Single-dose immunization with a chimpanzee adenovirus-based vaccine induces sustained and protective immunity against SARS-CoV-2 infection

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Abstract

The development of an effective vaccine against SARS-CoV-2, the causative agent of pandemic coronavirus disease-2019 (COVID-19), is a global priority. Here, we present three chimpanzee adenovirus vaccines that express either the full-length spike (ChAdTS-S), or receptor-binding domain (RBD) with two different signal sequences (ChAdTS-RBD and ChAdTS-RBDs). Single-dose intranasal or intramuscular immunization induced robust and sustained neutralizing antibody responses in BALB/c mice, with ChAdTS-S being superior to ChAdTS-RBD and ChAdTS-RBDs. Intranasal immunization appeared to induce a predominately Th2-based response whereas intramuscular administration resulted in a predominately Th1 response. The neutralizing activity against several circulating SARS-CoV-2 variants remained unaffected for mice serum but reduced for rhesus macaque serum. Importantly, immunization with ChAdTS-S via either route induced protective immunity against high-dose challenge with live SARS-CoV-2 in rhesus macaques. Vaccinated macaques demonstrated dramatic decreases in viral RNA in the lungs and nasal swabs, as well as reduced lung pathology compared to the control animals. Similar protective effects were also found in a golden Syrian hamster model of SARS-CoV-2 infection. Taken together, these results confirm that ChAdTS-S can induce protective immune responses in experimental animals, meriting further development toward a human vaccine against SARS-CoV-2.
Introduction

The rapid and global spread of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the causative agent of coronavirus disease-2019 (COVID-19), calls for urgent development of safe, effective and equitably accessible vaccines. Since the release of the genome sequence of SARS-CoV-2 in early January of 2020, scientists and industrial partners around the world have been working tirelessly to develop various vaccines based on traditional and innovative platforms, each of which is expected to result in different safety and efficacy profiles in humans. For example, the mRNA vaccines developed by Pfizer/BioNtech and Moderna have recently been approved for emergency use by several regulatory agencies. The entire pipeline relies on innovative technologies for both antigen design and gene expression, which offer unprecedented speed and flexibility. However, the long-term safety, efficacy and durability of protection require further characterization. The rare-serotype adenovirus-based vaccines, being developed by AstraZeneca/Oxford University (ChAdOX1) and Johnson & Johnson (Ad26), have also incorporated novel features for antigen- and vector optimization. Given their prior experience in similar vector-based vaccines against Middle East Respiratory Syndrome Coronavirus (MERS-CoV) and human immunodeficiency virus type I (HIV-1), this strategy is likely to be superior in large-scale manufacturing, distribution and administration of vaccines, which may practically translate into larger accessibility and greater protection at the population level. Furthermore, killed vaccines have also been developed, demonstrating impressive safety and efficacy profiles in phase III human trials around the world. This classic form of vaccine has a proven record of success in inducing protective immunity in humans against various viral pathogens, although potential risks exist when manufacturing and handling industrial amounts of live infectious particles. Additional vaccine platforms,
based on DNA, recombinant proteins, or novel viral vectors, are also being
developed\textsuperscript{7-10}. The reported vaccine candidates incorporate either the full-length spike (S) protein of
SARS-CoV-2 or only its receptor-binding domain (RBD) as the immunogen, since the
RBD of S plays critical roles in mediating viral entry\textsuperscript{11-13} and inducing a protective
antibody response in infected individuals as well as experimental animals\textsuperscript{14-18}. Similar
to other coronaviruses, the S protein of SARS-CoV-2 consists of a globular S1 domain,
an N-terminal region, a membrane-proximal S2 domain, and a transmembrane
domain\textsuperscript{11}. The RBD, which is located within the S1 domain, determines the host range
and cellular tropism, while the S2 domain mediates membrane fusion\textsuperscript{11}. SARS-CoV-2
infects airway epithelial cells via an interaction of the RBD with the cellular receptor
angiotensin-converting enzyme 2 (ACE2) \textsuperscript{19}. We and others recently resolved the
crystal structure of the SARS-CoV-2 RBD bound to ACE2, which revealed that the
overall ACE2-binding mode is nearly identical to that of the SARS-CoV RBD, which
also utilizes ACE2 as the cellular receptor\textsuperscript{13,19-21}. This suggests that agents capable of
disrupting this binding interaction could serve as candidates to block the entry of
SARS-CoV-2 into target cells. Indeed, both polyclonal and monoclonal antibodies
directed against the RBD and the S protein in general have been shown to inhibit
SARS-CoV-2 infection\textsuperscript{14,18,22}. A large number of neutralizing monoclonal antibodies
that target the RBD of SARS-CoV-2 also provide protection against infection in
experimental animal models\textsuperscript{15,22-25}. Crystal structure analysis of these neutralizing
antibodies revealed their spatial overlaps and competition with ACE2 for binding to
RBD, thereby disrupting the viral cell-entry process\textsuperscript{14,22,25,26}. Importantly, the reported
vaccines that are being investigated or have been approved for emergency use target
either the S protein or the RBD, and have shown to elicit protective immunity in various
animal models and humans, reinforcing the scientific rationale to incorporate S or the RBD as vaccine immunogens.

After weighing the advantages and disadvantages of various strategies, we sought to develop a novel SARS-CoV-2 vaccine based on the less commonly used rare-serotype adenovirus vector ChAdTS, a derivative of replication-defective chimpanzee adenovirus type 68 (AdC68). Apart from low pre-existing immunity in humans, this vector was demonstrated to induce strong and long-lasting immunity against various viral antigens in experimental animals\(^{27-29}\). For example, the parental AdC68 vector was used by our group to develop a vaccine candidate expressing the full-length MERS-CoV S glycoprotein. Single-dose intranasal administration of this vaccine completely protected human DPP4 knock-in mice from lethal MERS-CoV challenge. Passive transfer of immune sera also conferred a survival advantage in lethal challenge mouse models\(^{27}\). This is in great contrast to the immunity induced by other candidate vaccines, for which immunogenicity is short-lived and multiple rounds of immunization are required to induce detectable levels of neutralizing antibodies or to confer protection against viral challenge\(^{1,30}\). Furthermore, recombinant vaccines based on AdC68 and other rare serotype chimpanzee adenoviral vectors, such as ChAd63 and ChAd3, have recently been engineered to express various antigens, with some demonstrating impressive safety and immunogenicity profiles in clinical studies\(^{31-35}\). These unique features and our prior experience in handling the chimpanzee adenovirus vectors provided a critical foundation and rationale for the development of a SARS-CoV-2 vaccine based on ChAdTS.

