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Intranasal vaccination with a lentiviral vector protects against SARS-CoV-2 in preclinical animal models

Graphical Abstract

Highlights
- A lentiviral vector encoding Spike shows preclinical efficacy as a COVID-19 vaccine
- Targeting the immune response to the upper respiratory tract provides critical protection
- Intranasal vaccination induces protective mucosal immunity against SARS-CoV-2 in rodents
- Lung anti-Spike IgA responses correlate with protection and reduced inflammation

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In Brief
Ku et al. present a lentiviral vaccination vector that encodes a full-length, membrane-anchored form of SARS-CoV-2 Spike glycoprotein and induces neutralizing antibodies and T cell responses. An intranasal boost strategy triggers a localized immune response in the upper respiratory tract that provides disease protection in mouse and hamster COVID-19 models.
Intranasal vaccination with a lentiviral vector protects against SARS-CoV-2 in preclinical animal models

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SUMMARY

To develop a vaccine candidate against coronavirus disease 2019 (COVID-19), we generated a lentiviral vector (LV) eliciting neutralizing antibodies against the Spike glycoprotein of SARS-CoV-2. Systemic vaccination by this vector in mice, in which the expression of the SARS-CoV-2 receptor hACE2 has been induced by transduction of respiratory tract cells by an adenoviral vector, confers only partial protection despite high levels of serum neutralizing activity. However, eliciting an immune response in the respiratory tract through an intranasal boost results in a >3 log10 decrease in the lung viral loads and reduces local inflammation. Moreover, both integrative and non-integrative LV platforms display strong vaccine efficacy and inhibit lung deleterious injury in golden hamsters, which are naturally permissive to SARS-CoV-2 replication and closely mirror human COVID-19 physiopathology. Our results provide evidence of marked prophylactic effects of LV-based vaccination against SARS-CoV-2 and designate intranasal immunization as a powerful approach against COVID-19.

INTRODUCTION

The new severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) that emerged in late 2019 in Wuhan, China, is extraordinarily contagious and fast spreading across the world (Guo et al., 2020). Compared with the previously emerged SARS or Middle East respiratory syndrome (MERS) coronavirus, SARS-CoV-2 poses an unprecedented threat to global health and has tremendous socio-economic consequences. Therefore, developing effective prophylactic vaccines against SARS-CoV-2 is absolutely imperative to contain the spread of the epidemic and to prevent the development of coronavirus disease 2019 (COVID-19)-associated symptoms, such as deleterious inflammation and progressive respiratory failure (Amanat and Krammer, 2020).

Coronaviruses are enveloped, non-segmented positive-stranded RNA viruses characterized by their envelope-anchored Spike (S) glycoprotein (Walls et al., 2020). The SARS-CoV-2 S (S<sub>CoV-2</sub>) is a (180 kDa) homotrimeric class I viral fusion protein that engages the carboxypeptidase angiotensin-converting enzyme 2 (ACE2) expressed on host cells. The monomer of S<sub>CoV-2</sub> possesses an ectodomain, a transmembrane anchor domain, and a short internal tail. S<sub>CoV-2</sub> is activated by a two-step sequential proteolytic cleavage that initiates fusion with the host cell membrane. After S<sub>CoV-2</sub>-ACE2 interaction, through a conformational reorganization, the extracellular domain of S<sub>CoV-2</sub> is first cleaved at the highly specific furin 682 RRAR685 site (Guo et al., 2020; Walls et al., 2020), a key factor determining the pathological features of the virus and linked to furin ubiquitous expression (Wang et al., 2020). The resulting subunits are...
models provided the proof-of-principle evidence of marked prophylactic effects of LV-based vaccine strategies against SARS-CoV-2 and the requirement for mucosal immunization to reach vigorous protective lung immunity against COVID-19.

RESULTS

Induction of antibody responses by LVs encoding SARS-CoV-2 Spike protein variants

To develop a vaccine candidate able to induce NAbs against S\textsubscript{CoV-2}, we generated LVs harboring, under the transcriptional control of the cytomegalovirus (CMV) immediate-early promoter, codon-optimized sequences encoding (1) S\textsubscript{1} alone (LV::S\textsubscript{1}), (2) the S1-S2 ectodomain without the transmembrane or C-terminal short internal tail (LV::S\textsubscript{1}-S\textsubscript{2}), and (3) the full-length, membrane-anchored form of S (LV::S\textsubscript{FL}), all of which harbor the RBD (Figure 1A; Figure S1), albeit with potential conformational heterogeneities (Yuan et al., 2020). To evaluate the humoral responses induced by these vectors, we immunized C57BL/6 mice (n = 4 per group) by a single intraperitoneal (i.p.) injection of \(1 \times 10^7\) transduction units (TU) of either an LV or a GFP-encoding LV as a negative control. Anti-S\textsubscript{CoV-2} Ab responses were investigated in the sera at weeks 1, 2, 3, 4, and 6 after immunization (Figure 1B). In LV::S\textsubscript{FL} or LV::S\textsubscript{1}-S\textsubscript{2}-immunized mice, anti-S\textsubscript{CoV-2} immunoglobulin G (IgG) was detectable as early as 1 week after immunization and increased progressively until week 6 after immunization to achieve a mean titer (1/dilution) ± SEM of (4.5 ± 2.9) \times 10^6 or (1.5 ± 1) \times 10^6, respectively. In comparison, anti-S\textsubscript{CoV-2} IgG titers were 2 orders of magnitude lower, i.e., (7.1 ± 6.1) \times 10^4, in their LV::S\textsubscript{1}-immunized counterparts. We then evaluated sera for their capacity to neutralize SARS-CoV-2 by using a reliable neutralization assay based on NAb-mediated inhibition of hACE2\textsuperscript{+} cell invasion by non-replicative LV particle surrogates pseudo-typed with S\textsubscript{CoV-2} (Sterlin et al., 2020). Such S\textsubscript{CoV-2}-pseudo-typed LV particles harbor a luciferase reporter gene, which allows quantitation of the hACE2\textsuperscript{+} host cell invasion. In this assay, the luciferase activity is inversely proportional to the neutralization efficiency of NAb present in biological fluids. Fine comparison, by 50% effective concentration (EC\textsubscript{50}) assays, of the sera from LV::S\textsubscript{1}-, LV::S\textsubscript{1}-S\textsubscript{2}-, or LV::S\textsubscript{FL}-immunized mice clearly established that LV::S\textsubscript{FL} was the most potent vector at inducing anti-S\textsubscript{CoV-2} NAb responses (Figure 1C). Moreover, neutralization activities were correlated with anti-S\textsubscript{CoV-2} IgG titers only in the sera of LV::S\textsubscript{FL}-immunized mice (Figure 1D). These results suggest that the conformation of relevant B cell epitopes is likely to be preserved in S\textsubscript{FL} but not necessarily in S\textsubscript{1} or S\textsubscript{1}-S\textsubscript{2} polypeptides, which could be crucial for giving rise to NAb responses. Comparison of the sera from the LV::S\textsubscript{FL}-immunized mice and a cohort of mildly symptomatic infected individuals living in Crêpy-en-Valois, one of the first epidemic zones to appear in France, revealed similar mean neutralizing activities (Figure 1E). These data predict a potentially protective humoral response induced by LV::S\textsubscript{FL}.

LV::S\textsubscript{FL}-immunized C57BL/6 mice (n = 3) also displayed strong anti-S\textsubscript{CoV-2} T cell responses, as detected 2 weeks after immunization by IFN-γ ELISPOT-based epitope mapping applied to splenocytes stimulated with distinct pools of 15-mer peptides spanning the full-length S\textsubscript{CoV-2} (Figure 2A). Significant amounts of responding T cells were detected in 6 of 16
peptide pools. Deconvolution of these positive pools allowed identification of S:256–275 (SGWTAGAAAYYVGYLQPRTF), S:536–550 (NKCVNFNFNGLTGTG), and S:576:590 (VRDPQTLEILDITPC) immunodominant epitopes, giving rise to >2,000 spot forming units (SFU) per $1 \times 10^6$ splenocytes (Figure 2B). These epitopes elicited CD8+ (but not CD4+) T cells, as assessed by intracellular cytokine staining (Figure 2C). The predominant CD8+ phenotype of these T cells is in accordance with the favored orientation of LV-encoded antigens to the MHC-I presentation pathway (Hu et al., 2011). We also identified S:441–455 (LDSKVGGNYNYLYRL), S:671–685 (CASYQTNNSPRRAR), and S:991–1005 (VQIDRLITGRLQSLQ) subdominant epitopes, which gave rise to <2,000 SFU per $1 \times 10^6$ splenocytes in the ELISPOT assay (Figure 2B).

