advances made in the field of soft matter physics
encompassing lipid membranes and polymers have contribu-
ted to a better understanding of complex molecular structures
and cell biology as a whole. The aim of this work was, there-
fore, to provide an interdisciplinary understanding of how lung
surfactant phospholipids may function as an innate barrier against host pathogen infection, focusing on IAV. More precisely, based on the physical principle that single chain lipids are more prone to generate curvature upon their
insertion into bilayer membrane, lipid surfactant analogues
were tested against specific influenza viruses, i.e. H2N3, H3N8 and H1N1. The overarching conclusion is that lyso-
lipids, in our case LPG and LPC, have indeed a more pro-
nounced impact on virus infectivity independently of the
cell type used (summarized in tables 1 and 2). It is remarkable
that while concepts from soft-matter physics or biophysics
have been successfully applied to a wide range of complex
living systems allowing a better understanding of those sys-
tems, very little has been done in the field of virology.

In normal conditions H2N3 has a spherical shape show-
ing outer glycoprotein spikes evenly distributed around the
outer surface of the virus. In the presence of lipids, H2N3 is
transformed into a giant/swollen structure and on some
occasions, the viral envelope is ruptured. A less quantifiable
but similar impact was observed on H3N8, a filamentous
virus, suggesting that lipid interaction with the viral mem-
brane is independent of the initial virus morphology.
Previous studies have demonstrated that influenza virus
binds efficiently with mixed liposomes composed of phos-
pholipids within minutes [22] and that depleting the
cholesterol from the viral envelope affects virus morphology,
with virions showing nicks or holes in parts of the viral enve-
lope [23]. Our study is therefore aligned with the latter
conclusions that lipids interact with viruses and the envelope
morphology is impacted upon changing its composition.

Those morphological alterations were associated with a
decrease in the ability of viruses to interact with the cell mem-
brane independently of the cell type. However, the ability of
the virus to replicate inside cells was not affected, as the kin-
etics of replication were the same between the control and
treated viruses. The influence of lipids on virus infectivity
in the current study is in agreement with previous
observations showing that the pulmonary surfactant phos-
phatidyglycerol (PG), which functions as the main innate
immunity regulator in the lungs, blocks pulmonary viral
infections [24] such as respiratory syncytial virus (RSV).
This could provide an explanation for the higher concen-
tration of phosphatidylglycerol in the lung compared to
other body organs [7]. The airway epithelium is a primary
barrier between the environment and the host, enhancing
inflammatory and chemokine production targeting influenza
viral infections and improving innate antiviral immunity [25].
Accordingly, and in agreement with a drop in infectivity,
alterations in IL-8 and TNF-α cytokine transcription levels
were measured together with a change in the number of
mRNA copies for the viral M-gene 24 h post infection with
influenza viruses.

Within the scope of this study our results demonstrate
that pulmonary surfactant lyso-lipids can be used against
H2N3, H3N8 and H1N1. The physical impact of lyso-lipids
on viruses is expected to be the same across all enveloped
viruses provided the presence of a bilayer membrane.
While the use of lipids to treat acute respiratory distress has
met with mixed results [26], the results presented here
suggest that such treatments need to be rethought using
optimized lyso-lipids with soft-matter/biophysical ‘function-

3. Discussion

Tables 1 and 2 summarize the reduction in virus infectiv-
ity that was measured for each combination of lipid, virus
and cell line.