The immunogen was either the full-length spike (ChAdTS-S), the receptor-binding domain with the cognate signal sequence (ChAdTS-RBD), or the RBD with the secretory signal sequence from mouse IgG (ChAdTS-RBDs). Single-dose intranasal
or intramuscular immunization with these three vaccine candidates induced a robust and sustained neutralizing antibody response in BALB/c mice, with ChAdTS-S being superior to ChAdTS-RBD and ChAdTS-RBDs. ChAdTS-S immunization through the IN route induced a predominately Th2-biased response, while the IM route resulted in a predominately Th1 response. Such biased immune responses are further supported by the analysis of T-cell responses in immunized animals. Importantly, either administration route was able to induce robust and protective immunity against a live-virus challenge in both golden Syrian hamsters and rhesus macaques, supporting the further development of ChAdTS-S into a clinical vaccine against SARS-CoV-2 infection in humans.

**Results**

**Generation and characterization of recombinant ChAdTS vaccines expressing the spike and RBD of SARS-CoV-2** We generated three recombinant ChAdTS vaccines, expressing either the full-length S protein with the original signal peptide (ChAdTS-S), the RBD with the original signal peptide of the S protein (ChAdTS-RBD), or the RBD with a secretory signal peptide (ChAdTS-RBDs) (Fig. 1a). The coding sequences of S or RBD were inserted into the E1 region of the ChAdTS vector under the control of the CMV promoter and terminated with a bovine growth hormone (BGH) polyadenylation signal sequence. HEK293T cells infected with ChAdTS-S, ChAdTS-RBD, ChAdTS-RBDs showed the desired expression of S or RBD according to both western blot (Fig. 1b) and flow cytometry analysis (Fig. 1c). Dose-dependent expression of S, S1, and RBD with the expected molecular weight was detected in all infected cells, while none was found in cells infected with the empty vector ChAdTS
(Fig. 1b), as expected. Surface expression of S and intracellular expression of the
RBD was further confirmed by staining with the RBD-specific mAbs 1F11 and 2F6,
initially isolated from SARS-CoV-2 infected individuals (Fig. 1c). As both of these
mAbs recognize conformational epitopes on the RBD, the positive signals indicated
proper expression and presentation of RBD epitopes by the infected cells. Here again,
dose-dependent expression was also found by both western blot and flow cytometry
analysis, while no signal was detected using the negative control antibody VRC01 (Fig.
1c).

To study the immunogenicity of the recombinant vaccines, we inoculated 18 groups of
six-week-old BALB/c mice (n = 5 per group) with ChAdTS-S, ChAdTS-RBD, or
ChAdTS-RBDs. Each regimen consisted of three single doses (10^{10}, 10^{9}, or 10^{8} vp)
administered either through the intranasal (IN) or intramuscular (IM) route. The
negative control animals only received a dose of 10^{10} vp of the ChAdTS empty vector.
Serum samples were collected every 2 weeks for 8 weeks, and their binding and
neutralizing activities against SARS-CoV-2 pseudovirus were analyzed. As shown in
Fig. 1, all three recombinant vaccines were able to induce strong and durable antibody
responses after a single immunization. Binding affinity for the RBD and neutralizing
activity against pseudovirus became detectable 2 weeks after immunization and
continued to rise throughout the 8-week period, particularly for the ChAdTS-S group.
The speed and magnitude of the antibody response appeared to be more favorable
for those in the IN than in the IM group. Furthermore, dose-dependent responses were
found for all vaccine candidates, although the effect was less pronounced in the IN
group. By week 8 after immunization, the ChAdTS-S animals had an overall average
binding ED50 of 6286.0 in the IN and 8585.4 in the IM group. Among the ChAdTS-
RBD and ChAdTS-RBDs animals, the corresponding values were 6515.4 and 3527.2,
as well as 7384.2 and 3948.5, respectively. However, the ChAdTS-S animals displayed higher neutralizing activities than those vaccinated with ChAdTS-RBD and ChAdTS-RBDs. Among the animals that received the highest dose, ChAdTS-S elicited an average neutralizing ID50 of 1508.5 in the IN group and 1253.9 in the IM group, which were approximately 10-fold higher than in the animals vaccinated with ChAdTS-RBD (ID50 187.5 for IN and 172.5 for IM) and ChAdTS-RBDs (ID50 165.2 for IN and 115.0 for IM). No detectable binding and neutralizing activities were found among the negative control animals. These results indicate that ChAdTS-S, ChAdTS-RBD, and ChAdTS-RBDs are all immunogenic, but ChAdTS-S appeared to be superior in inducing a binding and neutralizing antibody response in BALB/c mice.

ChAdTS-S induced potent and durable immune responses against SARS-CoV-2 in BALB/c mice We next conducted a more in-depth analysis of the antibody and T-cell responses in animals immunized with the highest dose of ChAdTS-S ($10^{10}$ vp) (Fig. 2). Consistent with what was detected in the pseudovirus neutralization assay, the neutralizing activity against live SARS-CoV-2 became detectable 2 weeks after immunization and continued to rise throughout the 8-week period, although the IN group (FRNT ID50 2260.1) demonstrated higher levels than the IM group (FRNT ID50 957.3). No neutralizing activity was detected in animals vaccinated with the empty ChAdTS vector (Fig. 2a), as expected. The IN group also showed an increasing saliva IgA antibody response against S trimer, which was temporally correlated with the serum neutralizing activity against live SARS-CoV-2. However, no such response was detected in the IM group (Fig. 2b). In terms of total serum IgG binding to the S trimer, the IN and the IM groups displayed similar dynamics in terms of speed and magnitude. By week 8 after immunization, the binding ED50 in the IN group reached as high as
19869.4, and that of the IM group reached 23058 (Fig. 2c). Interestingly, the binding profile of IgG subtypes for the S trimer were dramatically different between the two groups. In the IN group, IgG1 was dominant, followed by IgG2a and IgG2b throughout the entire study period (Fig. 2d). By contrast, IgG2a was dominant in the IM group, followed by IgG1 and IgG2b (Fig. 2e). As a result, the average IgG2a/IgG1 ratios in the IN group remained well below 1, while they were above 1 in the IM group, and continued to increase to as high as 2.8 by week 8 after immunization (Fig. 2f). These results suggest that ChAdTS-S immunization through the IN route induced a predominately Th2-biased response, whereas the IM route elicited a predominately Th1 response. Such biased (dichotomous) immune responses are further supported by the analysis of T-cell responses in immunized animals. Splenocytes collected 1 week after immunization were subjected to interferon (IFN)-γ ELISPOT and intracellular cytokine staining (ICS) analysis. In the IM group, a large number of spot-forming cells (593 SFU per $10^6$ splenocytes) were detected after simulation with an overlapping peptide pool of the S protein, whereas those in the IN group (89 SFU per $10^6$ splenocytes) demonstrated no discernable differences from the negative control ChAdTS group (33 SFU per $10^6$ splenocytes) (Fig. 2g). The ICS results also showed the same trend in the number of CD8+ and CD4+ T cells producing IFN-γ, tumor necrosis factor (TNF)-α, and interleukin (IL)-2 (Fig. 2h and i). These results indicate that a single immunization with ChAdTS-S induces strong antibody and cellular immune responses. The IN route appeared to elicit a predominately Th2 immune response, whereas the IM route elicited an immune response that was more biased toward Th1 cells.