Establishment of a murine model expressing hACE2 in respiratory tracts

Because $S_{CoV-2}$ does not interact well with murine ACE2, wild-type laboratory mice are not permissive to replication of SARS-CoV-2 clinical isolates. Therefore, we sought to develop a murine model in which human ACE2 (hACE2) expression was induced in the respiratory tracts and pulmonary mucosa to evaluate the efficacy of the LV::SFL vaccine. This method has been successfully used to establish the expression of human DPP4 for the study of mouse infection with MERS-CoV (Zhao et al., 2014) and also for hACE2 during the preparation of this manuscript (Sun et al., 2020). We generated an Ad5 vector to deliver the gene encoding hACE2 under the transcriptional control of the CMV promoter (Ad5::hACE2). We first checked in vitro the potential of the Ad5::hACE2 vector to transduce HEK293T cells by reverse-transcriptase PCR (RT-PCR) (Figure 3A). To achieve in vivo transduction of respiratory tract cells, we instilled 2.5 $\times$ $10^9$ infectious genome units (IGU) of Ad5::hACE2 into C57BL/6 mice via intranasal (i.n.) administration. Four days later, hACE2 protein expression was detectable in the lung cell homogenate by western blot (Figure 3B). To get more insights into the in vivo expression profile of a transgene administered under these conditions, we instilled the same dose of an Ad5::GFP reporter
vector into C57BL/6 mice via i.n. administration. As evaluated by flow cytometry, 4 days after instillation, the GFP reporter was expressed not only in the lung epithelial EpCam+ cells but also in lung immune cells, as tracked by the CD45 pan-hematopoietic marker (Figure 3C), showing that through this approach the transduction was efficiently achieved in epithelial cells, although not restricted to these cells.

To evaluate the permissibility of such hACE2-transduced mice to SARS-CoV-2 infection, 4 days after i.n. pretreatment with either Ad5::hACE2 or an empty Ad5 control vector, we inoculated C57BL/6 mice with 1 × 10^5 TCID_{50} (median tissue-culture infectious dose) of a SARS-CoV-2 clinical isolate (BetaCoV/France/IDFO372/2020) via i.n. administration (Lescure et al., 2020). The lung viral loads, determined 2 days post-inoculation (dpi) by RT-PCR and quantitative real-time PCR (qRT-PCR), were as high as (4.4 ± 1.8) × 10^5 copies of SARS-CoV-2 RNA in Ad5::hACE2-pretreated mice but only (6.2 ± 0.5) × 10^5 copies in empty-Ad5-pretreated mice or (4.0 ± 2.9) × 10^5 copies in non-pretreated mice (Figure 3D). In the latter two control groups, these copy numbers corresponded to the input viral RNA, as determined in Ad5::hACE2-pretreated mice inoculated with equivalent amounts of heat-killed viral particles (Figure 3D). The increased viral load from 10^6 copies to >10^9 copies per lung shows the occurrence of active replication of SARS-CoV-2 in vivo. At 4 dpi, the lung viral loads were maintained in Ad5::hACE2-pretreated mice (2.8 ± 1.3 × 10^8 copies), whereas a drop to (1.7 ± 2.3) × 10^8 or (3.9 ± 5.1) × 10^6 copies was observed in empty-Ad5-pretreated or non-pretreated mice, respectively. In Ad5::hACE2-pretreated mice, the viral loads were still detectable at 7 dpi [(1.33 ± 0.9) × 10^6 copies].

Ad5::hACE-2 i.n. instillation induced CD45+ cell recruitment to the lungs (Figure 3E). However, no pro-inflammatory effect was seen with a lower dose of 4 × 10^6 IU/mouse (Figure 3E), which still conferred full permissibility to SARS-CoV-2 replication (Figure 3F), and this dose was therefore chosen for the subsequent experiments described below. In this model of permissive mice, SARS-CoV-2 infection resulted in widespread infiltration of the lung interstitium by mononuclear inflammatory cells, i.e., lymphocytes and macrophages, at 3 dpi (Figure S2). However, in the Ad5::hACE2-pretreated and SARS-CoV-2-inoculated mice, we did not observe any weight loss or clinical signs of disease over the period of 2–7 days after virus inoculation.

These results show that pretreatment of mice with appropriate doses of Ad5::hACE2 can render mice permissive to SARS-CoV-2 replication without inducing Ad5-mediated inflammation, thus providing a valuable model for vaccine or drug studies.

**Intranasal boost with LV::SFL protects strongly against SARS-CoV-2 in mice**

To investigate the prophylactic potential of LV::SFL against SARS-CoV-2, we gave C57BL/6 mice (n = 4 or 5 per group) an i.p. injection of a single dose of 1 × 10^7 TU of LV::SFL or a negative control LV (sham). Seven weeks after immunization, mice were pretreated with Ad5::hACE2 and, 4 days later, inoculated with 1 × 10^5 TCID_{50} of SARS-CoV-2 via i.n. administration (Figure S3A). At 3 dpi, the lung viral loads in LV::SFL-vaccinated mice were reduced by ~6.5 fold, i.e., to a mean ± SD of (5.5 ± 3.8) × 10^8 SARS-CoV-2 RNA copies versus (3.1 ± 1.9) × 10^9 or (4.3 ± 3.0) × 10^7 copies in the un- or sham-vaccinated mice, respectively (Figure S3B). Therefore, a single i.p. LV::SFL injection provided only partial protection in the lung despite intense serum NAB activity.

To further improve the prophylactic effect, we evaluated the prime-boost and prime-target approaches. C57BL/6 mice (n = 4 or 5 per group) were primed via the i.p. route with 1 × 10^7 TU of LV::SFL or a control LV at week 0 and then boosted at week 3 with (1) 1 × 10^7 TU of the same LV via the i.p. route (LV::SFL i.p.-i.p., prime-boost) or (2) 3 × 10^7 TU via the i.n. route (LV::SFL i.p.-i.n., prime-target) to attract the mediators of systemic immunity to the lung mucosa (Figure 4A). Systemic i.p. boosting with LV::SFL resulted in a significant increase in the anti-S_{COV-2} IgG titers, which was more obvious when the binding was evaluated against the foldon-trimerized full-length S (Figure 4B, left) than against the S1 or RBD fragment (Figure S4A). In contrast, mucosal targeting with i.n. LV::SFL did not lead to a statistically significant improvement of anti-S_{COV-2} IgG titers at the systemic level (Figure 4B, left; Figure S4A). In terms of serum neutralization potential, even though a trend to increase was observed after the i.p. or i.n. boost, the differences did not reach statistical significance (Figure 4B, right).

All mice were then pretreated with Ad5::hACE2 and challenged with 3 × 10^5 TCID_{50} of SARS-CoV-2 via i.n. administration 4 weeks after priming. At 3 dpi, the lung viral loads were significantly lower in LV::SFL-i.p.-i.p.-immunized mice (mean ± SD of [2.3 ± 3.2] × 10^6 copies of SARS-CoV-2 RNA) than in sham-vaccinated mice ([13.7 ± 7.5] × 10^6 copies) (Figure 4C). This reduced viral load was similar to that obtained with a single LV::SFL administration (Figure S3B). Most importantly, after i.n. LV::SFL target immunization, a >3 log_{10} decrease in viral load was observed, and a qRT-PCR assay determined that two of five mice harbored undetectable lung viral loads. We also determined the viral loads in the lungs of these mice by performing a plaque-forming unit (PFU) assay (Figure S4C), which confirmed the same trend in reduction. However, the PFU approach underestimates the amounts of infectious viral particles in the lungs of immunized mice because of the high activity of anti-S NABs in the lung homogenates of these animals. Anti-S_{COV-2} IgG was in fact detected in the clarified lung homogenates of the partially (LV::SFL i.p.-i.p.) or fully (LV::SFL i.p.-i.n.) protected mice. In contrast, anti-S_{COV-2} IgA was detectable only in the fully protected LV::SFL i.p.-i.n. mice (Figure 4D). Higher neutralizing activity was detected in the clarified lung homogenates of LV::SFL i.p.-i.n. mice than in their LV::SFL i.p.-i.p. counterparts (Figure 4E). Therefore, increasing the titers of NABs of the IgG

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**Figure 2. Induction of T cell responses by LV::SFL**

C57BL/6 mice (n = 3) were immunized with 1 × 10^7 TU of LV::SFL or a negative control LV via i.p. injection.