![Figure 5. Cytokine transcription levels and M-gene copy number 24 h post infection of MDCK cells with H2N3 (MOI 5) pre-treated with 500 µM LPG or DPPG. (a) IL-8, (b) TNF-α, (c) M-gene copy number. Means and standard errors of at least three experiments are represented. Altogether those results confirm that the cells’ response to infection is diminished when viruses are pre-treated with phospholipids. (*p < 0.05; **p < 0.01; ***p < 0.001; †p < 0.0001; one-way ANOVA.)](https://royalsocietypublishing.org/journal/rsif)
virus infections. Future treatments with lyso-lipids could therefore be used to control the 'peak' of infectivity for enveloped virus infections of the respiratory tract, including SARS-CoV 2. This could delay disease progression, and, in combination with other therapeutics, reduce disease morbidity and mortality.
Figure 7. MDCK cells infected with equine influenza H3N8 following treatment with phospholipids. (a) TEM images of H3N8 virus without treatment show filamentous particles. (b) Phospholipid-treated H3N8 reveals observable changes in virus morphology including envelope rupture (scale bar, 200 nm). (c) Detection of H3N8 via FITC-labelled anti-HA antibody following virus binding (MOI 5) for 10 min at 4°C and (d) following further incubation for 30 min at 37°C. (e) Immunocytochemical staining of cells for virus NP 6 h post infection with H3N8 (MOI 1) or following treatment with LPG or DPPG and (f) with each lipid at 5, 50 or 500 µM resulting in a decrease in virus focus forming units that is concentration dependent (*p < 0.05; **p < 0.01; one-way ANOVA). (g) Cytokine transcription levels of IL-8 and TNF-α from H3N8 infected cells (MOI 1) with or without lipid treatment 24 h post infection. Means and standard errors of at least three experiments are represented. The results demonstrate that the infectivity of filamentous viruses is also affected by phospholipids. However, differences were observed and measured between spherical and filamentous viruses that may be related to the amount of membrane involved and the fact that the impact of lipids is diluted when the amount of membrane constituting the virus is larger. (*p < 0.05; **p < 0.01; ***p < 0.001; #p < 0.0001; one-way ANOVA.)
Table 2. Reduction in the percentage of virus infectivity for H2N3, H1N1 and H3N8 in MDCK cells following DPPG or LPG treatment.

| lipid         | concentration (µM) | H2N3                  | H1N1                  | H3N8                  |
|---------------|--------------------|-----------------------|-----------------------|-----------------------|
| virus only    | n.a.               | 563.0 ± 53.0 (100%)   | 678.3 ± 35.4 (100%)   | 688.2 ± 165.0 (100%) |
| DPPG          | 5                  | 301.3 ± 64.3 (53.5%)  | 485.7 ± 21.0 (71.6%)  | 517.1 ± 53.1 (75.1%)  |
|               | 50                 | 226.7 ± 69.3 (40.2%)  | 315.5 ± 41.5 (46.5%)  | 374.8 ± 36.1 (54.4%)  |
|               | 500                | 181.0 ± 86.0 (32.1%)  | 31.3 ± 12.0 (4.6%)    | 356.4 ± 42.6 (52.0%)  |
| LPG           | 5                  | 278.7 ± 76.9 (49.5%)  | 521.0 ± 34.0 (76.8%)  | 486.4 ± 31.0 (70.7%)  |
|               | 50                 | 264.5 ± 71.1 (46.9%)  | 38.2 ± 10.6 (5.6%)    | 101.4 ± 15.5 (14.7%)  |
|               | 500                | 20.0 ± 7.4 (3.5%)     | 9.2 ± 4.6 (1.3%)     | 51.2 ± 22.9 (7.4%)    |

In short, this paper is a call, or plea, to urge physicists to consider viruses as an important and central physical system to study.

4. Methods

4.1. Viruses

AI H2N3 (A/mallard duck/England/7277/06) and pandemic influenza A (H1N1) virus (A/California/7/2009) were kindly provided by Dr Ian Brown (Animal and Plant Health Agency (APHA), UK); a filamentous influenza, equine influenza H3N8 (A/equine/Newmarket/5/03), was kindly provided by Dr Debra Elton (Animal Health Trust, UK). Viruses were propagated in the allantoic fluid of embryonated hens’ eggs. Virus titre was determined by immunocytochemistry as previously described [27] with aliquots stored at −80°C before use.

4.2. Cells

MDCK and A549 cells were grown and maintained in Dulbecco’s modified Eagle’s medium (DMEM, Glutamax; Invitrogen) containing 10% fetal calf serum (FCS; Invitrogen) and 1% penicillin/streptomycin (DMEM-10).

4.3. Lipids

Lipids 1,2-dipalmitoyl-sn-glycero-3-phospho-(1’-rac-glycerol) (DPPG); 1,2-dipalmitoyl-sn-glycero-3-phosphocholine (DPPC); 1-palmitoyl-2-hydroxy-sn-glycero-3-phospho-(1’-rac-glycerol) (LPG); 1-palmitoyl-2-hydroxy-sn-glycero-3-phosphocholine (LPC) (Avanti Polar Lipids, Inc.) were dried under nitrogen flow (Peak Scientific Instruments Ltd, UK) and resuspended in infection medium (IM) comprising serum free Ultraculture medium (Lonza, UK) supplemented with 1% penicillin/streptomycin (Invitrogen), 1% L-glutamine and 2 µg ml⁻¹ tosyl phenylalanyl chloromethyl ketone (Sigma) prior to sonication at 60 kHz for 10 min to create a lipid emulsion.