To further study how long the elicited antibody response can persist after a single immunization, we intranasally or intramuscularly inoculated 2 groups of six-week-old...
BALB/c mice (n = 10 per group). Within each group, five animals received $10^{10}$ vp of ChAdTS-S and the remaining five were administered $10^{10}$ vp of the empty ChAdTS vector as a negative control. Blood samples were collected every two weeks from the time of immunization until 36 weeks thereafter. As shown in Fig. 2j, the titer of binding antibodies against the S trimer increased dramatically two weeks after immunization, continued to increase and remained at high levels in the ensuing period of time. IN and IM immunization appeared to induce equivalently high levels of binding antibodies without an appreciable decline throughout the experiment. Furthermore, the neutralizing antibody response, measured using the pseudovirus neutralization assay, also demonstrated a similar trend of increase and persistence after a single immunization. IN immunization appeared to induce relatively higher levels of neutralizing antibodies than the IM route, although the differences were not statistically significant. These results indicate that a single immunization with ChAdTS-S through either the IN or IM route is able to induce strong and durable antibody response, a unique and advantageous feature compared to other reported strategies.

ChAdTS-S elicited potent and protective immune responses against SARS-CoV-2 in rhesus macaques A total of eight adult rhesus macaques (6 to 9 years old) were immunized with a single inoculum comprising $10^{11}$ vp of ChAdTS-S either via the IN (n = 4) or the IM route (n=4). Within each group, two animals received ChAdTS-S and the remaining two received the empty ChAdTS vector without any insert (Fig. 3a). From two weeks post vaccination, IgG responses against trimeric S and RBD were detected in all serum samples from the ChAdTS-S-vaccinated animals. In the IN group, the IgG response appeared to be faster and higher than that in the IM group, where a relatively slower but steadily increasing response was observed until challenge (Fig.
Trimeric S-specific IgA antibodies were only detectable in serum samples from animals immunized through the IN route (Fig. 3d). However, the IgA response seemed to be short-lived and exhibited a clear decline at two to four weeks after vaccination. Consistent with the binding activity, serum neutralizing antibodies against the pseudovirus and live SARS-CoV-2 became detectable two weeks post vaccination. The IN group also demonstrated faster and higher levels than the IM group, but experienced a dramatic decline in the ensuing period while the antibody titer in the IM group continued to increase or remained relatively stable up to 8 weeks post vaccination (Figs. 3e and f). In addition, all vaccinated and control animals developed low levels of ChAdTS vector-specific neutralizing antibodies, although the IN vaccinated macaques developed relatively lower levels of anti-vector neutralizing antibodies than the IM vaccinated macaques (Fig. 3g), similar to what was reported in Ad5 immunized animals. Finally, serum cytokine profiling at two weeks post vaccination revealed that the IM-vaccinated animals had detectable levels of IFNγ and IL-2, while the IN animals demonstrated high levels of IL-6 and IL-18 (Fig. 3h). At four weeks post vaccination, almost all these cytokines exhibited a precipitous decline, but IL-6 showed an increase instead (Fig. 3i).

To evaluate the protective potential of ChAdTS-S, immunized rhesus macaques were challenged with $10^6$ plaque-forming units (PFU) of live SARS-CoV-2 by intratracheal inoculation (Fig. 3a). As the animals in the IN and IM groups responded differently to vaccination, they were challenged either at 4 weeks post vaccination (IN group) or 8 weeks post vaccination (IM group) (Fig. 3a). All animals were euthanized on day 7 (IM group) or 8 (IN group) after the challenge. Viral loads in lung tissues and nasal swabs were quantified, and histopathological analysis of lung tissues was conducted. The absolute number of viral genomic RNA (gRNA) and sub-genomic RNA (sgRNA, an
indication of replicating virus) were measured by droplet PCR, which is more sensitive and accurate than conventional RT-PCR\textsuperscript{37}. As shown in Fig. 4a, no detectable levels of gRNA or sgRNA were found in the lung tissues of ChAdTS-S-vaccinated macaques, while in the control group the viral load reached as high as 2.37×10\textsuperscript{8} and 1.17×10\textsuperscript{5} copies per gram, respectively. ChAdTS-S-vaccinated animals also demonstrated transient levels of gRNA or sgRNA in the nasal swabs, while the control animals showed persistent high levels throughout the experiments. For example, one animal in the IN group demonstrated gradually decreasing gRNA and sgRNA levels from day 2 up to day 7 after the challenge, although the gRNA levels were substantially higher than the sgRNA levels. Similarly, one animal in the IM group also had detectable levels of gRNA and sgRNA on day 2 after challenge, but the viral load became undetectable afterwards. This result may indicate that only a small fraction of detected gRNA copies in the nasal swab are relevant to viral replication. Finally, we performed histopathological analysis on the lung sections collected on day 7 or 8 after the viral challenge. As shown in Fig. 4d, the ChAdTS-S-vaccinated macaques in both the IN and IM groups maintained a normal lung structure with mild infiltration of interstitial lymphocytes and macrophages recruited to the alveolar space. By contrast, the control animals showed severe interstitial pneumonia in all lobes, as evidenced by the infiltration of monocytes and lymphocytes in most alveoli, as well as edema in a proportion of alveoli (Fig. 4d). These results indicate that ChAdTS-S was able to elicit potent and protective immune responses against SARS-CoV-2 infection in rhesus macaques.

