Prophylactic intranasal administration of a TLR2/6 agonist reduces upper respiratory tract viral shedding in a SARS-CoV-2 challenge ferret model

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ARTICLE INFO
Article History:
Received 24 August 2020
Revised 16 October 2020
Accepted 13 November 2020
Available online xxx

Keywords:
Ferret
COVID-19
SARS-CoV-2
Viral shedding
TLR-2
INNA-051

ABSTRACT
Background: The novel human coronavirus SARS-CoV-2 is a major ongoing global threat with huge economic burden. Like all respiratory viruses, SARS-CoV-2 initiates infection in the upper respiratory tract (URT). Infected individuals are often asymptomatic, yet highly infectious and readily transmit virus. A therapy that restricts initial replication in the URT has the potential to prevent progression of severe lower respiratory tract disease as well as limiting person-to-person transmission.

Methods: SARS-CoV-2 Victoria/01/2020 was passaged in Vero/hSLAM cells and virus titre determined by plaque assay. Challenge virus was delivered by intranasal instillation to female ferrets at 5.0 £10^6 pfu/ml. Treatment groups received intranasal INNA-051, developed by Ena Respiratory. SARS-CoV-2 RNA was detected using the 2019-nCoV CDC RUO Kit and QuantStudioTM 7 Flex Real-Time PCR System. Histopathological analysis was performed using cut tissues stained with haematoxylin and eosin (H&E).

Findings: We show that prophylactic intra-nasal administration of the TLR2/6 agonist INNA-051 in a SARS-CoV-2 ferret infection model effectively reduces levels of viral RNA in the nose and throat. After 5 days post-exposure to SARS-CoV-2, INNA-051 significantly reduced virus in throat swabs (p<0.0001) by up to a 24 fold (96% reduction) and in nasal wash (p=0.0107) up to a 15 fold (93% reduction) in comparison to untreated animals.

Interpretation: The results of our study support clinical development of a therapy based on prophylactic TLR2/6 innate immune activation in the URT, to reduce SARS-CoV-2 transmission and provide protection against COVID-19.

Funding: This work was funded by Ena Respiratory, Melbourne, Australia.
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1. Introduction

Coronaviruses (CoV) are pleomorphic, positive-sense, single-stranded RNA-enveloped viruses, members of the Coronaviridae family, that mainly infect wild animals and cause mild human disease [1]. In addition, seven human CoVs that belong to either the Alphacoronavirus- or Betacoronavirus-genus have now been identified. Four of these human CoVs usually cause a mild, upper respiratory
China. Unlike the highly pathogenic SARS or MERS CoVs, SARS-respiratory tract illness (common cold). Another three novel human CoVs have emerged in the past two decades through transmission to humans via an intermediate animal host [2], and caused outbreaks of significant respiratory morbidity and mortality: in 2003, Severe Acute Respiratory Syndrome (SARS) CoV in China [3], in 2012 Middle Eastern Respiratory Syndrome (MERS) CoV in Saudi Arabia [4] and in December 2019 a novel CoV, SARS-CoV-2, identified in the lower respiratory tract of patients presenting viral pneumonia in Wuhan, China [5]. Unlike the highly pathogenic SARS or MERS CoVs, SARS-CoV-2 infections have spread rapidly around the globe, causing broad spectrum respiratory symptoms, from very mild to severe, life-threatening disease (COVID-19) mostly in at risk populations such as the elderly and those with comorbidities.

As with other respiratory CoVs, SARS-CoV-2 primarily spreads via the airborne route, with respiratory droplets expelled by infected individuals [6]. Virus can be transmitted from symptomatic, as well as pre- or asymptomatic individuals [7,8], with asymptomatic individuals being able to shed virus, and therefore being capable to transmit the disease, for longer than those with symptoms [9]. As with other respiratory viruses such as influenza, recent evidence suggests that, the epithelium of the upper respiratory tract (URT) is the initial site of SARS-CoV-2 infection [10,11]. This is consistent with the abundant nasal epithelial cell expression of the SARS-CoV-2 receptor, angiotensin-converting enzyme 2 (ACE2) and its decreasing expression throughout the lower respiratory tract [11].