4.4. Transmission electron microscopy

MDCK cells were infected with H2N3 or H3N8 at an MOI of 1.0 with incubation at 37°C with 5% CO₂ for 72 h. The virus-containing supernatant was then harvested, clarified by centrifugation at 1200g for 2 min, and mixed with phospholipid solutions for 30 min at 37°C with 5% CO₂. The final solutions were then pipetted onto holey carbon support film with 3.05 mm copper grids including 300 hexagonal mesh (EM Resolution, UK) and stained with 2% phospho-tungstic acid (PTA) at pH 7.4 (Sigma, UK). A Tecnai G212 BioTwin Digital TEM system was used to image the samples from different areas of the grids and Image J (Image J 1.47v, National Institute of Health, USA) was used to measure virus aspect ratio and surface area to quantify virus morphology.

4.5. Flow cytometry to determine potential impact of lipids on MDCK cells

MDCK cells were cultured in DMEM-10 until 90-95% confluent. The medium was discarded and cells washed twice with cold PBS at 4°C. Cells were then incubated in verseine (ethylenediamine-tetraacetic acid, Gibco, UK) at 37°C for 10 min to detach them. DMEM-10 was used to neutralize verseine and the cell suspension was centrifuged at 1200g for 6 min. The pellet was resuspended at a concentration of 1 × 10⁶ cells ml⁻¹ in IM. Single cell suspensions were then incubated with phospholipids at the desired concentration for 10 min at 37°C. To minimize the potential toxicity of phospholipids on cells, lipids were then removed by ultrafiltration using an Amicon Ultra-15 filter (50 K MWCO, Fisher Scientific, UK). Cells were then fixed in 1% PFA (Fisher, UK) for 20 min at room temperature and washed three times with cold PBS. The samples were then examined with an FC500 flow cytometer (Beckman Coulter, UK), recording 10,000 events, and the data analysed using Weasel software program v. 3.0 (Walter and Eliza Hall Institute, Australia).

4.6. Measurement of cell metabolic activity

MDCK or A549 cells were seeded in 96 well plates (Nunc) at 5 × 10⁵ cells well⁻¹ with DMEM-10 at 37°C and 5% CO₂ to reach 95% confluence. The supernatant was discarded and the cells washed three times with PBS prior to being incubated with 100 µl of phospholipid-IM solution at 37°C for 30 min, 60 min or 6 h. Triton 1% was used as a positive control. The cell metabolic activity was determined by using MTT reagent (CellTiter 96 Aqueous one solution for cell proliferation assay; Promega) following the manufacturer's instructions. Absorbance was read at 492 nm using a plate reader (Labtech International Ltd).

4.7. Flow cytometry to measure H2N3 binding and uptake using MDCK cells

H2N3 virus (at a concentration to allow infection at MOI of 5.0) was mixed with phospholipid solution at the desired concentration for 30 min at 37°C (5% CO₂) prior to ultrafiltration to remove free lipids. MDCK cells were prepared as previously described and resuspended at a concentration of 1 × 10⁶ cells ml⁻¹ in IM. Prepared cells and lipid-treated virus were then mixed together and incubated at 10 min at 4°C to allow virus binding, followed by 10, 20 or 30 min incubation at 37°C before centrifugation at 1200g for
2 min to recover the cell pellet that was then fixed using 1% PFA for 20 min at 4°C. Fixed cells were then blocked in 1% BSA (Fisher Scientific, UK) for 1 h at room temperature and incubated with specific chicken H3 antiserum (kindly provided by Ian Brown, APHA, UK) for 1 h at room temperature. After three washes with PBS, cells were incubated for 1 h with FITC-conjugated goat anti-chicken IgY (H + L) polyclonal secondary antibody (Novex, UK). Cells were then examined using the flow cytometer as described above.

4.8. Fluorescence microscopy to visualize H2N3 binding and internalization in MDCK and A549 cells and H3N8 binding and internalization in MDCK cells

MDCK or A549 cells were grown to 90–95% confluence on 13 mm glass coverslips and incubated with H2N3 or H3N8 (MOI 5) for 10 min at 4°C to allow virus binding, followed by 10, 20 or 30 min incubation at 37°C. Virus was incubated with phospholipid or IM prior to cell infection. Cells were then incubated with Alexa Fluor 488 (goat anti-chicken IgY (H + L), Invitrogen). Then coverslips were mounted with Prolong Gold Anti-Fade Reagent with 4,6-diamidino-2-phenylindole (DAPI; Life Technologies) and imaged using a Leica DM 5000B epifluorescence microscope.