\textbf{ChAdTS-S protected golden Syrian hamsters from SARS-CoV-2 infection} We went further to evaluate the protective potential of ChAdTS-S in another animal model
of SARS-CoV-2 infection using golden Syrian hamsters. A total of eight hamsters were intramuscularly inoculated with either $10^{10}$ vp of ChAdTS-S (n=4) or saline as control (n=4) and monitored for their serum antibody response 2 and 4 weeks after immunization. At 6 weeks post immunization, the animals were challenged with $10^5$ PFU of SARS-CoV-2 (HKU-13 strain, GenBank accession no: MT835140) and assessed for body weight changes, viral RNA titer and immunohistochemical staining for viral antigens in the lungs up to 4 days after the challenge. As shown in Fig. 5a and 5b, ChAdTS-S elicited a strong antibody response at 2 and 4 weeks after immunization. Immunization protected the animals from body weight loss (Fig. 5c) and significantly reduced the viral RNA load in the lungs compared to the control animals (Fig. 5d). Immunohistochemistry analysis failed to detect viral nucleocapsid protein in lung sections of ChAdTS-S immunized animals, while immunopositivity was abundantly visible and widely distributed along the alveolar epithelia cells in the control group (Fig. 5e, f). Collectively, these results show that ChAdTS-S is able to elicit protective immunity in the golden Syrian hamster model of SARS-CoV-2 infection.

Neutralization of SARS-CoV-2 variants by immune sera elicited by ChAdTS-S To determine whether immune sera from ChAdTS-S-vaccinated animals could neutralize the SARS-CoV-2 variants of concern, we tested their neutralizing activities against 4 pseudoviruses carrying either the wild type Wuhan-Hu-1 strain, UK501Y.V1 variant (GISAID: EPI_ISL_601443), SA501Y.V2 variant (GISAID: EPI_ISL_792681), and BR501Y.V3 variant (GISAID: EPI_ISL_700450). Both SA501Y.V2 and BR501Y.V3 contain three RBD mutations of concern at positions K417, E484 and N501. The immune sera from BALB/c remains largely unaffected in neutralizing the variants, although the average ID50 against BR501Y.V3 (ID50 1076.9 for IM and 2712.4 for IN)
was relatively higher than that to wildtype Wuhan-Hu-1 (ID50 626.9 for IM and 2207.5 for IN) (Fig. 6a). However, the immune sera from the rhesus macaque sera demonstrated 1 to 3.0-fold decline against SA501Y.V2 and BR501Y.V3, although about 1 to 3.7-fold increase was found against UK501Y.V1 (Fig. 6b). Such discrepancy between mice and rhesus macaque may due to different immune responses induced by the same immunogen ChAdTS-S or limited number of animals studied here. Regardless, immunogenicity and protectivity of the vaccines against newly emerged SARS-CoV-2 variants warrant further investigation.

Discussion

SARS-CoV-2 continues to rage around the world with no sign of abating. One contributing factor is the high prevalence of asymptomatic COVID-19 patients, particularly among the young and healthy population38,39. While many public health measures have proven successful in containing viral spread, the ultimate control will rely on the successful development and deployment of safe and effective vaccines against SARS-CoV-2. In this study, we constructed recombinant viral vectors expressing either the S-protein or its RBD based on the rare serotype chimpanzee adenovirus ChAdTS as potential vaccine candidates against SARS-CoV-2 infection. We show that a single IN or IM immunization elicited a robust and sustained neutralizing antibody response in BALB/c mice, with ChAdTS-S being superior to ChAdTS-RBD and ChAdTS-RBDs. IN immunization induced a predominately Th2-biased response, whereas the IM route resulted in a predominately Th1 response in mice. Such biased immune responses are further supported by the analysis of T-cell
responses in the same immunized animals. Interestingly, the neutralizing activity against several circulating SARS-CoV-2 variants remained unaffected for mice serum but reduced for rhesus macaque serum. Such discrepancy between two animal species may reflect different immune responses towards the ChAdTS-S or limited number of animals studied here. Regardless, immunogenicity and protectivity of the vaccines against newly emerged SARS-CoV-2 variants warrant further investigation. Critically, both immunization routes were able to elicit robust and protective immunity in rhesus macaques. Similarly, IM immunization protected golden Syrian hamsters from weight loss, and significantly reduce viral burdens in the lungs and respiratory tract. These results show that ChAdTS-S can induce protective immune responses in experimental animals, confirming that it is a promising candidate for further development of a vaccine against SARS-CoV-2 infection in humans.

A couple of unique aspects of our study can be highlighted here. First and foremost, the ChAdTS vector has desirable properties such as a large capacity for foreign genes and low pre-existing immunity in humans\textsuperscript{30,40,41}. This is clearly superior to many adenoviral vectors that are widely used in vaccine development, such as human adenovirus 5 (HuAd5), whose seroprevalence in the normal human population reaches as high as 75-80\%\textsuperscript{40-44}. This would undoubtedly impose a steep immune barrier for the vector and lead to compromised immunogenicity of the expressed antigen. The only way to overcome such pre-existing immunity is to increase the inoculation dose, but this increases the risk of undesired adverse effects. By contrast, the seroprevalence of ChAdTS is only 0-15\%, and even when it is positive, the serum antibody titer is generally low\textsuperscript{41,43,44}. This unique feature allows enhanced immunogenicity at low doses of antigens expressed by ChAdTS, thus reducing the likelihood of adverse effects. In fact, the parental AdC68 vector has been engineered
to express antigens from a wide range of pathogens such as Ebola virus, HIV-1 and influenza A virus, demonstrating impressive safety and immunogenicity profiles in preclinical studies. In addition, vaccines based on other serotypes of chimpanzee adenovirus demonstrated similar profiles of low pre-existing immunity. In particular, the ChAdOx1 vaccine expressing the full-length of SARS-CoV-2 S protein (AZD1222), developed by AstraZeneca/Oxford University, has already entered phase III human trials and showed promising safety and efficacy against SARS-CoV-2 infection despite some trial participants only receiving a partial dose. Furthermore, the same adenovirus vector expressing the full-length S protein of MERS-CoV was also evaluated in a phase I human trial and demonstrated good safety and tolerability. ChAd3 vectors expressing Ebola Zaire glycoprotein (ChAd3-EBO-Z) elicited strong immune responses in clinical trial participants. These results highlight the favorable safety, tolerability, and immunogenicity profiles of chimpanzee adenovirus vector-based vaccines in humans, which hold great promise for vaccine development.