(A) Splenocytes collected 2 weeks after immunization were subjected to an IFN-γ ELISPOT with 16 distinct pools of 15-mer peptides spanning the entire S_{COV-2} (1-1273 aa) and overlapping each other by 10 as residues. SFU, spot-forming units.

(B) Deconvolution of the positive peptide pools by ELISPOT applied to splenocytes pooled from three LV::SFL-or control-LV-immunized mice.

(C) Intracellular IFN-γ versus IL-2 staining of CD4+ or CD8+ T splenocytes after stimulation with individual peptides encompassing the immunodominant epitopes.
isotype at the systemic level did not improve the protection against SARS-CoV-2. However, a mucosal i.n. target immunization (with the potential to attract immune effectors to the entry point of the virus to the host organism and to induce local IgA Abs) correlated with the inhibition of SARS-CoV-2 replication.

Because of the importance of innate immune hyperactivity in the pathophysiology of COVID-19 (Vabret et al., 2020), we investigated the lung innate immune cell subsets in the non-infected controls and sham-vaccinated or LV::SFL-vaccinated mice inoculated with SARS-CoV-2 (Figure 5A). At 3 dpi, we detected no difference between the proportions of basophils or natural killer (NK) cells and total lung CD45+ cells among the experimental groups (Figure 5B). In sharp contrast, we detected significantly higher proportions of alveolar macrophages, dendritic cells, neutrophils, and mast cells and a trend toward increased Ly6C+ monocytes and macrophages versus total lung CD45+ cells in sham-vaccinated mice. These observations demonstrate that in this mouse model, lung SARS-CoV-2 loads correlate with the expansion of several inflammation-related innate immune cell subsets, whereas vaccine-mediated protection dampens or prevents the inflammatory reaction.

This was corroborated by the reduced inflammatory cytokine and chemokine contents in the RNA extracted from total lung homogenates of mice immunized with LV::SFL i.p.-i.p. (Figure 5C, top) or LV::SFL i.p.-i.n. (Figure 5C, bottom) and protected against SARS-CoV-2. Significant reductions of TNF-α, TGF-β, IL-1β, IL-12p40, IL-17A, IL-33, CCL2, CCL5, CXCL9, and CXCL10 contents were observed in the lungs of LV::SFL i.p.-i.p.- versus sham-i.p.-i.p.-vaccinated and challenged mice. Significant reductions were notably detected in the IL-1β, IL-6, CCL2, CCL3, and CXCL5 contents in the lungs of LV::SFL i.p.-i.n.- versus sham-i.p.-i.n.-vaccinated and challenged mice. Therefore, the conferred protection also reduced pulmonary inflammation mediated by SARS-CoV-2 infection, as demonstrated by cytometric and RT-PCR approaches.
Intranasal vaccination with LV::SFL strongly protects against SARS-CoV-2 in golden hamsters

Outbred *Mesocricetus auratus*, so-called golden hamsters, provide a suitable pre-clinical model for studying COVID-19 pathology because the ACE2 ortholog of this species interacts productively with SCoV-2 by supporting host cell invasion and viral replication (Sia et al., 2020). We thus investigated in this model the protective effect against SARS-CoV-2 infection of vaccination by LV::SFL and by an integrase-deficient, non-integrative version of this vector (NILV) with the prospect of application in future human clinical trials.

For assessment of the prophylactic effect of vaccination after the prime-boost or prime-target regimen, *M. auratus* hamsters (*n* = 6 per group) were (1) primed with 1 x 10^6 TU of LV::SFL via i.p. injection and boosted at week 5 with 4 x 10^7 TU of LV::SFL via i.n. injection (int LV::SFL i.p.-i.n. low), (2) primed with 1 x 10^7 TU of LV::SFL via i.p. injection and boosted at week 5 with 4 x 10^7 TU of LV::SFL via i.n. injection (int LV::SFL i.p.-i.n. high), or (3) primed with 1 x 10^8 TU of NILV::SFL via intramuscular (i.m.) injection and boosted at week 5 with 1 x 10^9 TU of NILV::SFL via intramuscular (i.m.) injection (NILV::SFL i.m.-i.n.) (Figure 6A). Sham-vaccinated controls received the same amounts of an empty LV via i.p. and then i.n. routes. Strong and comparable anti-SCoV-2 IgG Abs were detected by ELISA in the sera of hamsters from the three vaccinated groups before and after the i.n. boost (Figure 6B). After boosting and targeting, serology detected neutralization activity in all groups, but the highest EC_{50} average was observed in the int LV::SFL i.p.-i.n. high individuals. Such levels were comparable to those detected in COVID-19 cases in humans (Figure 6C). All hamsters were challenged with 0.3 x 10^5 TCID_{50} of SARS-CoV-2 via i.n. injection at week 5. Up to 16% weight loss was progressively reached at 4 dpi in sham-vaccinated individuals, whereas there was a non-significant loss in the

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**Figure 6. Intranasal boost with LV::SFL strongly protects against SARS-CoV-2 in mice**

(A) Timeline of the LV-based prime-boost strategy, followed by Ad5::hACE2 pretreatment and SARS-CoV-2 challenge.

(B) Titers of anti-SCoV-2 IgG as quantitated by ELISA in the sera of C57BL/6 mice primed at week 0 via the i.p. route and boosted at week 3 via the i.p. or i.n. route (left). Titers were determined as a mean endpoint dilution before the boost (week 3) and challenge (week 4). ***p < 0.001, ****p < 0.0001; two-way ANOVA followed by Sidak’s multiple-comparison test. NS, not significant. The neutralization capacity of these sera is indicated as EC_{50} (right). See also Figure S4A.

(C) Lung viral loads at 3 dpi in mice primed (i.p. route) and boosted (i.p. or i.n. route) with LV::SFL. Sham-vaccinated mice received an empty LV. The red line indicates the detection limit. Statistical significance of the differences in the viral loads was evaluated by a two-tailed unpaired t test; *p < 0.0139, ***p < 0.0088.

(D) Titers of anti-SCoV-2 IgG and IgA Abs determined in the clarified lung homogenates by ELISA with foldon-trimerized SCoV-2 for coating. See also Figure S4B.

(E) Neutralizing activity of the clarified lung homogenates as determined for 1/5 dilution. Statistical significance of the difference was evaluated by a Mann-Whitney U test (*p < 0.0159). See also Figure S3.
LV::SFL-vaccinated groups (Figure 6D). As assessed by qRT-PCR, at 2 dpi, the lung viral loads of the int LV::SFL i.p.-i.n. low, int LV::SFL i.p.-i.n. high, and NILV::SFL i.m.-i.n. groups were ~1.2–2 log_{10} lower than those of sham-vaccinated hamsters (Figure 6E). At 4 dpi, the magnitude of viral-load reductions in the vaccinated groups, compared with the sham group, were statistically significant, even if there was one outlier hamster per group (Figure 6E). Using a PFU assay at 2 and 4 dpi, we detected (6.56 ± 1.88) and (2.94 ± 3.16) × 10^6 PFU per lung of sham-vaccinated controls. At 2 dpi, the decrease in viral loads was largely statistically significant in all of the LV::SFL-vaccinated hamsters, and at 4 dpi, with the exception of one NILV::SFL i.m.-i.n. hamster, no vaccinated individuals possessed detectable infectious viral particles (Figure S5A).

As evaluated by qRT-PCR in the total lung homogenates of the protected int LV::SFL i.p.-i.n. low, int LV::SFL i.p.-i.n. high, and NILV::SFL i.m.-i.n. groups, the expression of inflammatory IFN-γ and IL-6 cytokines, anti-inflammatory IL-10 cytokine, and CCL2, CCL3, and CXCL10 chemokines was substantially lower at 4 dpi than in their unprotected sham-vaccinated counterparts (Figure 6F). The other inflammatory mediators tested were not significantly modified (Figure S5B).