4.9. Infectivity of influenza A virus pre-treated with phospholipids in MDCK cells or A549 cells

Influenza A viruses H2N3 and H3N8 were pre-treated with phospholipids at a range of concentrations. After ultrafiltration, as described above, the viruses were resuspended in infection medium and incubated with confluent MDCK cells (H2N3 or H3N8) or A549 cells (H2N3) at MOI of 1 for 2 h at 37°C. Cells were then washed three times with PBS at room temperature and incubated for a further 4 h in IM at 37°C before being fixed at 4°C in 1:1 acetone–methanol for 10 min at room temperature. Immunocytochemical staining was performed using a primary mouse monoclonal antibody (AA5H, Abcam, UK) for 1 h to detect influenza viral nucleoprotein which was then visualized with an Envision+ horseradish peroxidase phosphate (HRP) kit (DAB; Dako, Ely, UK) following the manufacturer’s instructions. Finally, the proportion of cells stained for NP was calculated using counts from five fields captured using light microscopy assisted by Image J software, allowing calculation of focus forming units of virus per microlitre.

4.10. Influenza viral and host gene expression by quantitative RT-PCR

Viral RNA was extracted from cells using a QIAamp viral RNA purification kit (Qiagen) according to the manufacturer’s instructions. The reverse transcription reaction of the viral RNA to
Table 3. Primers used for RT-PCR amplification.

| gene name | forward primer (5’-3’) | reverse primer (3’-5’) | species |
|-----------|------------------------|-----------------------|---------|
| 18S       | TGGCGGTCAGAGGTGAAAATT  | TGGCAAATGCTTGGCTTT    | human   |
| 18S       | AGAACAGCCTACACATCC     | CACCGACCTGGCTCCA      | canine  |
| IL-8      | GGTTTGGAGGGGCTGAG      | TTTGCTTGAAGTTCTCAGG   | human   |
| IL-8      | TAAAAGAGCTGAGAACAC     | TTTGAAGTCCATGGCATC    | canine  |
| TNF-α     | AGGAGCAGATCACATTC      | TTAATCTCAGTGCACAG     | human   |
| TNF-α     | TAGCTCATGGTGGACAAAC    | AGTATGAGGAGTACAACCC   | canine  |
| IFN-β     | CCAGGTCAGAAGGAGGCA     | TGGCCAGGTTGAAAGTTC    | canine  |

cDNA was performed using a SensiFAST cDNA Synthesis Kit (Bioline) following manufacturer’s instructions. Amplification of mRNA for IL-8, TNFα and IFN-β using human (for RNA extracted from A549 cells) or canine (for RNA extracted from MDCK cells) gene specific primers was performed using a SensiFAST SYBR No-ROX Kit (Bioline). 18S rRNA was used as a reference gene, previously shown to be suitable for analysis of cells infected with influenza A [28]. All primers were synthesized by Sigma Aldrich (table 3). Influenza Matrix gene (M gene) was quantified using a SensiFAST probe No-ROX Kit (Bioline). A forward primer (5’-3’) AGATGTA GTCTTAA CGG AGT CG and reverse primer TGG AAC ATC TTC AAG TCT, with a hydrolysis probe (5’-3’) [6FAM]TCAGGCCCTCCCTAAAGGCCG AG[BHQ1] were used to amplify a 101 bp region of the M-gene as previously described [29,30].

4.11. Statistical analysis
Statistical analyses were performed using GraphPad Prism Student (v. 6.07). A one-way ANOVA with Tukey’s multiple comparisons test was used to determine significant differences between different treatments. Differences were considered significant for p < 0.05.

Data accessibility. Data are available upon request. Data are available from the electronic supplementary material [31].

Authors’ contributions. L.A.: formal analysis, investigation, methodology, validation, visualization; S.P.D.: conceptualization, data curation, formal analysis, investigation, methodology; C.R.: conceptualization, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, supervision, validation, writing—original draft, writing—review and editing.

All authors gave final approval for publication and agreed to be held accountable for the work performed therein.