The second unique aspect of this study is the exceptional durability and strength of the protective immunity elicited by ChAdTS-S in experimental animals. In BALB/c mice, a single IN or IM dose of ChAdTS-S elicited robust and sustained systemic and mucosal immune responses lasting for up to 36 weeks post immunization. Such high and persistent levels of antibodies are superior to many vaccines for which immunogenicity is short-lived and multiple rounds of immunization are required to induce a detectable and protective neutralizing antibody response. While the underlying mechanism is currently unclear, it is possible that the broad host-cell tropism of ChAdTS and the expression of membrane-anchored instead of soluble S protein by the recombinant ChAdTS-S played critical roles. Interestingly, IN
immunization appeared to induce relatively faster and stronger mucosal and systemic antibody responses than IM immunization, although a bias toward the Th2 response was noted. The mucosal response is particularly relevant for protection against SARS-CoV-2 infection, since viral transmission and replication largely occurs in the upper respiratory tract. Intramuscular immunization, however, appeared to induce a more durable antibody response and higher levels of Th1 cells than intranasal immunization. We are currently uncertain whether these differences translate into different levels of protection in mice. However, it is reassuring to find that a single IN or IM dose of ChAdTS-S elicited protective immune responses against SARS-CoV-2 challenge in rhesus macaques. ChAdTS-S-vaccinated macaques had no detectable levels of SARS-CoV-2 gRNA or sgRNA in the lung tissues, with only transient and low levels in the nasal swabs. By contrast, the control animals showed persistent high levels of viral RNA throughout the experiment. ChAdTS-S-vaccinated animals also maintained normal lung structure, while the control animals showed severe interstitial pneumonia in all lobes. Finally, single-dose intramuscular immunization of golden Syrian hamsters with ChAdTS-S also induced robust and protective immunity, which prevented body weight loss and inhibited viral replication in lung tissues according to viral RNA loads and immunohistochemical staining for the SARS-CoV-2 NP protein. Taken together, these results highlight that ChAdTS-S can elicit potent and protective immune responses against SARS-CoV-2 in experimental animals, meriting further translational research toward a clinical vaccine against SARS-CoV-2 infection in humans.

Materials and Methods
Construction, rescue, amplification and purification of ChAdTS-based vaccines

The ChAdTS, derived from AdC68, is an E1- and E3-deleted replication-deficient adenoviral vector. It also has a partial deletion in E4, which was replaced by the corresponding E4 region of the human adenovirus 5 AdHu5\textsuperscript{51}. The codon-optimized gene encoding the spike (S) protein from SARS-CoV-2 (Wuhan-Hu-1, GenBank: MN908947.3) was synthesized by Tsingke Biological Technology, China. The coding sequence of the RBD region, which spans the residues Arg319–Phe541 of the S protein, was obtained by PCR from the codon-optimized spike gene. The full-length S protein, RBD with the original signal peptide from the S protein, and RBD with the secretory signal peptide from the mouse Ig heavy chain were inserted into the E1-deleted region of the ChAdTS vector by isothermal assembly to obtain the vectors pChAdTS-S, pChAdTS-RBD, and pChAdTS-RBDs, respectively. The primer pair for ChAdTS-S included the forward primer (5′-gctagcgtttaacgggccGCCCCGGGCTTTATAAgccaccATGTTCCTGCTGACCACC-3′) and the reverse primer (5′-agcggtttaaacgggccGCCCCGGGCTTTATAACTAGGTGTAGTGCAAGCTTCA-3′). The primer pair for ChAdTS-RBD included the forward primer (5′-gctagcgtttaacgggccGCCCCGGGCTTTATAAgccaccATGTTCGTGTTCCTGGTGCT-3′) and the reverse primer (5′-agcggtttaaacgtgtaGCCCCGGGCTTTATAACTAGGTGTAGTGCAAGCTTCA-3′). The primer pair for ChAdTS-RBDs included the forward primer (5′-gctagcgtttaacgggccGCCCCGGGCTTTATAAgccaccATGGGATGGTCATGTATC-3′) and the reverse primer (5′-agcggtttaaacgggccGCCCCGGGCTTTATAACTAGGAATCCACGCACCTTAT-3′). Empty ChAdTS with no insertion in the E1 deletion region was employed as the negative
control vaccine. The vectors pChAdTS-S, pChAdTS-RBD, and pChAdTS-RBDs were linearized and transfected into HEK293 cells (ATCC) to rescue ChAdTS-S, ChAdTS-RBD, and ChAdTS-RBDs. Theses recombinant adenoviruses as well as empty control ChAdTS were propagated and purified by CsCl gradient ultracentrifugation before quantification by spectrophotometry\textsuperscript{51,52}.

Detection of S and RBD protein expression HEK293T cells cultured in 6-well plates were infected with ChAdTS-S, ChAdTS-RBD, ChAdTS-RBDs, and ChAdTS at doses of $10^{10}$, $10^9$, and $10^8$ vp per well. After 24 h, the cells were harvested and lysed with 100 μL of buffer containing a protease inhibitor cocktail. The cell lysates were subjected to SDS-PAGE, followed by western blotting with a primary anti-SARS-CoV-2 S antibody (Sino Biological, China) and detected using an HRP-conjugated secondary anti-rabbit IgG (Promega, USA). Beta-actin was used as a loading control.

In the flow cytometry assay, S and RBD expression was detected by surface and intracellular staining, respectively. Two human monoclonal antibodies (P2C-1F11 and P2B-2F6)\textsuperscript{14} specific for the RBD isolated from SARS-CoV-2 infected individuals were incubated with infected cells at a final concentration of 10 μg/mL at 4°C for 30 min. After extensive washing, the cells were further incubated with anti-human IgG-PE (BioLegend, USA) at a 1:50 dilution and analyzed using a BD Calibur FACS instrument (BD, USA). The VRC01 antibody specific for human immunodeficiency virus type I (HIV-1), was used as a negative control.

Mouse immunization and sample collection A total of 120 female BALB/c mice aged 6–8 weeks were randomly divided into 24 groups ($n = 5$ in each group) and immunized with a single dose comprising $10^{10}$, $10^9$, or $10^8$ vp of ChAdTS-S, ChAdTS-
RBD, ChAdTS-RBDs, or ChAdTS via either the intranasal (IN) or intramuscular (IM) route. Sera were collected every 2 weeks until 8 weeks following the immunization, heat-inactivated at 56°C for 30min, and stored at -80°C before analysis for SARS-CoV-2 specific antibodies. T cell responses were measured in the spleens of 20 additional female BALB/c mice one week after immunization. To further study the durability of the elicited antibody response, 20 additional female BALB/c mice aged 6–8 weeks were intranasally or intramuscularly immunized with $10^{10}$ vp of ChAdTS-S and followed up to 36 weeks. Blood samples were collected every 2 weeks, heat-inactivated at 56°C for 30min, and stored at -80°C before analysis for SARS-CoV-2-specific binding and neutralizing antibodies.