In sham-vaccinated and challenged hamsters, overall marked multifocal degenerative changes of the bronchiolar epithelium, moderate effacement of the epithelium (associated with mild to moderate mixed inflammation in the airway lumen), and minimal multifocal interstitial mononuclear cell inflammation were detected at both time points (Figure S6A). At 4 dpi, effacement of the respiratory epithelium and mixed inflammation in the alveoli were more severe; moderate fibrin deposits were also noted. In vaccinated hamsters, regardless of the vaccine dose, lung lesions were clearly of lower incidence and severity (Figure S6B). In these animals, only minimal degenerative lesions and minimal effacement of the bronchiolar epithelium and occasionally mixed inflammation in the airway lumen of alveoli and mononuclear infiltrates in the interstitium were observed. These microscopic findings were similar after vaccination with the int LV or NILV.

In an additional experiment (Figure 7A), we showed that (1) a single i.m. injection of NILV::SFL induced high titers of serum anti-S Abs (Figure 7B) and initiated significant—but partial—levels of protection in the lungs (Figure 7C) and (2) an i.n. boost with NILV::SFL did not improve the serum NAb activity (Figure 7D), induced significantly improved protection against SARS-CoV-2, as determined by the lung viral loads detected via qRT-PCR (Figure 7C) and PFU (Figure S7A), as well as weight loss (Figure S7B), at 4 dpi. At 4 dpi, in sham-vaccinated and challenged hamsters, substantial pulmonary lesions, severe parenchyma inflammation, consolidation of pulmonary parenchyma, marked alteration of the bronchiolar epithelium, and moderate effacement of the bronchiolar epithelium were detected (Figure 7E; Figure S7C). In their NILV::SFL-vaccinated counterparts, boosted or not, pulmonary lesions were clearly less severe (Figures 7E–7G).

Altogether, on the basis of a complete set of virological, immunological, and histopathological data, the LV::SFL vector elicited anti-S CoV NAb and T cell responses and robust protection against SARS-CoV-2 infection in two pertinent animal models, particularly when mucosal i.n. administration was included in the vaccinal scheme.

**DISCUSSION**

Prophylactic strategies are necessary to control SARS-CoV-2 infection, which, 9 months into the pandemic, still continues to spread exponentially without signs of slowing down. It has now been demonstrated that primary infection with SARS-CoV-2 in rhesus macaques leads to protective immunity against re-exposure (Chandrashekar et al., 2020). Numerous vaccine candidates, based on naked DNA (Yu et al., 2020) or mRNA, recombinant protein, replicating or non-replicating viral vectors (including adenoviral Ad5 vector; Zhu et al., 2020), or alum-adjuvanted inactivated virus (Gao et al., 2020), are under active development for COVID-19 prevention. Our immunological rationale for selecting a LV to deliver the gene encoding S_{CoV-2} antigen was based on its potential to induce in situ expression of heterologous genes and on the ability of this immunization platform to elicit sustained humoral and cell-mediated responses. Unique to LVs is the ability to transduce proliferating and non-dividing cells such as dendritic cells (Esslinger et al., 2002; Firat et al., 2002; He et al., 2005) and thereby serve as a powerful vaccination strategy (Belgnon et al., 2009; Buffa et al., 2006; Coutant et al., 2012; Gallinaro et al., 2018; Iglesias et al., 2006) to provoke strong and long-lasting adaptive responses (Cousin et al., 2019; M.-W.K. et al., unpublished data). Notably, in net contrast to many other viral vectors, LVs do not suffer from pre-existing immunity in populations because of their pseudo-typing with VSV-G, to which humans are barely exposed. We recently demonstrated that a single injection of an LV expressing the Zika envelope provides a rapid and durable sterilizing protection against Zika infection (Ku et al., 2020). Our recent comprehensive comparison of LVs to the gold-standard Ad5 immunization vector also documented the superiority of LVs in inducing multifunctional and central memory T cell responses in the mouse model and their strong immunogenicity in outbred rats (M.-W.K. et al., unpublished data), underlining the suitability of LVs for vaccinal applications.

Because laboratory mice are not naturally susceptible to SARS-CoV-2 infection, we set up an *in vivo* transduction murine model in which hACE2 expression is induced in the respiratory...
tracts by an i.n. Ad5::hACE2 pretreatment prior to SARS-CoV-2 inoculation. Our system, very close to a recently published protocol (Sun et al., 2020), confirmed that mice become largely permissive to SARS-CoV-2 replication in the lungs and thus represent a model for assessment of vaccine or drug efficacy against this virus. Even though the Ad5::hACE2 model might not fully mimic the physiological ACE2 expression profile and thus might not reflect all the aspects of the pathophysiology of SARS-CoV-2 infection, it provides a pertinent model for evaluating in vivo the effects of anti-viral drugs, vaccine candidates, various mutations, or genetic backgrounds on the SARS-CoV-2 replication. Using a low dose of Ad5::hACE2, we detected no CD45+ cell recruitments 4 days after instillation, indicative of an absence of Ad5-related inflammation before the inoculation of SARS-CoV-2.

In the transduced mouse model that allows high rates of SARS-CoV-2 replication, vaccination by a single i.p. administration of 1 × 10^7 T U of LV::SFL 6 weeks before virus inoculation was sufficient to inhibit the viral replication by ~6-fold. Compared with a single administration, further boosting via the systemic route did not afford an improved protection rate. However, systemic priming followed by a mucosal boost efficiently inhibited viral replication and prevented lung inflammation. This protection was correlated with high titers of anti-S_{SARS-CoV-2} IgG and IgA and strong neutralization activity in sera. Additional experiments with appropriate transgenic mice or adoptive immune cell transfer will be necessary for identifying the immunological pathways that contribute to disease severity or protection against SARS-CoV-2. NAbS and cell-mediated immunity, very efficaciously induced with the LV-based vaccine candidate, could synergize to block infection and viral replication.

As this manuscript was being prepared, the laboratory of M.S. Diamond reported that i.n. administration of a Chimpanzee adenoviral-36 vector encoding a stabilized form of S_{SARS-CoV-2} protected upper and lower respiratory tracts against SARS-CoV-2 both in BALB/c mice expressing hACE2 in the lung after i.n. delivery of Ad5::hACE2 and in hACE2-expressing transgenic C57BL/6 mice (Hassan et al., 2020). The authors mentioned that immunization and challenge studies in hamsters and non-human primates (NHPs) are planned and that investigations in humans will also be required for evaluating cross-immunity between this Chimpanzee adenoviral-36 vector and adenoviruses circulating in humans. The latter can in fact be a source of pre-existing immunity, which can interfere with the efficacy of Chimpanzee adenoviral-36 vector. No such limitation exists in the case of VSV-G-pseudotyped LV vaccinal vector (Hu et al., 2011), as mentioned above.

Ab-dependent enhancement (ADE) of coronavirus entry into host cells has been evoked as a potential obstacle in vaccination against coronaviruses. With DNA (Yu et al., 2020) or inactivated SARS-CoV-2 virus (Gao et al., 2020) vaccination in macaques, no immunopathological exacerbation has been observed. Long-term observation even after a decrease in Ab titer could be necessary for excluding such a hypothesis. In the case of MERS-CoV, it has been reported that one particular RBD-specific neutralizing monoclonal Ab (Mersmab1) could mediate in vitro ADE of MERS-CoV into host cells by mimicking the viral receptor human DPP4 and inducing conformational rearrangements of S_{MERS} (Wan et al., 2020). We believe that it is difficult to compare the polyclonal Ab response and its paratope repertoire complexity with the singular properties of a monoclonal Ab that cannot be representative of the polyclonal response induced by a vaccine. In addition, very contradictory results from the same team reported that a single-dose treatment with a humanized version of Mersmab1 afforded complete protection of a human transgenic mouse model from lethal MERS challenge (Qiu et al., 2016). Therefore, even with an Ab that might facilitate host-cell invasion in vitro in some conditions, not only is there no exacerbation of the infection in vivo, but there is also a notable protection.

It is noteworthy that all vaccine candidates currently under phase 2 and 3 clinical development are planned to be administered via the i.m. route and will thus induce systemic—but not mucosal—immunity. Our results evidenced a strong correlation between the presence of mucosal anti-S_{SARS-CoV-2} IgA and local NAb activity in the respiratory tracts and very robust pulmonary protection. Therefore, vaccine administration via the i.n. route, which is the main entry door of SARS-CoV-2, has to be taken into account when COVID-19 vaccination protocols are established, especially because this route is non-invasive and particularly suitable for mass vaccination of children and elderly people.