A topical treatment of the URT that boosts anti-viral immunity and restricts viral replication is a promising method to promote viral clearance, reduce viral shedding and transmission. The TLRs are key microbe-recognition receptors with a crucial role in activation of host defence and protection from infections and therefore attractive drug targets against infectious diseases [12–14]. Synthetic agonists of the intracellular viral DNA/RNA-recognising TLR molecules, TLR3, TLR7/8 and TLR9, are capable of boosting protective innate immune responses against respiratory viruses [15]. However, their success in the clinic has been limited, due to short-duration of benefit or induction of adverse effects, related to the release of pro-inflammatory cytokines and activation of the type-1 Interferon pathway [16–19]. TLRs expressed on the cell surface such as TLR2 offer an alternative approach. TLR2 dimerizes with TLR1 or TLR6 to recognize a broad variety of commensal and pathogenic microbial molecules and its activation is tightly regulated to maintain immune homeostasis [20]. A series of novel synthetic molecules, named the INNA compounds, have been developed with TLR2/6 agonist properties. Importantly, TLR2/6 agonists of the INNA compound series do not directly activate Type-1 Interferons (unpublished data).

Airways administration of INNA compounds has been shown to protect from lethal influenza virus infection, prevent viral transmission and secondary bacterial superinfections in mouse disease models [21–23]. Intranasal (i.n.) treatment with INNA compounds also reduces viral load and lung inflammation in mouse models of rhinovirus infection (unpublished data). The demonstrated prophylactic benefit is associated with fast TLR2/6-mediated up-regulation of a series of innate immune response elements in airway epithelial cells, defined by early, rapid expression of NF-κB-regulated anti-microbial genes, including IFN-λ and chemokines, that precede immune cell recruitment and support prolonged antiviral defence, suppresses viral load and virus-induced pulmonary inflammation (unpublished data).

Ferret challenge models are commonly used to understand human respiratory virus-induced diseases and to evaluate the efficacy of related vaccines and drugs [24,25]. Use of ferrets is appropriate in the case of SARS-CoV-2 infection, as they express the virus entry ACE2 receptor in their airways [26–28] and SARS-CoV-2 i.n. inoculation in ferrets results in virus replication in the URT and dose-dependent viral shedding [28,29].

To determine whether TLR2/6 agonists are also active against SARS-CoV-2, we used prophylactic i.n. administration of the novel compound INNA-051, in a SARS-CoV-2 challenge ferret model [29].

2. Methods

2.1. Animals and ethics

Twenty-four healthy, female outbred ferrets (Mustela putorius furo) aged 6–8 months were obtained from a UK Home Office accredited supplier. The mean weight at the time of first INNA-051 treatment was 845 g/ferret (range 740–1040 g). Animals were housed in social groups of six prior to and post INNA-051 treatment at Advisory Committee on Dangerous Pathogens (ACDP) containment level 2. 

Evidence before the study

Toll Like Receptors (TLR), the sentinel stimulators of the host immune defence against invading microbes, are recognised as targets for the development of broad-spectrum antivirals. Ena Respiratory’s candidate TLR2/6 agonists, including INNA-051, had been shown to reduce virus levels in the upper respiratory tract and lungs in mouse challenge models of rhinovirus and influenza. When the COVID-19 pandemic was declared, we raised the question “Would priming the innate immunity at the site of infection with a TLR2/6 agonist reduce SARS-CoV-2 virus levels and be of potential use to stop disease progression and community transmission?” We searched PubMed for articles in English before 23 March 2020, using the search terms (“SARS-CoV-2” OR “COVID-19”) AND (“TLR agonists”) in all fields. We found no original research articles. To answer this question, a collaboration was set up between Ena Respiratory and Public Health England, who had established a SARS-CoV-2 virus challenge model in ferrets.

Added value of this study

Our study provides evidence that a TLR2/6 agonist delivered topically to the respiratory tract is highly effective against SARS-CoV-2 (COVID-19) in the ferret challenge model, reducing viral levels in the nose and throat of treated animals by up to 96%. The results of this study support the clinical rationale of using a TLR2/6 agonist as prophylaxis, to reduce SARS-CoV-2 transmission and to protect against COVID-19 disease progression.

Implications of all the available evidence

Vaccination is the most attractive approach for long-term protection against SARS-CoV-2 and other viral respiratory infections that cause serious health complications and spread quickly and widely in the community. In a global push to create SARS-CoV-2 vaccines, there are concerns that vaccines in clinical development might have limited ability to reduce viral shedding and virus transmission in the community, as well as limited efficacy in at risk populations such as elderly individuals. There remains an immediate and pressing need for complementary approaches to stop viral community transmission and disease progression. Adjunct to effective social distancing, face masks and vaccine approaches, priming the innate immunity at the site of infection with a pharmaceutical agent, such as INNA-051, is a promising approach to fight SARS-CoV-2, particularly to health care providers, vulnerable individuals and compromised patient groups.