**Rhesus macaque immunization and challenge with live SARS-CoV-2**

Eight adult rhesus macaques aged between 5 and 9 years were intranasally or intramuscularly vaccinated with $10^{11}$ vp of ChAdTS-S or empty ChAdTS. Peripheral blood samples were collected every 2 weeks for antibody and cytokine profiling. The vaccinated animals were intratracheally challenged with $10^6$ PFU of live SARS-CoV-2 after 4 weeks after intranasal or 8 weeks after intramuscular immunization. Nasal swabs were collected on days 0, 1, 2, 3, 5, and 7 after the viral challenge. The animals were euthanized on either day 7 or 8 after the challenge and the lungs were collected for viral load analysis and histopathological examination. Viral gRNA and sgRNA in the lung tissues and nasal swabs were measured by droplet digital PCR\textsuperscript{37}, using a COVID-19 digital PCR detection kit (TargetingOne, China). The kit allows the detection of the ORF1ab gene, N gene, and a positive reference gene. The limit of detection is 3 copies/test for ORF1ab gene and 5 copies/test for N gene. The same detection kit, with different primers and probes to target the E gene was used for the detection of
sgRNA. For histopathological analysis, lung tissues were collected and fixed in 10% neutral buffered formalin, embedded in paraffin, and sectioned (5 μm) for standard hematoxylin and eosin staining.

Golden Syrian hamster immunization and challenge with live SARS-CoV-2

The entire procedure was performed as described previously. Briefly, 6- to 8-week-old male and female hamsters were obtained from the Chinese University of Hong Kong Laboratory Animal Service Centre through the HKU Laboratory Animal Unit and kept in Biosafety Level-2 (BSL-2) housing until virus challenge in the BSL-3 animal facility. The hamsters (n=4 per group) were immunized with ChAdTS-S or saline as control via the IM route. Six weeks later, each hamster was intranasally challenged with a single dose comprising $10^5$ PFU of live SARS-CoV-2 (HKU-13 strain, GenBank accession no: MT835140) under intraperitoneal ketamine (200 mg/kg) and xylazine (10 mg/kg) anesthesia. The weights of the hamsters were monitored daily and the animals were sacrificed at day 4 post-challenge. Half of each hamster’s lung was used for viral load determination using a quantitative SARS-CoV-2 RdRp/β-actin reverse transcription-polymerase chain reaction assay. For immunostaining, lung tissues were fixed and incubated with a monoclonal antibody against SARS-CoV-2 nucleocapsid (Sino Biological, China).

ELISA of binding antibodies in the sera of immunized mice, rhesus macaques and golden Syrian hamsters

For the immunized mice, the serum samples were serially diluted and added to 96-well plates pre-coated with recombinant SARS-CoV-2 S trimer or RBD produced in HEK 293F cells (100 ng/well). After incubation at 37°C for 1 h, the plates were washed three times with phosphate-buffered saline containing
0.1% Tween® 20 (PBST) and incubated with a secondary horseradish peroxidase (HRP)-conjugated antibody against mouse IgG (1:4000, Promega, USA), IgG1, IgG2a, IgG2b, or IgA (1:40000; Abcam, UK) at 37°C for 1 h. The samples were further washed 3 times with PBST before the substrate TMB (3’,3’,5’,5’-tetramethyl benzidine) was added and the reaction was stopped by adding 1M H₂SO₄. Absorbance at 450nm was measured using an ELISA plate reader. The ED50 value was calculated based on binding curves drawn in Prism 8.0 software (GraphPad Inc., USA). For immunized rhesus macaques, the IgG response specific to the SARS-CoV-2 S trimer or RBD was measured using an HRP-conjugated anti-monkey IgG (1:6000; Southern Biotech, USA), but otherwise same as the protocol for the immunized mice (see above). The endpoint antibody titer of the macaque sera was defined as the optical value three times above that of the naïve serum. For immunized golden Syrian hamsters, the antibody response was measured using an HRP-conjugated anti-hamster IgG (Invitrogen, USA).

Pseudovirus and live virus neutralization assays Neutralizing titers of the immunized sera were determined using SARS-CoV-2 pseudovirus and live virus neutralization assays as previously reported. The pseudovirus was generated by co-transfection of HEK293T cells with the HIV backbone expressing firefly luciferase (pNL43R-E-luciferase) and pcDNA3.1 (Invitrogen, USA) encoding S proteins from SARS-CoV-2 variants. The wild type S protein was from Wuhan-Hu-1, GenBank: MN908947.3. The variant UK501Y.V1 (GISAID: EPI_ISL_601443) was constructed with mutations Δ69-70, Δ144, N501Y, A570D, D614G, P681H, T716I, S982A and D1118H. The variant SA501Y.V2 (GISAID: EPI_ISL_792681) was constructed with mutations L18F, D80A, D215G, Δ242-244, S305T, K417N, E484K, N501Y, D614G,
A701V and R246I. The variant BR501Y.V3 (GISAID: EPI_ISL_700450) was constructed with mutations L18F, T20N, P26S, D138Y, R190S, K417N, E484K, N501Y, D614G, H655Y, T1027I and V1176F. After 48h, the cell supernatant containing the pseudovirus was collected, measured and stored at -80°C until further use. Serum samples were serially diluted 3-fold in 96-well cell culture plates before SARS-CoV-2 pseudovirus was added and incubated at 37°C for 1h. Approximately 1.5 × 10^4 Huh7 cells were then added to the serum-pseudovirus mixture and incubated at 37°C for an additional 60 h. The ID50 values were calculated based on the relative light units (Bright-Glo Luciferase Assay Vector System, Promega, USA) using Prism 8.0 (GraphPad Software Inc., USA). For the live virus assay, we used live SARS-CoV-2 initially isolated from an infected patient in China and the focus reduction neutralization was performed in a certified BSL3 facility at Shenzhen Third People’s Hospital, China. Briefly, serial dilutions of sera were mixed with SARS-CoV-2 and incubated for 1 h at 37°C. The mixtures were then transferred to 96-well plates seeded with Vero E6 cells and incubated for 1 h at 37°C. After changing the medium, the plates were incubated at 37°C for an additional 24 h. The cells were then fixed, permeabilized, and incubated with cross-reactive rabbit anti-SARS-CoV-N IgG (Sino Biological, Inc., China) for 1 h at room temperature before adding an HRP-conjugated goat anti-rabbit IgG antibody (Jackson ImmunoResearch, USA). The reactions were developed using KPL TrueBlue peroxidase substrate (Seracare Life Sciences Inc., USA). The number of SARS-CoV-2 foci was quantified using an EliSpot reader (Cellular Technology Ltd. USA).