On the basis of largely diverse manifestations of COVID-19 in humans, distinct animal models mimicking various settings of human infection are needed. After SARS–CoV-2 inoculation, rhesus macaques develop transient and mild disease (Muñoz-Fontela et al., 2020). It has been shown that an i.m.-i.m. prime-boost strategy with a DNA-based anti-S_{SARS-CoV-2} vaccine candidate in these animals results in lower infectious virus titers in bronchoalveolar lavages and nasal swabs than do sham controls (Yu et al., 2020). In parallel, golden hamsters develop severe lung pathology and rapid weight loss, which sets a high bar for demonstrating prophylactic efficacy (Muñoz-Fontela et al., 2020). In
Figure 7. Protective efficacy of NILV::SFL in a systemic prime and i.n. boost regimen in hamsters

(A) Timeline of the NILV::SFL prime-boost or prime-target immunization regimen and challenge. (B) Profile of serum anti-S CoV-2 IgG response following a single (i.m.) injection or a prime (i.m.)-boost (i.n.) immunization with NILV::SFL. (C) Lung viral loads at 4 dpi with SARS-CoV-2 in controls or NILV::SFL-vaccinated hamsters. Statistical significance was evaluated by a two-tailed unpaired t test; **p < 0.01. (D) EC50 serum neutralizing titers after the boost-target regimen. (E) Lung histological H&E analysis shown in a heatmap recapitulating the histological scores for each parameter defined in Figure S7C and determined for individuals of various groups at 4 dpi. (F and G) Representative whole-lung section from NILV::SFL i.m.-NILV::SFL i.n. (F) or sham i.m.-sham i.n. (G) hamsters. See also Figure S7.

Prophylactic vaccination is the most cost-effective and efficient strategy against infectious diseases in general and against emerging coronaviruses in particular. Our results firmly establish that the SCoV-2-encoding LV used in a prime-target protocol is a prominent vaccine strategy against COVID-19 and should be considered for future human clinical trials.

Limitations of study

Multiple animal models that mimic various manifestations of human COVID-19 are required. One can hypothesize that the expression of hACE2 in the transduced lungs of Ad5::hACE2-pretreated mice could be inconsistent. However, the routine use of this mouse model in our hands shows that (1) all pretreated animals express hACE2 mRNA (as detected in total lung extracts) and (2) the range of favorable for use in mucosal immunization through the i.n. route, which is essential in immune protection against SARS-CoV-2. Prophylactic vaccination is the most cost-effective and efficient strategy against infectious diseases in general and against emerging coronaviruses in particular. Our results firmly establish that the SCoV-2-encoding LV used in a prime-target protocol is a prominent vaccine strategy against COVID-19 and should be considered for future human clinical trials.

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STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.chom.2020.12.010.

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DECLARATION OF INTERESTS

P.C. is the founder and COO of Theravectys. M.-W.K., P.A., J.L., K.N., F.M., A.N., B.V., F.N., and F.A. are employees of Theravectys. M.-W.K., M.B., P.A., N.E., L.M., and P.C. are inventors of a pending patent directed to a vaccine candidate against SARS-CoV2.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE  | SOURCE                           | IDENTIFIER          |
|----------------------|----------------------------------|---------------------|
| **Antibodies**       |                                  |                     |
| Goat anti-human ACE-2| R&D systems                      | Cat# AF933; RRID: AB_35572 |
| HRP-conjugated anti-goat | Abcam                          | Cat# ab6741         |
| HRP-conjugated anti-β-actin | Abcam                        | Cat# ab197277       |
| Streptavidin-HRP     | Mabtech                          | Cat# 3310-8-1000    |
| Rat anti-mouse IgA-HRP | Southern Biotech                | Cat# 1165-05; RRID: AB_2794657  |
| Rabbit anti-hamster IgG-HRP | Jackson ImmunoResearch Labs   | Cat# M37470         |
| Goat anti-mouse IgG-HRP | Jackson ImmunoResearch Labs  | Cat# 115-035-075; RRID: AB_2338508   |
| Rat anti-IFN-γ, biotin conjugated, clone R4-6A2 | BD Biosciences                     | Cat# 551506; RRID: AB_394224 |
| Rat anti-IFN-γ, unconjugated, clone AN-18 | BD Biosciences                     | Cat# 551309; RRID: AB_394145 |
| Near IR live/dead   | Thermo Fisher Scientific          | Cat# L34976         |
| Rat anti-CD16/CD32, clone 2.4G2 | BD Biosciences                    | Cat# 553142; RRID: AB_394657 |
| APC rat anti-IFN-γ, clone XM1.2 | BD Biosciences                    | Cat# 554413; RRID: AB_398551 |
| APC-H7 rat anti-mouse CD8a, clone 53-6.7 | BD Biosciences                    | Cat# 560182; RRID: AB_1645237 |
| PE-Cy7 rat anti-CD4, clone RM4-5 | BD Biosciences                     | Cat# 552775; RRID: AB_394461 |
| BV605 rat anti-CD45, clone 30-F11 | BD Biosciences                     | Cat# 563053; RRID: AB_2737976 |
| PE rat anti-CD11b, clone M1/70 eBioscience | BD Biosciences                     | Cat# 12-0112-82; RRID: AB2734869 |
| APC rat anti-CD11b, clone M1/70 | BD Biosciences                     | Cat# 553312; RRID: AB_398535 |
| PE-Cy7 hamster anti-CD11c, clone N418 | Thermo Fisher Scientific            | Cat# 25-0114-82; RRID: AB_469590 |
| ef450 hamster anti-CD11c, clone N418 | Thermo Fisher Scientific            | Cat# 48-0114-82; RRID: AB_1548654 |
| BV711 hamster anti-CD103, clone 2E7 | BioLegend                           | Cat# 121435; RRID: AB_2686970 |
| AF700 rat anti-MHC-II, clone M5/114.15.2 | BioLegend                           | Cat# 107621; RRID: AB_493726 |
| PerCP-Cy5.5 rat anti-Ly6C, clone HK1.4 | Thermo Fisher Scientific            | Cat# 45-5932-80; RRID: AB_2723342 |
| APC Hu anti-Ly-6G, clone REA526 | Miltenyi Biotec                     | Cat# 130-107-913; RRID: AB_2652821 |
| PE rat anti-Ly6G, clone 1A8 | BioLegend                           | Cat# 127608; RRID: AB_1186099 |
| FITC rat anti-CD24, clone M1/69 | BD Biosciences                       | Cat# 561777; RRID: AB_10896486 |
| BV421 rat anti-Siglec-F, clone E50-2440 | BD Biosciences                       | Cat# 562681; RRID: AB_2723081 |
| BV421 Mo anti-CD64, clone X54-S7.1 | BioLegend                           | Cat# 139309; RRID: AB_2562694 |
| PE-Cy7 rat anti-CD117, clone 2B8 | BD Biosciences                       | Cat# 561681; RRID: AB_10893022 |
| APC hamster anti-FcR1, clone MAR-1 | BioLegend                           | Cat# 134316; RRID: AB_10640121 |
| AF700 rat anti-NKp46, clone 29A1.4 | BD Biosciences                       | Cat# 561169; RRID: AB_10561840 |
| FITC rat anti-CCR3, clone J073E5 | BioLegend                           | Cat# 144509; RRID: AB_2561608 |
| **Bacterial and virus strains** |                                  |                     |
| LV::S1               | this paper                        | N/A                 |
| LV::S1-S2            | this paper                        | N/A                 |
| LV::SFL              | this paper                        | N/A                 |
| LV::GFP or empty     | this paper                        | N/A                 |
| Non-replicative LV::SFL | this paper                     | N/A                 |
| LV::GFP              | Zennou et al. (2000)              | N/A                 |
| LV::empty            | this paper                        | N/A                 |
| Ad5::hACE2           | this paper                        | N/A                 |
# Cell Host & Microbe