2. Methods

2.1. Animals and ethics

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Group sizes of 6 ferrets were used to satisfy the UK Home Office approved project licence requirements for reduction, but to allow determination of statistical significance between groups. Animals were transferred to ACDP containment level 3 and housed in pairs post SARS-CoV-2 challenge. Cages met with the UK Home Office Code of Practice for the Housing and Care of Animals Bred, Supplied or Used for Scientific Procedures (December 2014). Access to food and water was *ad libitum* and environmental enrichment was provided. Animals were sedated by intramuscular injection of ketamine/xylazine (17.9 mg/kg and 3.6 mg/kg bodyweight) for administering of treatments, in-life sampling and viral challenge.

All experimental work was conducted in accordance with and under the authority of a UK Home Office approved project licence that had been subject to local ethical review at PHE Porton Down by the Animal Welfare and Ethical Review Body (AWERB) as required by the *Home Office Animals (Scientific Procedures) Act 1986.*

### 2.2. INNA-051 treatment

INNA-051 belongs to a series of closely-related, pegylated synthetic analogues of the diacylated lipopeptide, S-[2,3-bis(palmitoyl oxoy)propyl] cysteine (Pam2Cys) (INNA compound series), with selective TLR2/TLR6 agonist activity. Pam2Cys is inherently insoluble and has been rendered soluble by others through addition of the amino acid motif SK4 [30]. Oligo lysine sequences have, however, been shown to be toxic, albeit at high concentration [31] and to modulate viral infection processes independent of TLR activation [32]. Any off-target effects were mitigated by incorporating polyethylene glycol as a solubilising agent, in the INNA compound series [32]. The EC50s for INNA-051 for the human TLR2/6 receptor is calculated at 40.1 pg/mL of -19 PM. Freeze dried INNA-051 provided by Ena Respiratory, Melbourne, Australia was reconstituted in phosphate buffered saline (PBS) (1 mg/ml) and stored 2–8 °C. Immediately prior to treatment, INNA-051 (1 mg/ml) was further diluted in PBS to the required treatment doses; high dose (100 μg/ml), low dose (20 μg/ml) and mixed dose (20 μg/ml first dose and 100 μg/ml second dose).

### 2.3. Viral challenge inoculum

SARS-CoV-2 Victoria/01/2020 [33] was generously provided by Peter Doherty Institute for Infection and Immunity, Melbourne, Australia at P1 and passaged twice in Vero/hSLAM cells [ECACC Cat# 04091501, RRID:CVCL_037], obtained from the European Collection of Authenticated Cell Cultures (ECACC) PHE, Porton Down, UK. Whole genome sequencing was performed, on the challenge isolate, SISPA protocol and then sequenced using Nanopore as described previously [34]. Virus titre was determined by plaque assay on Vero/E6 cells [ECACC Cat# 85020206, RRID:CVCL_0574]. Challenge substance dilutions were conducted in Phosphate Buffer Saline (PBS). Challenge virus was delivered by intranasal instillation (1.0 ml total, 0.5 ml per nostril) at 5.0 × 10^6 pfu/ml.

### 2.4. Experimental design

Experimental design and viral challenge dose were informed by a previous dose-dependent ferret study [29]. Prior to commencing the experiment, animals were randomly assigned to the four treatment groups, to minimise bias. Assignment was not based on weight or any other notable characteristics. To assign animal IDs to groups, ID numbers were randomised by assigning a value to each using RAND() function in Excel and ordering them low to high. A temperature/ID chip (Bio-Thermo Identichip, Animalcare Ltd, UK) was inserted subcutaneously into the dorsal cervical region of each animal.