Adenovirus neutralization assay The levels of chimpanzee adenovirus-specific neutralizing antibodies were measured as described previously^42. Serum samples
were 3-fold serially diluted in 96-well cell culture plates and mixed with 1500 TCID$_{50}$ of ChAdTS-GFP vector, which expresses green fluorescent protein. After incubation at 37 °C for 1 h, approximately $1.5 \times 10^4$ HEK293 cells were added to the antibody-virus mixture and co-cultured for an additional 24 h. The green fluorescent protein levels were examined using an Opera Phenix (PerkinElmer, USA) to determine the levels of vector-specific neutralizing antibodies. The ID$_{50}$ values of the sera for ChAdTS were defined as the dilution required to reduce the number of GFP-expressing cells by 50% compared with wells treated with the virus alone.

**Assessment of T-cell responses in mice** Cellular immune responses in the vaccinated mice were assessed using the IFN-γ pre-coated ELISPOT kit (MabTech, Sweden), according to the manufacturer’s protocol. Splenocytes from mice were stimulated with a peptide pool covering the SARS-CoV-2 S protein (GenScript, USA) at a concentration of 2 μg/mL of each peptide. Phorbol myristate acetate/ionomycin was used as a positive control and RPMI 1640 medium as a negative control. After incubation at 37°C for 28 h, the plates were washed extensively before a biotinylated anti-mouse IFN-γ antibody was added to each well and incubated further for 2 h at room temperature. The substrate AEC was added and the spots in each well were read using the automated ELISPOT reader (AID, USA). The number of spot-forming units (SFUs) per 1,000,000 cells was calculated. For T cell proliferation analysis, approximately 1,000,000 mouse splenocytes were stimulated with the same SARS-CoV-2 S peptide pool as above (2 μg/mL of each peptide) and brefeldin A (GolgiPlug; BD, USA) for 6 h at 37 °C in 5% CO$_2$. Following two washes with PBS, the splenocytes were permeabilized and stained with the fluorescently conjugated antibodies CD4-FITC (GK1.5; BioLegend, USA), CD8-PE/Cyanine7 (53-6.7; BD), CD19-
APC/Cyanine7 (1D3; BD), INFγ-BV421 (XMG1.2; BD), TNFα-APC (MP6-XT22; BioLegend), and IL2-PE (JES6-5H4, BioLegend). Dead cells were stained using the Zombie Yellow Fixable Viability Kit (BioLegend, USA). The data were collected using a Cytek Aurora FACS instrument (Cytek, USA) and analyzed using FlowJo software.

Ethics statement All experiments were carried out in strict compliance with the Guide for the Care and Use of Laboratory Animals of the People’s Republic of China and approved by the Committee on the Ethics of Animal Experiments of Tsinghua University, Chinese Academy of Medical Sciences, and University of Hong Kong. Mouse immunization and characterization were conducted in the animal facility of Tsinghua University. Golden Syrian hamster experiments involving live SARS-CoV-2 were conducted in the ABSL-3 facility at the University of Hong Kong. Rhesus macaque experiments involving live SARS-CoV-2 were performed in the ABSL-4 facility of the Kunming National High-level Biosafety Primate Research Center, Yunnan, China, and approved by the institutional biosafety committee.

Statistical analysis Prism 8.0 software (GraphPad, USA) was used for statistical analysis and data plotting. Unless specified otherwise, the data are presented as means ± SEM. Analysis of unpaired Students’ t-test (two-tailed) was used to determine the statistical significance of differences among different groups (*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001; n.s., not significant).

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Author Contributions

D.Z. and L.Z. conceived, designed, and supervised the entire study. M.L., G.J., H.S., M.X. and W.J. constructed, produced and purified the vaccines and carried out all immunogenicity evaluations in mice. S.L., H. L., and X. P. performed the immunization and protection experiment in rhesus macaques. R. Z., P.W., L.L., S.L., H.C. and Z. C. performed the immunization and protection experiments in golden Syrian hamsters. Q.L., L.F., Q.Z. and X.S. assisted the mouse immunization and evaluation experiments. L.C., B.J., and Z.Z. carried out the live virus neutralization assays. N.W. and Y.G. carried out the droplet digital PCR. Y.W. and H. Q. assisted the intracellular cytokine staining assays. R.W. made the construction of S variants. Z.H., X.T. and L.Y. are employees of Walvax Biotechnology Co., Ltd. M.L., G.J., D.Z. and L.Z. wrote the manuscript. All other authors reviewed and edited the manuscript.
Competing Interests

The authors declare that L.Z., D.Z., M.L. and X.S. are co-inventors on pending patent applications related to the ChAdTS-S, ChAdTS-RBD, and ChAdTS-RBDs vaccine candidates. Z.H., X.T. and L.Y. are employees of Walvax Biotechnology Co., Ltd. The remaining authors declare no competing interests.

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Figure legends

Figure 1. Generation and characterization of recombinant ChAdTS expressing the full-length spike protein or RBD of SARS-CoV-2. (a) Schematic diagram of the recombinant ChAdTS expressing the full-length S protein with its original signal peptide (ChAdTS-S), RBD with the original signal peptide from the S protein (ChAdTS-RBD), or RBD with the secretory signal peptide from mouse IgG (ChAdTS-RBDs). The coding sequence of the S protein or RBD was inserted into the E1 region of the ChAdTS vector under the control of the CMV promoter and terminated using a bovine growth hormone (BGH) polyadenylation signal sequence. (b) Western blot analysis of S and RBD expression in HEK293T cells infected with the recombinant ChAdTS (10^8, 10^9 and 10^10 vp). (c) Flow cytometry analysis of S and RBD expression using the SARS-CoV-2-specific mAbs P2C-1F11 and P2B-2F6. HEK293T cell lysates and intact HEK293T cells infected with the empty ChAdTS vector (10^10 vp) were used as negative controls. VRC01 was used as a negative control antibody. Serum binding activity to SARS-CoV-2 RBD (d, e, f) and neutralizing activity against pseudotyped SARS-CoV-2 (g, h, i) over an 8-week period after single-dose immunization with ChAdTS-S, ChAdTS-RBD or ChAdTS-RBDs. For each recombinant ChAdTS construct, three different doses (10^{10}, 10^9, and 10^8 vp) were administered through either the intranasal (IN) or intramuscular (IM) route, which is indicated by different
symbols and colors. Data points corresponding to animals in the empty ChAdTS vector control groups \(10^{10}\) vp are shown in grey. All data are presented as means ± SEM.