## Article

| **Continued** | **REAGENT or RESOURCE** | **SOURCE** | **IDENTIFIER** |
|---------------|--------------------------|------------|---------------|
| **Non-replicative S<sub>CoV-2</sub> pseudo-typed LV particles** | Anna et al. (2020) | N/A |
| **SARS-CoV-2 (BetaCoV/France/IDF0372/2020)** | clinical isolate; Lescure et al. (2020) | N/A |
| **Chemicals, peptides, and recombinant proteins** | | |
| **Stabilized foldon-trimerized SARS-CoV-2 Spike** | Grzelak et al. (2020) | N/A |
| **S1 monomer** | this paper | N/A |
| **RBD subdomain** | this paper | N/A |
| **S<sub>CoV-2</sub>-derived 15-mer peptide library** | this paper; Hoffmann et al., 2020; http://www.mimotopes.com | N/A |
| **Critical commercial assays** | | |
| **ViraPower Adenoviral Promoterless Gateway Expression Kit** | Thermo Fisher Scientific | ViraPower Adenoviral Promoterless Gateway Expression Kit |
| **Adeno-X rapid Maxi purification kit** | Takara Bio | Cat# 631533 |
| **QIAGEN Viral RNA Mini Kit** | QIAGEN | Cat# 52906 |
| **Experimental models: cell lines** | | |
| **HEK293T cells** | ATCC | HEK293T cells |
| **HEK293T cells transduced to stably express human ACE2** | Grzelak et al. (2020) | N/A |
| **Experimental models: organisms/strains** | | |
| **Mouse: C57BL/6J** | Janvier, Le Genest Saint Isle, France | Cat#000664; RRID: IMSR JAX:000664 |
| **Hamster: *Mesocricetus auratus* golden** | Janvier, Le Genest Saint Isle, France | Cat# HAMST-M79 |
| **Oligonucleotides** | | |
| **pFLAP plasmid: forward 5'-TGG AGG AGA TAT GAG GG-3'** | this paper | N/A |
| **pFLAP plasmid: reverse 5'-CTG CTG CAC TAT ACC AGA CA-3'** | this paper | N/A |
| **gadph gene: forward 5'-TCT CCT CTG ACT TCA ACA GC-3'** | this paper | N/A |
| **gadph gene: reverse 5'-CCC TGC ACT TTT TAA GAG CC-3'** | this paper | N/A |
| **Primers and probes for SARS-CoV-2 load determination** | this paper | Table S1 |
| **Ad5: forward 5'-TTG TGG TTC TTG CAG ATA TGG C-3'** | Gallaher and Berk (2013) | N/A |
| **Ad5: reverse 5'-TCG GAA TCC CGG CAC C-3'** | Gallaher and Berk (2013) | N/A |
| **Ad5: probe 5'-[FAM]-CTC ACC TGC CGC CTC CGT TTC C-[BHQ1]-3'** | Gallaher and Berk (2013) | N/A |
| **Mouse and hamster cytokine and chemokine primers** | this paper | Table S2 |
| **Recombinant DNA** | | |
| **pFlap-ieCMV-WPREm PC** | Zennou et al. (2000) | N/A |
| **pMK-RQ_S-2019-nCoV** | this paper | N/A |
| **Software and algorithms** | | |
| **GraphPad Prism** | GraphPad | v9.0.0 |
| **FlowJo** | FlowJo LLC | v10 |
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Laleh Majlessi (laleh.majlessi@pasteur.fr).

Materials availability
All plasmids and LV generated in this study will be available under MTA for research use, given a pending patent directed to a vaccine candidate against SARS-CoV2.

Data and code availability
The published article includes all datasets generated and analyzed during this study.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice
Female C57BL/6JRj mice were purchased from Janvier (Le Genest Saint Isle, France) and used between the age of 6 and 10 weeks. They were housed under specific pathogen-free conditions in individually ventilated cages during the immunization period and were then transferred into isolator for SARS-CoV-2 inoculation. Mice were housed in individually filtered cages in isolator until the end of the experiment at the Institut Pasteur animal facilities. Mice were vaccinated with the indicated TU of LV and sera were collected at various time points post-immunization to monitor binding and neutralization activities. Previous to i.m. or i.n. instillations, mice were anesthetized by i.p. injection of a mixture of Ketamine (Imalgene, 50 mg/kg) and Xylazine (Rompun, 50 mg/kg).

Hamsters
Male Mesocricetus auratus golden hamsters (Janvier, Le Genest Saint Isle, France) were purchased mature, i.e., 80-90 gr weight. At the beginning of the immunization regimen they weighed 100 to 120 gr. Hamsters were housed under specific pathogen-free conditions in individually ventilated cages during the immunization period and were transferred into isolators for SARS-CoV-2 inoculation. Hamsters were housed in individually filtered cages inside isolators until the end of the experiment at the Institut Pasteur animal facilities. Hamsters were vaccinated with the indicated TU of LV and their sera were collected at the time points indicated after immunization to monitor Ab activities. Prior to i.m. or i.n. injections, hamsters were anesthetized by i.p. injection of Ketamine (Imalgene, 50 mg/kg) and Xylazine (Rompun, 50 mg/kg).

Ethical approval of animal studies
Experimentation on animals was performed in accordance with the European and French guidelines (Directive 86/609/CEE and Decree 87-848 of 19 October 1987) subsequent to approval by the Institut Pasteur Safety, Animal Care and Use Committee, protocol agreement delivered by local ethical committee (CETEA #DAP20007) and Ministry of High Education and Research APAFIS#24627-2020031117362508 v1.

Patient samples
Serum samples were taken from COVID-19 patients at a period where they were asymptomatic, pauci-symptomatic, symptomatic or hospitalized, as recently described elsewhere (Grzelak et al., 2020). Each participant provided written consent to participate in the study, which was approved by the regional investigational review board (IRB; Comité de Protection des Personnes Ile-de-France VII, Paris, France), according to European guidelines and the Declaration of Helsinki. The study was registered as ClinicalTrials.gov (NCT04325646).

Cell culture
HEK293T and Vero cells were cultured in DMEM media (GIBCO Thermo Fisher Scientific) complemented with 10% heat-inactivated serum (Corning), 100 U/mL penicillin and 100 μg/mL streptomycin (GIBCO Thermo Fisher Scientific) and were incubated at 37°C, 5% CO2.

METHOD DETAILS

Construction of transfer pFLAP plasmids coding for SFL, S1-S2, or S1 proteins
A codon-optimized full-length S (1–1273) sequence was amplified from pMK-RQ_S-2019-nCoV and inserted between BamHI and XhoI sites of pFlap-ieCMV-WPREm. Sequences encoding for S1–S2 (1–1211) or S1 (1–681) were amplified by PCR from the pFlap-ieCMV- SFL-WPREm plasmid and sub-cloned into pFlap-ieCMV-WPREm between the BamHI and XhoI restriction sites (Figure S1). Each of the PCR products were inserted between the native human ieCMV promoter and a mutated Woodchuck Post-transcriptional Regulatory Element (mWPRE) sequence, in which the atg starting codon was mutated to avoid transcription of the downstream truncated “X” protein of Woodchuck Hepatitis Virus, in order to improve the vector safety. Plasmids were amplified
in *Escherichia coli* DH5α in Lysogeny Broth supplemented with 50 μg/mL of kanamycin and purified using the NucleoBond Xtra Maxi EF Kit (Macherey Nagel) and resuspended in Tris-EDTA Endotoxin-Free buffer overnight. Plasmid were quantified with a NanoDrop 2000c spectrophotometer (Thermo Fisher Scientific), aliquoted and stored at −20°C. Plasmid DNA were verified by enzymatic digestion and by sequencing the region proximal to the transgene insertion sites.

**Production and titration of LV**

Non-replicative LV were produced in Human Embryonic Kidney (HEK)-293T cells, as previously detailed (Zennou et al., 2000). Briefly, lentiviral particles were produced by transient calcium phosphate co-transfection of HEK293T cells with the vector plasmid pTrIP/SE, a VSV-G Indiana envelope plasmid and an encapsidation plasmid (p8.74 or pD64V for the production of integration-proficient or integration-deficient vectors respectively). Supernatants were harvested at 48 h post-transfection, clarified by 6-min centrifugation at 2500 rpm at 4°C. LV were aliquoted and stored at −80°C. Vector titers were determined by transducing 293T cells treated with aphidicolin. The titer, proportional to the efficiency of nuclear gene transfer, is determined as Transduction Unit (TU)/mL by qPCR on total lysates at day 3 post-transduction, by use of forward 5'-TGG AGG AGG AGA TAT GAG GG-3' and reverse 5'-CTG CAC TAT ACC AGA CA-3' primers, specific to pFLAP plasmid and forward 5'-TCT CCT CTG ACT TCA ACA GC-3' and reverse 5'-CCC TGC ACT TTG TAA GAG CG-3' primers specific to the host housekeeping gene *gadph* as previously described (Iglesias et al., 2006).