Conflict of interest declaration. The authors declare that there are no conflicts of interest.

Funding. L.A.’s PhD studentship was funded by a grant from the Iraqi Ministry of Higher Education and Scientific Research.

Acknowledgements. We would like to thank Denise Christie from the School of Biomedical Science, University of Nottingham, for her support with the EM and Dr David Onion and Nicola Croxall from the School of Life Sciences, University of Nottingham, for their help concerning the flow cytometry work.

References

1. Bernhard W. 2016 Lung surfactant: function and composition in the context of development and respiratory physiology. Ann. Anat. 208, 146–150. (doi:10.1016/j.jaanat.2016.08.003)
2. Buckingham S, Avery ME. 1962 Time of appearance of lung surfactant in the fetal mouse. Nature 193, 688–689. (doi:10.1038/193688a0)
3. Malloy JL, Veldhuizen RAW, Thibodeaux BA, O’Callaghan RJ, Wright JR. 2005 Pseudomonas aeruginosa protease IV degrades surfactant proteins and inhibits surfactant host defense and biophysical functions. Am. J. Physiol. Lung Cell Mol. Physiol. 288, 409–418. (doi:10.1152/ajplung.00322.2004)
4. Han S, Mallampalli RK. 2015 The role of surfactant in lung injury and host defense against pulmonary infections. Ann. Am. Thorac. Soc. 12, 765–774. (doi:10.1513/AnnalsATS.201411-307FR)
5. Penin J, Crouzier D, Spelner D, Debozy JC, Garin D, Crance JM, Favier A-L. 2011 Lung surfactant DPPG phospholipid inhibits vaccinia virus infection. Antiviral Res. 89, 89–97. (doi:10.1016/j.antiviral.2010.11.009)
6. Nakamura M, Kuronuma K, Mitsuzawa H, Voelker DR. 2009 Anti-inflammatory and anti-viral properties of phosphatidylglycerol in lung surfactant. Chem. Phys. Lipids 160, 515–516. (doi:10.1016/j.chemphyslip.2009.06.135)
7. Numata M, Nagashima Y, Moore ML, Berry KZ, Chan M, Kandasamy P, Peebles RS, Murphy RC, Voelker DR. 2013 Phosphatidylglycerol provides short-term prophylaxis against respiratory syncytial virus infection. J. Lipid Res. 54, 2133–2143. (doi:10.1194/jlr.M037077)
8. Numata M, Kandasamy P, Voelker DR. 2012 Anionic pulmonary surfactant liquid regulation of innate immunity. Expert Rev. Respir. Med. 6, 243–246. (doi:10.1586/ers.12.21)
9. Numata M, Chua HW, Dakhamab A, Voelker DR. 2010 Pulmonary surfactant phosphatidylglycerol inhibits respiratory syncytial virus-induced inflammation and infection. Proc. Natl Acad. Sci. USA 107, 320–325. (doi:10.1073/pnas.0909361107)
10. Li Q, Sun X, Liu Y, Vavricka CJ, Qi J, Gao GF. 2012 Structural and functional characterization of neumaminidase-like molecule N10 derived from bat influenza A virus. Proc. Natl Acad. Sci. USA 109, 18 897–18 902. (doi:10.1073/pnas.1211037109)
11. Thaa B, Herrmann A, Veit M. 2009 The polybasic region is not essential for membrane binding of the matrix protein M1 of influenza virus. Virology 383, 150–155. (doi:10.1016/j.virol.2008.10.001)
12. Sukharev S, Sachs F. 2012 Molecular force transduction by ion channels: diversity and unifying principles. J. Cell Sci. 125, 3075–3083. (doi:10.1242/jcs.092353)
13. Osman Z, Scalef P, Stewart S, Rauch C. 2017 Physical and biological characteristics of multi drug resistance (MDR): an integral approach considering pH and drug resistance in cancer. Semin. Cancer Biol. 43, 42–48. (doi:10.1016/j.semcancer.2017.01.002)
14. Rauch C, Farge E. 2000 Endocytosis switch controlled by transmembrane osmotic pressure and phospholipid number asymmetry. Biophys. J. 78, 3036–3047. (doi:10.1016/S0006-3495(00)78642-1)
15. Baba T et al. 2001 Clathrin-dependent and clathrin-independent endocytosis are differentially sensitive to insertion of poly(ethylene glycol)-derivatized cholesterol in the plasma membrane. Traffic 2, 501–512. (doi:10.1034/j.1600-0854.2001.20707.x)
16. Israelachvili JN. 2015 Intermolecular and surface forces, 3rd edn. San Diego, CA: Academic Press.
17. Numata M, Kandasamy P, Nagashima Y, Posey J, Hartskorn K, Woodland D, Voelker DR. 2012 Phosphatidylglycerol suppresses influenza A virus infection. *Am. J. Respir. Cell Mol. Biol.* **46**, 479–487. (doi:10.1165/rcmb.2011-0194OC)