INNA-051 was delivered by intranasal instillation (1.0 ml total, 0.5 ml per nostril) to three groups (*n*=6) of ferrets 4 days and 1 day prior to challenge. On each day, group 1 received a high dose [100 μg/ml], group 2 a low dose [20 μg/ml] and group 3 received a 20 μg/ml dose 4 days prior to challenge and a 100 μg/ml dose 1 day before challenge. PBS was delivered to control group ferrets (*n*=6) 4 days and 1 day prior to challenge. Two ferrets each from the high dose, low dose and control groups were scheduled for euthanasia on day 3 (*n*=6) to assess pathology at early stage infection. Remaining ferrets (*n*=18) were scheduled for euthanasia on days 12–14; high and low dose [day 12 = 1, day 13 = 2, day 14 = 1], mixed dose [n=2 days 12–14] and control [n=2 days 12 and 14].

Nasal washes and throat swabs for all ferrets were taken prior to first treatment, at days 1 and 3 pc (*n*=24) and at days 5, 7, 10 and 12 pc for surviving ferrets (*n*=18). At necropsy, tissue samples were taken for histopathology and analysed by PCR. Nasal washes were obtained by flushing the nasal cavity with 2 ml PBS. Cotton throat swabs (Koehler Technische Produkte, VWR) were gently stroked across the back of the pharynx in the tonsillar area and retained in viral transport media (VTM). Throat swabs were processed, and aliquots were stored in AVL at -80 °C until assay.

### 2.5. Clinical and euthanasia observations

Animals were monitored for clinical signs of disease twice daily (approximately 8 hours apart) for the entirety of the experiment. Clinical signs of disease were assigned a score based upon the following criteria. Activity was scored as follows: 0 = alert and playful, 1 = alert, playful when stimulated, 2 = alert, not playful when stimulated, 3 = not alert or playful. No clinical signs were noted throughout the experiment. To meet the requirement of the project license, immobility, neurological signs or a sudden drop in temperature were predetermined automatic euthanasia criteria. Animals were also deemed to have reached a humane endpoint if their body weight was at or below 30% baseline. If any ferret reached any of these three criteria, they were to be immediately euthanised using a UK Home Office approved Schedule 1 procedure. No animals reached these end-points during this study.

Temperature was taken using a microchip reader and implanted temperature/ID chip. Temperature was recorded at each clinical scoring point using the chip to ensure any peak of fever was recorded. Animals were weighed at the same time each day throughout the experiment.

### 2.6. Necropsy procedures

Ferrets were anaesthetised with ketamine/xylazine (17.9 mg/kg and 3.6 mg/kg bodyweight) and exsanguination was effected via cardiac puncture, followed by injection of an anaesthetic overdose (sodium pentobarbitone Dolelethal, Vetquinol UK Ltd, 140 mg/kg). A necropsy was performed immediately after confirmation of death. The left lung was dissected and used for subsequent virology procedures.

### 2.7. RNA extraction

RNA was isolated from nasal wash, throat swabs and lung tissue. Weighed lung tissue was homogenised and inactivated in RLT (Qiagen) supplemented with 1% (v/v) Beta-mercaptoethanol. Tissue homogenate was then centrifuged through a QIAshredder homogeneizer (Qiagen) and supplemented with ethanol as per manufacturer’s instructions. Non-tissue samples were inactivated in AVL (Qiagen) and ethanol. Downstream extraction on all inactivated samples was then performed using the BioSprint™96 One-For-All vet kit (Indical) and Kingfisher Flex platform as per manufacturer’s instructions.
2.8. Quantification of Viral Loads by RT-qPCR

Reverse transcription-quantitative polymerase chain reaction (RT-qPCR) targeting a region of the SARS-CoV-2 nucleocapsid (N) gene was used to determine viral loads and was performed using TaqPath™ 1-Step RT-qPCR Master Mix, CG (Applied Biosystems™), 2019-ncov CDC RUO Kit (Integrated DNA Technologies) and QuantoStudio™ 7 Flex Real-Time PCR System. Sequences of the N1 primers and probe were: 2019-ncov_N1-forward, 5’ GACCC-CAAAATCAGCAGAAAT 3’; 2019-ncov_N1-reverse, 5’TCTGGTACTGC-CAGTTGTAAGTCG 3’; 2019-ncov_N1-probe, 5’ FAM-ACCCCGCAT TACGTTGTCGTTGAG-BHQ1 3’. The quantification standard was in vitro transcribed RNA of the SARS-CoV-2 N ORF (NCBI Reference Sequence: NC_045512.2) with quantification between 1 × 10^1 and 1 × 10^6 copies/μL. Positive samples detected below the lower limit of quantification (LLOQ) were assigned the value of 5 copies/μL, whilst undetected samples were assigned the value of < 2.3 copies/μL, equivalent to the assays lower limit of detection (LLOD).