**SARS-CoV-2 inoculation**

Hamsters or Ad5:hACE2-pretreated mice were anesthetized by i.p. injection of mixture Ketamine and Xylazine, transferred into a biosafety cabinet 3 where they were inoculated i.n. with BetaCoV/France/IDF0372/2020 SARS-CoV-2 clinical isolate (Lesure et al., 2020), amplified in VeroE6 cells. The strain BetaCoV/France/IDF0372/2020 was supplied by the National Reference Centre for Respiratory Viruses hosted by Institut Pasteur (Paris, France) and headed by Pr. Sylvie van der Werf. The human sample from which strain BetaCoV/France/IDF0372/2020 was isolated has been provided by Dr. X. Lesure and Pr. Y. Yazdanpanah from the Bichat Hospital, Paris, France. During the progression of our experiments, we observed that reducing the input SARS-CoV-2 dose from 1 × 10^6 to 0.3 × 10^6 TCID_{50}/individual did not impact the plateau of the lung viral loads reached at 2–4 dpi. Therefore, in many studies we used the lower dose to mimic better the natural infectious dose. The viral inoculum was contained in 20 μL for mice and in 50 μL for hamsters. Animals were then housed in an isolator in BioSafety Level 3 animal facilities of Institut Pasteur. The organs and fluids recovered from the animals infected with live SARS-CoV-2 were manipulated following the approved standard operating procedures of these facilities.

**Recombinant S\textsubscript{CoV-2} proteins**

Codon-optimized nucleotide fragments encoding a stabilized foldon-trimerized version of the SARS-CoV-2 S ectodomain (aa 1–1208), the S1 monomer (aa 16–681) and the RBD subdomain (aa 331–519) both preceded by a murine Igk leader peptide and followed by an 8xHis Tag were synthesized and cloned into pcDNA3.1/Zeo\textsuperscript{+} expression vector (Thermo Fisher Scientific). Proteins were produced by transient co-transfection of exponentially growing Freestyle 293-F suspension cells (Thermo Fisher Scientific) using polyethylenimine (PEI)-precipitation method as previously described (Lorin and Mouquet, 2015). Recombinant S\textsubscript{CoV-2} proteins were purified by affinity chromatography using the Ni Sepharose Excel Resin according to manufacturer’s instructions (Thermo Fisher Scientific). Protein purity was evaluated by in-gel protein silver-staining using Pierce Silver Stain kit (Thermo Fisher Scientific) following SDS-PAGE in reducing and non-reducing conditions using NuPAGE 3%–8% Tris-Acetate gels (Life Technologies). Purified proteins were dialyzed overnight against PBS using Slide-A-Lyzer dialysis cassettes (10 kDa MW cutoff, Thermo Fisher Scientific). Protein concentration was determined using the NanoDrop One instrument (Thermo Fisher Scientific).

**ELISA**

Ninety-six-well Nunc Polysorp plates (Nunc, Thermo Fisher Scientific) were coated overnight at 4°C with 100 ng/well of purified S\textsubscript{CoV-2} proteins in carbonate-bicarbonate buffer (pH 9.6). The next day, plates were blocked with carbonate buffer containing 1% BSA for 2 h at 37°C. Wells were then washed with PBS containing 0.05% Tween 20 (PBS-T), 1:100-diluted sera or 1:10-diluted lung homogenates in PBS-T containing 1% BSA and four serial ten-to-ten dilutions were added and incubated during 2 h at 37°C. After PBS-T washings, plates were incubated with 1,000-fold diluted peroxidase-conjugated goat anti-mouse IgG (Jackson ImmunoResearch Europe Ltd, Cambridgeshire, United Kingdom) for 1 h. Plates were revealed by adding 100 μL of 3,3',5,5'-tetracolchin benzidine chromogenic substrate (Eurobio Scientific). Following a 30 min incubation, reaction was stopped by adding 100 μL of 2N H2SO4 and optical densities were measured at 450nm/620nm on a PR3100 reader.

**NAb detection**

Serial dilutions of heat inactivated sera or clarified lung homogenates were assayed for NABs via an inhibition assay which uses HEK293T cells transduced to stably express human ACE2 and non-replicative S\textsubscript{CoV-2} pseudo-typed LV particles which harbor the reporter *luciferase firefly* gene, allowing quantitation of the host cell invasion by mimicking fusion step of native SARS-CoV-2 virus (Sterlin et al., 2020). First, 1.5 × 10^5 TU of S\textsubscript{CoV-2} Pseudo-typed LV were pre-incubated, during 30 min at room temperature, in U-bottom plates, with serial dilutions of each serum in a final volume of 50 μL in DMEM-glutamax, completed with 10% heat-inactivated FCS and 100 U/mL penicillin and 100 μg/mL streptomycin. The samples were then transferred into clear-flat-bottom 96-well-black-plates (Corning), and each well received 2 × 10^3 hACE2+ HEK293-T cells, counted in a NucleoCounter NC.200 system (Chemometec, Denmark) contained
in 50 µL. After 2 days incubation at 37°C 5% CO₂, the transduction efficiency of hACE2+ HEK293-T cells by pseudo-typed LV particles was determined by measuring the luciferase activity, using a Luciferase Assay System (Promega) on an EnSpire plate reader (PerkinElmer), as detailed elsewhere (Sterlin et al., 2020). Results are expressed as percentages of inhibition of luciferase activity compared to the maximum of luciferase activity in the absence of NAbs. Sera were heat-inactivated by 30 min incubation at 56°C and were manipulated in BSL2 conditions.

S₅FL T cell epitope mapping

In order to map the immuno-dominant epitopes of S CoV-2, peptides spanning the whole S FL (Mimotopes, Australia) were pooled by 16, each containing 15 aa residues overlapping by 10 aa peptides were dissolved in DMSO at a concentration of 2 mg/mL and diluted before use at 1 µg/mL for ELISPOT or 2-5 µg/mL for ICS (Intracellular Cytokine Staining) in RPMI – 1640 medium supplemented with 10% FCS, 100 U/mL penicillin, 100 µg/mL streptomycin, 1 x 10⁻⁶ M non-essential amino-acids, 1% vol/vol HEPES, 1 x 10⁻³ M sodium pyruvate and 5 x 10⁻⁵ M of β-mercapto-ethanol. IFN-γ ELISPOT and ICS assays were performed as described previously (Bourgine et al., 2018; Sayes et al., 2016). For ICS, cells were acquired in an Attune NxT Flow cytometer (Thermo Fisher Scientific) and the data were analyzed by FlowJo Software (TreeStar).

Generation of Ad5 gene transfer vectors

The Ad5 gene transfer vectors were produced using the ViraPower Adenoviral Promoterless Gateway Expression Kit (Thermo Fisher Scientific). The sequence containing CMV promoter, BamH1/Xho1 restriction sites and WPRE was PCR amplified from the pTRIP-ΔU3CMV plasmid, by use of (1) forward primer, encoding the attB1 in the 5’ end, and (2) reverse primer, encoding both the attB2 and SV40 polyA sequence in the 5’ end. The attB-PCR product was cloned into the gateway pDONR207 donor vector, via BP Clonase reaction. The hACE2 was amplified from a plasmid derivative of hACE2-expressing pcDNA3.1, while gfp was amplified from pTRIP-ΔCMV-ΔgFP-WPRE. The amplified PCR products were cloned into the pDONR207 plasmid via the BamH1 and Xho1 restriction sites. To obtain the final Ad5 plasmid, the pDONR207 vector, harboring hACE2 or gfp genes, was further inserted into pAd/PL-DEST vector via LR Clonase reaction (Figure S8).