18. Farge E, Devaux PF. 1993 Size-dependent response of liposomes to phospholipid transmembrane redistribution: from shape change to induced tension. *J. Phys. Chem.* **97**, 2958–2961. (doi:10.1021/j100114a022)

19. Bialas KM, Bussey KA, Stone RL, Takimoto T. 2014 Specific nucleoprotein residues affect influenza virus morphology. *J. Virol.* **88**, 2227–2234. (doi:10.1128/JVI.00835-07)

20. Seladi-Schulman J, Steel J, Lowen AC. 2013 The genetics of virus particle shape in equine influenza A virus. *Influenza Other Respir. Viruses* **7**, 81–89. (doi:10.1111/irv.12197)

21. Elton D et al. 2013 The genetics of virus particle shape in equine influenza A virus. *Influenza Other Respir. Viruses* **7**, 81–89. (doi:10.1111/irv.12197)

22. Maeda T, Kawasaki K, Ohnishi SI. 1981 Interaction of influenza virus hemagglutinin with target membrane lipids is a key step in virus-induced hemolysis and fusion at pH 5.2. *Proc. Natl Acad. Sci. USA* **78**, 4133–4137. (doi:10.1073/pnas.78.7.4133)

23. Barman S, Nayak DP. 2007 Lipid raft disruption by cholesterol depletion enhances influenza A virus budding from MDCK cells. *J. Virol.* **81**, 12 169–12 178. (doi:10.1128/JVI.00835-07)

24. Quinteno OA, Wright JR. 2000 Metabolism of phosphatidylglycerol by alveolar macrophages in vitro. *Am. J. Physiol. Lung Cell Mol. Physiol.* **279**, 399–407. (doi:10.1152/ajplung.2000.279.2.L399)

25. Sanders CJ, Doherty PC, Thomas PG. 2011 Respiratory epithelial cells in innate immunity to influenza virus infection. *Cell Tissue Res.* **343**, 13–21. (doi:10.1007/s00441-010-1043-z)

26. Hite RD, Morris PE. 2001 Acute respiratory distress syndrome: pharmacological treatment options in development. *Drugs* **61**, 897–907. (doi:10.2165/00003495-200161070-00001)

27. Kuchipudi SV, Dunham SP, Nelli R, White GA, Coward VJ, Slomka MJ, Brown IH, Chang KC. 2012 Rapid death of duck cells infected with influenza: a potential mechanism for host resistance to H5N1. *Immunol. Cell Biol.* **90**, 116–123. (doi:10.1038/icb.2011.17)

28. Kuchipudi SV et al. 2012 18S rRNA is a reliable normalisation gene for real time PCR based on influenza virus infected cells. *Virul. J.* **9**, 230. (doi:10.1186/1743-422X-9-230)

29. Spackman E, Senne DA, Myers TJ, Bulaga LL, Garber LP, Perdue ML, Leohman K, Daum LT, Suarez DL. 2002 Development of a real-time reverse transcriptase PCR assay for type A influenza virus and the avian H5 and H7 hemagglutinin subtypes. *J. Clin. Microbiol.* **40**, 3256–3260. (doi:10.1128/JCM.40.9.3256-3260.2002)

30. Slomka MJ, Pavlidis T, Coward VJ, Voermans J, Koch G, Hanna A, Banks J, Brown IH. 2009 Validated RealTime reverse transcriptase PCR methods for the diagnosis and pathotyping of Eurasian H7 avian influenza viruses. *Influenza Other Respir. Viruses* **3**, 151–164. (doi:10.1111/j.1750-2659.2009.00083.x)

31. Al-dalawi L, Dunham SP, Rauch C. 2022 Lipid biophysics and/or soft matter-inspired approach for controlling enveloped virus infectivity. Figshare. (https://doi.org/10.6084/m9.figshare.c.5943792)