2.9. Histopathological analysis

Samples from the left cranial and left caudal lung lobe together with nasal cavity, were fixed by immersion in 10% neutral-buffered formalin and processed routinely into paraffin wax. Nasal cavity samples were decalcified using an EDTA-based solution prior to embedding. 4 μm sections were cut and stained with haematoxylin and eosin (H&E) and examined microscopically. In addition, samples were stained using the RNAscope technique to identify the SARS-CoV-2 virus RNA. Briefly, tissues were pre-treated with hydrogen peroxide for 10 min (room temperature), target retrieval for 15 min (98–101 °C) and protease plus for 30 min (40 °C) (Advanced Cell Diagnostics). A V-nCoV2019-S probe (Cat No. 848561, Advanced Cell Diagnostics) was incubated on the tissues for 2 h at 40 °C. Amplification of the signal was carried out following the RNAscope protocol using the RNAscope 2.5 HD Detection kit – Red (Advanced Cell Diagnostics).

2.10. Statistical analysis

Statistical analyses were performed using GraphPad Prism, version 8.4.2 (GraphPad Software, Inc., San Diego, CA). No data was excluded from analysis. Weight change from baseline was compared to vehicle-control Group 4 by two-way ANOVA with Dunnett’s multiple comparisons test. Viral titres from groups 1–3 combined were compared against control group 4 by two-way ANOVA with Dunnett’s multiple comparisons test. In each test a p-value <0.05 was considered statistically significant.

2.11. Role of the funding source

The funders, Ena Respiratory, contributed to study design, interpretation, writing of the manuscript and the decision to publish.

3. Results

3.1. Clinical observations in ferrets

Ferrets received two i.n. administrations of INNA-051 or vehicle-control (PBS), at 4 days before and 1 day prior to i.n. challenge with 5.0 × 10^4 plaque-forming units (PFU) of SARS-CoV-2 in 1 ml volume (day 0). Four groups (6 animals/group) were used: Group 1 received two 1 ml doses (100 μg/ml) INNA-051 (high dose); Group 2 received two 1 ml doses (20 μg/ml) INNA-051 doses (low dose); Group 3 received one 1ml dose (20 μg/ml) INNA-051 at -4 days and one 1 ml dose INNA-051 (100 μg/ml) at -1 day (mixed dose); Group 4 received two doses of vehicle (PBS) alone. After inoculation with SARS-CoV-2, ferrets were monitored for 12 days. In life samples were taken at days 1, 3, 5, 7, 10 and 12, with scheduled culls at days 3 and 12. By 5 dpc, the viral RNA titres from groups 1-3 combined were compared against control group 4 by two-way ANOVA with Dunnett’s multiple comparisons test. Viral titres were excluded from analysis. Weight change from baseline was compared to vehicle-control Group 4 by two-way ANOVA with Dunnett’s multiple comparisons test. Viral titres from groups 1–3 combined were compared against control group 4 by two-way ANOVA with Dunnett’s multiple comparisons test. In each test a p-value <0.05 was considered statistically significant.

Analysis of viral RNA in throat swabs provided further evidence of the capacity of INNA-051 prophylactic treatment, nasal wash and throat swab samples were taken 4 days before viral challenge and at 1, 3, 5, 7, 10 and 12-days post challenge (dpc). Analysis of viral RNA in nasal wash samples at 1 dpc confirmed infection in all treatment groups, with lower viral RNA levels detected in INNA-051 treatment Group 3 (Fig. 2A). Reduction of viral RNA in treatment Group 3 was also evident at 3 dpc (p=0.0155) (Fig. 2A). By 5 dpc, all INNA-051 treated groups had significantly reduced viral RNA compared to the vehicle-control group (2-way ANOVA Dunnett’s multiple comparison test: Group 1 p=0.0244; Group 2 p=0.0107; Group 3 p=0.0071 compared to vehicle-control Group 4) (Fig. 2A). On 5 dpc, the viral RNA levels in the nasal washes of the majority of INNA-051 treated animals remained low or below quantifiable limits throughout the course of infection. Viral RNA levels were found to be below the level of quantification in nasal washes of PBS-treated animals from 10 dpc onwards (Fig. 2A).