The Ad5 virions were generated by transfecting the E3-transcomplementing HEK293A cell line with pAd CMV-GFP-WPRE-SV40 polyA or pAd CMV-hACE2-WPRE-SV40 polyA plasmid by subsequent vector amplification, according to the manufacturer’s protocol (ViraPower Adenoviral Promoterless Gateway Expression Kit, Thermo Fisher Scientific). The Ad5 particles were purified using Adeno-X rapid Maxi purification kit and concentrated with the Amicon Ultra-4 10k centrifugal filter unit. Vectors were resuspended and stored at –80°C in PIPES buffer pH 7.5, supplemented with 2.5% glucose. Ad5 were titrated using a qPCR protocol, as described (Sallaber and Berk, 2013).

Western blot

Expression of hACE2 in the lungs of Ad5::hACE2-transduced mice was assessed by western blotting. One million cells from lung cell suspension were resolved on 4%–12% NuPAGE Bis-Tris protein gels (Thermo Fisher Scientific), then transferred onto a nitrocellulose membrane (Biorad, France). The nitrocellulose membrane was blocked in 5% non-fat milk in PBS-T for 2 h at room temperature and probed overnight with goat anti-hACE2 primary Ab at 1 µg/mL (AF933, R&D systems). Following three wash intervals of 10 min with PBS-T, the membrane was incubated for 1 h at room temperature with HRP-conjugated anti-goat secondary Ab and HRP-conjugated anti-β-actin (ab197277, Abcam). The membrane was washed with PBS-T thrice before visualization with enhanced chemiluminescence via the super signal west femto maximum sensitivity substrate (Thermo Fisher Scientific) on ChemiDoc XRS+ (Biorad, France). PageRuler Plus prestained protein ladder was used as size reference.

Determination of SARS-CoV-2 viral loads in the lungs

Half of each lung lobes were removed aseptically and frozen at –80°C. Organs were thawed and homogenized for 20 s at 4.0 m/s, using lysing matrix M (MP Biomedical) in 500 µL of ice-cold PBS. The homogenization was performed in an MP Biomedical Fastprep 24 Tissue Homogenizer. The homogenates were centrifuged 10 min at 2000 g for further RNA extraction from the supernatants. Particulate viral RNA was extracted from 70 µL of such supernatants using QIaamp Viral RNA Mini Kit (QIAGEN) according to the manufacturer’s procedure. Viral load was determined by measuring the luciferase activity, using a Luciferase Assay System (Promega) on an EnSpire plate reader (PerkinElmer), as detailed elsewhere (Sterlin et al., 2020). Results are expressed as percentages of inhibition of luciferase activity compared to the maximum of luciferase activity in the absence of NAbs. Sera were heat-inactivated by 30 min incubation at 56°C and were manipulated in BSL2 conditions.

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Expression of hACE2 in the lungs of Ad5::hACE2-transduced mice was assessed by western blotting. One million cells from lung cell suspension were resolved on 4%–12% NuPAGE Bis-Tris protein gels (Thermo Fisher Scientific), then RNA was diluted to 10⁹ genome equivalents/mL in RNase-free water containing 100 µg/mL tRNA carrier and used to establish a standard curve in each assay. Thermal cycling conditions were (1) reverse transcription at 55°C for 10 min, (2) enzyme inactivation at 95°C for 3 min, and (3) 45 cycles of denaturation/amplification at 95°C for 15 s, 58°C for 30 s. Products were analyzed on an ABI 7500 Fast real-time PCR system (Applied Biosystems). PFU assay was performed as recently described (Case et al., 2020).
Cytometric analysis of lung cells

Lungs from individual mice were treated with 400 U/mL type IV collagenase and DNase I (Roche) for a 30-min incubation at 37°C and homogenized by use of GentleMacs (Miltenyi Biotec) mAbs. Cells were filtered through 100 μm-pore filters and centrifuged at 1200 rpm during 8 min. Cells were then treated with Red Blood Cell Lysing Buffer (Sigma) and washed twice in PBS. Cells were stained as follows. (1) To detect DC, monocytes, alveolar and interstitial macrophages: Near IR Live/Dead (Invitrogen), FcγII/III receptor blocking anti-CD16/CD32 (BD Biosciences), BV605-anti-CD45 (BD Biosciences), PE-anti-CD11b (eBioscience), PE-Cy7-antiCD11c (eBioscience), BV450-anti-CD64 (BD Biosciences), FITC-anti-CD24 (BD Biosciences), BV711-anti-CD103 (BioLegend), AF700-anti-MHC-II (BioLegend), PerCP-Cy5.5-anti-Ly6C (eBioscience) and APC anti-Ly-6G (Miltenyi) mAbs. (2) To detect neutrophils or eosinophils: Near IR DL (Invitrogen), FcγII/III receptor blocking anti-CD16/CD32 (BD Biosciences), PerCP-Vio700-anti-CD45 (Miltenyi), APC-anti-CD11b (BD Biosciences), PE-Cy7-anti-CD11c (eBioscience), FITC-anti-CD24 (BD Biosciences), AF700-anti-MHC-II (BioLegend), PE-anti-Ly6G (BioLegend), BV421-anti-Siglec-F (BD Biosciences), (3) to detect mast cells, basophils, NK: Near IR LD (Invitrogen), BV605-anti-CD45 (BD Biosciences), PE-anti-CD11b (eBioscience), eF450-anti-CD11c (eBioscience), PE-Cy7-anti-CD117 (BD Biosciences), APC-anti-FcεR1 (BioLegend), AF700-anti-NKp46 (BD Biosciences), FITC-anti-CCR3 (BioLegend), without FcγII/III receptor blocking anti-CD16/CD32. Cells were incubated with appropriate mixtures for 25 min at 4°C, washed twice in PBS containing 3% FCS and then fixed with Paraformaldehyde 4% by an overnight incubation at 4°C. The cells were acquired in an Attune NxT cytometer system (Invitrogen) and data were analyzed by FlowJo software (Treestar, OR, USA).

qRT-PCR detection of inflammatory cytokines and chemokines in the lungs of the mice and hamsters

Lung samples from mice or hamsters were added to lysing matrix D (MP Biomedical) containing 1 mL of TRizol reagent and homogenized at 30 s at 6.0 m/s twice using MP Biomedical Fastprep 24 Tissue Homogenizer. Total RNA was extracted using TRizol reagent (Thermo Fisher Scientific) according to the manufacturer’s procedure. cDNA was synthesized from 4 μg of RNA in the presence of 2.5 μM of oligo(dT)18 primers, 0.5 mM of deoxyribonucleotides, 2.0 U of RNase Inhibitor and SuperScript IV Reverse Transcriptase (Thermo Fisher Scientific) in 20 μL reaction. The real-time PCR was performed on QuantStudio 7 Flex Real-Time PCR System (Thermo Fisher Scientific). Reactions were performed in triplicates in a final reaction volume of 10 μL containing 5 μL of IQ SYBR Green Supermix (BioRad, France), 4 μL of cDNA diluted 1:15 in DEPC-water and 0.5 μL of each forward and reverse primers at a final concentration of 0.5 μM (Table S2). The following thermal profile was used: a single cycle of polymerase activation for 3 min at 95°C, followed by 40 amplification cycles of 15 s at 95°C and 30 s 60°C (annealing-extension step). Mice β-globin or hamster ribosomal protein L18 (RLP18) was used as an endogenous reference control to normalize differences in the amount of input nucleic acid. The average Ct values were calculated from the technical replicates for relative quantification of target cytokines/chemokines.

Lung histopathology

Samples from the lung were fixed in formalin for at least 7 days and routinely embedded in paraffin. Five μm thick paraffin sections were stained with hematoxylin eosin & saffron (HE&S) or HE. Microscopic changes were qualitatively described and when applicable scored semiquantitatively, by histopathologists blinded to the conditions, using (1) distribution qualifiers (i.e., focal, multifocal, locally extensive or diffuse), and (2) a five-scale severity grade, i.e., 1: minimal, 2: mild, 3: moderate, 4: marked and 5: severe.

Quantification and statistical analysis

Statistical parameters are presented as means ± SEM or ± SD and the exact number (n) of mice per group are annotated in the corresponding figure legend and in the text. Statistical analysis of virological and immunological data was performed using GraphPad Prism v9.0.0 (GraphPad Software, CA, USA). Differences between two groups were evaluated by using unpaired two-tailed nonparametric Mann-Whitney U test. For statistical analysis of multiple groups, the two-way ANOVA followed by Sidak’s multiple comparisons test was applied (*p < 0.05; **p < 0.01; ***p < 0.001; ****p < 0.0001; ns, not statistically significant).