Analysis of viral RNA in throat swabs provided further evidence of the capacity of INNA-051 treatment to reduce SARS-CoV-2 in the URT (Fig. 2B). On 3 dpc lower viral RNA levels were found in throat swabs of INNA-051 treated animals, with significantly greater reduction observed (p=0.0345, 2-way ANOVA Dunnett’s multiple comparison) in INNA-051 treatment Group 3. By 5 dpc, all groups treated with INNA-051 had significantly reduced viral RNA levels, compared to the vehicle control group (2-way ANOVA Dunnett’s multiple comparison: Group 1 p=0.0002, Group 2 p=0.0001 and Group 3 p=0.0039 compared to vehicle control Group 4). Highly significantly reduction
Fig. 1. Clinical observations. (a) Schematic of experimental design. Ferrets received INNA-051 and PBS treatments 4 days and 1 day prior to challenge with $5.0 \times 10^6$ pfu/ml SARS-CoV-2. Nasal wash and throat swabs were collected at days 1, 3, 5, 7, 10 & 12 post challenge (p.c.) for all treatment groups and control group. Scheduled culls were performed for 6/24 ferrets on day 3 p.c. (group 1 n=2, group 2 n=2, group 4 n=2) and 18/24 ferrets on days 12-14 p.c. (b) Temperatures were measured twice daily (approximately 8 hours apart), using implanted temperature/ID chips. Mean temperatures +/- standard error of the mean (SEM) are displayed. Temperature dip post SARS-CoV-2 challenge (*) was attributed to sedation. Group 1 animals were first to come out of sedation and therefore recover temperatures. (b) Weight was recorded daily and percentage change from the ferret weight prior to treatment plotted. Mean percentage weight change +/- SEM are displayed. Weight dips on days 4 and 8 p.c. (*) attributed to 2/4 ferrets in group 2 losing weight due to thirst after upturning water bowl overnight. Ferrets in group 3 (2/6 animals) lost weight for same reason on day 8 p.c.
in viral RNA in the throat of INNA-051 treated animals was also apparent on 7 dpc (2-way ANOVA Dunnett’s multiple comparison: Group 1 p=0.0014, Group 2 p=0.0001 and Group 3 p=0.0002 compared to vehicle control Group 4), while by 10 dpc to the end of the study, the levels of viral RNA were below the limit of quantitation in all treatment groups (Fig 2B).

Variation in viral titres of individual animals within groups is an expected outcome of using outbred ferrets and can be seen in a
similar ferret challenge experiments [29]. Additional analysis performed using Pearson’s correlation shows that the correlation between viral titers in the nose and throat for the individual animals is highly significant at days 5 and 7 p.c (supplementary figures).

Because all INNA-051 treatment groups exhibited reduced viral RNA in the nose and throat, we combined these groups into a single data set (supplementary figures) and compared to the group treated with vehicle. Using 2-way ANOVA Sidak’s multiple comparison test, significant reduction in nasal viral RNA was observed at 5 dpc ($p=0.0057$) and highly significant ($p<0.0001$), greater than 10-fold reduction in throat viral RNA was apparent from 5 to 7 dpc following INNA-051 i.n. treatment (Figure S2). After 5 days post-exposure to SARS-CoV-2, animals treated with INNA-051 in group 2 had statistically significant reduction of virus in throat swabs ($p=0.0001$, 2-way ANOVA Sidak’s multiple comparison test) with 24 fold (96% reduction) and nasal wash ($p=0.0107$, 2-way ANOVA Sidak’s multiple comparison test) 15 fold (93% reduction) in this group compared to untreated animals.

These results indicate a similar profile with the protective effects of natural acquired immunity in the ferret challenge model, as observed following re-challenge with SARS-CoV-2 [29].

To assess SARS-CoV-2 detected beyond the URT, lung tissue samples were collected, on scheduled cull day 3 (6/24 animals) and days 12 – 14 (18/24 animals) dpc and analysed for viral RNA levels. On day 3 dpc, two culled ferrets from the control vehicle group had detectable viral RNA levels ($7.42 \times 10^4$ and $2.86 \times 10^9$ copies/ml) (Fig 2C). There was one ferret in Group 1 showing detectable, but below the quantifiable limit, viral RNA, and no other INNA-051 treated ferrets showing detectable viral RNA in lung tissue on day 3 and days 12 – 14 dpc.

4. Discussion

This study provides evidence supporting a novel approach to prevent SARS-CoV-2 transmission, based on reduced viral shedding, following prophylactic i.n administration of INNA-051. Global efforts for prevention of SARS-CoV-2 infection have so far been mostly focused on social distancing and hygiene measures as well as on R&D efforts for the development of vaccines. Our data demonstrate, for the first time, in an in vivo SARS-CoV-2 infection model, that INNA-051 is highly effective at reducing URT viral shedding, providing the potential to control virus transmission and COVID-19 disease.

TLR2 stimulation at mucosal surfaces triggers rapid up-regulation of protective, innate immune defence responses, and also activates counter-regulatory signalling that suppresses development of excessive inflammation and tissue damage and promotes the integrity of local epithelial barrier function [36,37]. In addition, the INNA compounds have been specifically designed to exert TLR2-mediated pharmacological activity on mucosal epithelium, without being systemically absorbed (Ena Respiratory unpublished data), a property that is expected to facilitate their development as safe, antiviral drug candidates. The lack of obvious clinical signs of inflammation following the administration of two doses of INNA-051 administered i.n supports this view. Histopathology from the study indicates that i.n. INNA-051 administration does not exacerbate SARS-CoV-2 pathology in the ferret lung in this setting (Figure S3).

It has been previously shown that i.n administration of an INNA compound in a mouse model of influenza triggers a cascade of innate immune signals that results in reduction of viral load, prevention of lower-respiratory infection and viral transmission between animals [21–23].

Although the ferret SARS-CoV-2 model has limitations and may not represent the severe spectrum of COVID-19 disease, our findings are highly encouraging and indicative of the potential impact i.n. administration of INNA-051 prophylactically may have against SARS-CoV-2 in humans. Though the prophylactic effects of INNA-051 showed statistical significance across all INNA-051 treated groups, further work is needed to determine the optimal dosing. The fact that a significant reduction of URT viral RNA levels was observed in INNA-051-treated outbred ferrets during the peak of viral replication (5–7 days dpc) in this model [29] implies airway immunity priming and enhancement of antiviral host defence. The predictive value of antiviral effectiveness data from respiratory viral infection ferret models and translation into human infectious disease has been established [25,38]. For this reason, the SARS-CoV-2 ferret model has been used, during the current pandemic, to evaluate the therapeutic effect of a number of FDA-approved/repurposed drugs including, lopinavir-ritonavir, hydroxychloroquine sulfate, or emtricitabine-tenofovir [39]. These drugs were found to have no or only modest (~4 fold for emtricitabine-tenofovir) effect against SARS-CoV2 viral replication, as measured by viral titres in nasal wash from the ferrets [39].

Substantial reduction of SARS-CoV-2 viral shedding in the URT and therefore control of respiratory virus transmission may not be easily achievable without potentiation of airways antiviral immune defences [40]. Systemic antiviral drugs, as well as vaccines may not be effective in halting respiratory viral transmission even if they achieve suppression of clinical disease and in fact preliminary results from an experimental study with one of the leading SARS-CoV-2 vaccine candidates (an adenovirus-construct expressing SARS-CoV-2 spike protein) in non-human primates have shown little effect on the virus load in nasal washes [41]. To address these potential limitations, particularly during the urgent circumstances of an epidemic, parallel use of an i.n. administered innate immune modulator with the characteristics of INNA-051 may be highly appropriate to rapidly boost innate immunity at the primary site of respiratory infection which is protective within days of treatment. The use of i.n. INNA-051 for antiviral respiratory prophylaxis therefore offers several additional advantages, including fast-acting protection, and is in contrast to vaccines that take 2–4 weeks to mount a protective response. The limited risk for development of antiviral resistance, the option of self-administration and the non-prohibitive cost for large-scale manufacturing are also especially attractive factors.

In conclusion, this study provides evidence that prophylactic i.n. administration of the TLR2/6 agonist INNA-051 offers a promising approach for prevention and management of SARS-CoV-2 infection that can be used as a stand-alone method of antiviral prophylaxis and is complimentary to potential vaccination programs. This approach is particularly appealing to individuals at elevated risk of community transmission or development of severe disease, including front-line health care workers, vulnerable communities, the elderly, the immunocompromised and those with existing comorbidities.

Funding sources

This work was funded by Ena Respiratory, Melbourne, Australia.

Contributors

N.W.B, B.Y.C, C.D, D.C.J, F.M, P.P, D.T and M.W.C. contributed to the concept and design of the study. P.P and D.T. wrote the manuscript. N.W.B, B.Y.C, C.D, S.D, D.C.J, F.M and M.W.C. provided critical review. P.P co-ordinated the study and performed analysis on data generated. J.D. and M.G.C. provided virus strain. I.H, D.C.J and W.Z contributed to the concept and design of the study. P.P and D.T. wrote the manuscript. N.W.B, B.Y.C, C.D, S.D, D.C.J, F.M. and M.W.C. provided critical review. B.E. C, R,C, C.M.K.H, V.L, D.N, K.A.R and S.T. contributed to the processing of all animal samples at containment level 3. E.R directed histology

P.C. Proud et al. / EBioMedicine 63 (2021) 103153
experiments and performed critical assessment of pathology. M.J.A, N.I.W and R.J.W. performed the extraction and PCR of samples. G.S performed quality control and analytical assistance on PCR data. All authors have read and approved the final version of the manuscript.

Declaration of Competing Interests

Authors report grants from Ena Respiratory, during the conduct of the study. W. Zeng and D.C. Jackson reports grants from Ena Therapeutics, during the conduct of the study. D. Tsitoura, C. Demaison, F. Mercuri, I. Holmes and N.W. Bartlett reports personal fees and other from Ena Therapeutics, outside the submitted work. D.C. Jackson, W. Zeng and B.Y. Chua reports other from Ena Therapeutics, outside the submitted work. Dr. Holmes reports personal fees from Ena Therapeutics, outside the submitted work.

In addition, D. Tsitoura, C. Demaison and F. Mercuri have a patent AU 2020901709 pending to Ena Therapeutics. D.C. Jackson, W. Zeng and C. Demaison have a patent PCT/AU2011/001225 issued to Ena Therapeutics. D.C. Jackson, W. Zeng, I. Holmes and C. Demaison have a patent PCT/AU2020/050660 pending to Ena Therapeutics.

Acknowledgements

The authors gratefully acknowledge the support from the Biological Investigations Group at the National Infection Service, PHE, Porton Down, United Kingdom. Special thanks to Thomas Hender, Carrie Turner, Stephen Findlay-Wilson and Neil McLeod for assisting in providing RT-qPCR data for this work. The authors would like to express their gratitude to Jade Gouriet, Phillip Brown, Karen Gooch and Jemma Paterson for their help in processing of Containment Level 3 in vivo samples. The authors would like to thank Laura Hunter, Chelsea L. Kennard and Francisco J. Salguero for their contribution to the processing and critical review of pathology tissues. The authors are grateful to Michael G. Catton and Julian Druce from Victorian Infectious Disease Reference Laboratory for the generous donation of the SARS-CoV-2 strain. The authors gratefully acknowledge Ena Respiratory for funding this work.

Data sharing statement

The data that support the findings of this study are available on request from the corresponding author for Ena Respiratory, F.M. upon signing a material transfer agreement. The data are not publicly available due to animal study ethical restrictions.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.ebiom.2020.103153.

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Proud, PC; Tsitoura, D; Watson, RJ; Chua, BY; Aram, MJ; Bewley, KR; Cavell, BE; Cobb, R; Dowall, S; Fotheringham, SA; Ho, CMK; Lucas, V; Ngabo, D; Rayner, E; Ryan, KA; Slack, GS; Thomas, S; Wand, NI; Yeates, P; Demaison, C; Zeng, W; Holmes, I; Jackson, DC; Bartlett, NW; Mercuri, F; Carroll, MW

Title:
Prophylactic intranasal administration of a TLR2/6 agonist reduces upper respiratory tract viral shedding in a SARS-CoV-2 challenge ferret model.

Date:
2021-01

Citation:
Proud, P. C., Tsitoura, D., Watson, R. J., Chua, B. Y., Aram, M. J., Bewley, K. R., Cavell, B. E., Cobb, R., Dowall, S., Fotheringham, S. A., Ho, C. M. K., Lucas, V., Ngabo, D., Rayner, E., Ryan, K. A., Slack, G. S., Thomas, S., Wand, N. I., Yeates, P. ..., Carroll, M. W. (2021). Prophylactic intranasal administration of a TLR2/6 agonist reduces upper respiratory tract viral shedding in a SARS-CoV-2 challenge ferret model.. EBioMedicine, 63, pp.103153-. https://doi.org/10.1016/j.ebiom.2020.103153.

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