**Figure 2.** ChAdTS-S induces strong and durable immune responses in BALB/c mice. Animals immunized with \(10^{10}\) vp of ChAdTS-S were characterized in greater detail. (a) Neutralizing activity of the immunized serum against live SARS-CoV-2 over an 8-week period after immunization. (b) Binding activity of total saliva IgA and (c) immune serum total IgG with the SARS-CoV-2 S trimer. Binding activity of immune serum IgG subtypes IgG1, IgG2a, and IgG2b with the SARS-CoV-2 S trimer in the (d) IN or (e) IM group. (f) The IgG2a/IgG1 ratios in the IN and IM groups. (g) ELISPOT analysis for IFN-\(\gamma\)-positive splenocytes and FACS analysis of intracellular IFN-\(\gamma\), TNF-\(\alpha\), and IL-2 production in (h) CD8\(^+\) and (i) CD4\(^+\) splenic T cells two weeks after immunization. The durability of the antibody response was analyzed using (j) ELISA with the SARS-CoV-2 S trimer and (k) pseudovirus neutralization up to 36 weeks after single-dose immunization. Data corresponding to animals in the IN group are shown in red, those in the IM group in blue, and those in the control group in grey. All data are presented as the means ± SEM. Analysis of unpaired Students’ \(t\)-test (two-tailed) was used to determine the statistical significance of differences among different groups (*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001; n.s., not significant).

**Figure 3.** ChAdTS-S induces strong immune responses in rhesus macaques. (a) Timeline for vaccination followed by virological and immunological characterization in two groups of animals. In group one, four animals were intranasally immunized with \(10^{11}\) vp of ChAdTS-S (n=2) or \(10^{11}\) vp of the empty ChAdTS vector (n=2). After 4 weeks post-immunization, the animals were challenged with \(10^{6}\) PFU of live SARS-
CoV-2. In group two, the same protocol was followed except that immunization was conducted via the IM route and challenge was carried out 8 weeks after immunization. Blood samples were collected every 2 weeks before viral challenge. Blood samples and nasal swabs were also collected on days 0, 1, 2, 3, 5, and 7 after the challenge. All animals were euthanized on day 7 or 8 after the challenge to quantify the viral load in the lung tissue and conduct histopathological examinations. Binding activity of immune serum total IgG with the SARS-CoV-2 S trimer (b), SARS-CoV-2 RBD (c), or total IgA with the SARS-CoV-2 S trimer (d). Neutralizing activity of immunized serum against pseudovirus (e) and live SARS-CoV-2 (f). Neutralizing activity of immunized serum against the empty ChAdTS vector was assessed using a ChAdTS-GFP neutralization assay (g). Serum levels of cytokines were measured on week 2 (h) and week 4 (i) after immunization. Data corresponding to animals in the IN group are shown in red, those in the IM group in blue, and those in the control group in grey. All data are presented as the means ± SEM.

Figure 4. Vaccination with ChAdTS-S elicits a protective immune response against SARS-CoV-2 infection in rhesus macaques. (a) The total number of viral gRNA and sgRNA copies in lung tissue samples from six lobes was quantified by droplet digital PCR (TargetingOne, China). (b) Viral gRNA and (c) sgRNA in the nasal swabs were monitored and quantified using the same technique up to day 7 after viral challenge. (d) Histopathological comparison of tissues from animals vaccinated with ChAdTS-S and the empty ChAdTS vector animals. Representative tissue sections in standard hematoxylin and eosin staining are shown (10×). The scale bar is 100 μm. Data corresponding to animals in the IN group are shown in red, those in the IM group in blue, and those in the control group in grey. All data are presented as means ± SEM.
Figure 5. Vaccination with ChAdTS-S elicits a protective immune response against SARS-CoV-2 infection in golden Syrian hamsters. The hamsters were immunized with ChAdTS-S (n=4) or saline as control (n=4) via the IM route. (a) The binding ability of the immunized serum for SARS-CoV-2 RBD on week 2 and week 4 after immunization was assessed. (b) The neutralizing ability of the serum against SARS-CoV-2 pseudovirus. Intranasal challenge with $10^5$ plaque-forming units (PFU) of the HKU-13 strain of SARS-CoV-2 was carried out at 6 weeks post immunization. (c) Weight loss in SARS-CoV-2 challenged hamsters. Four days after the challenge, lung tissues were harvested to quantify the viral load using qPCR (d) and by immunostaining lung tissue sections using a specific monoclonal antibody against SARS-CoV-2 nucleocapsid protein (20×) (e,f). Data corresponding to animals in the IM group in blue, and those in the control group in grey. All data are presented as the means ± SEM. Analysis of unpaired Students' t-test (two-tailed) was used (*P < 0.05; **P < 0.01; ***P < 0.001).

Figure 6. Neutralization titers of immune sera from vaccinated BALB/c mice and rhesus macaques against wildtype and new variants of SARS-CoV-2. Sera samples were collected and analyzed against pseudoviruses carrying S protein either from wildtype Wuhan-Hu-1, UK501Y.V1, SA501Y.V2 or BR501Y.V3. (a) Neutralizing activity of immune sera from BALB/c mice 36 weeks post immunization and (b) from rhesus macaques 2 weeks post IN immunization and 6 weeks post IM immunization. All data are presented as the means ± SEM. Analysis of unpaired Students’ t-test (two-tailed) was used.
Fig. 1

a) Schematic diagrams of the constructs used in the study:
- ChAd
- ChAdTS-S
- ChAdTS-RBD
- ChAdTS-RBDs

b) Western Blot analysis showing bands for Spikes, S1, β-actin, and RBDs.

C) Cell Counts (Fluorescence intensity %) for different constructs and markers:
- ChAdTS-S
- ChAdTS-RBD
- ChAdTS-RBDs
- ChAdTS

- Cell Counts: 10^10 vp, 10^9 vp, 10^8 vp
- Fluorescence intensity %:
  - Spike
  - S1
  - β-actin
  - RBD

D, E, F) Graphs showing the immunogenicity of constructs over weeks post immunization:
- ChAdTS-S
- ChAdTS-RBD
- ChAdTS-RBDs
- IN and IM routes

G, H, I) Graphs showing the immunogenicity of constructs over weeks post immunization:
- D50 (Log10)
- D50 (Log10)
- D50 (Log10)
Fig. 3

a) Rhesus macaque (n=4) Immunization IN ChAdTS-S (n=2) ChAdTS (n=2) Challenge Euthanasia

b) SARS-CoV-2 S trimer
   ELISA titer (Log10)
   Weeks post immunization

c) SARS-CoV-2 RBD
   ELISA titer (Log10)
   Weeks post inoculation

d) SARS-CoV-2 S trimer
   IgA-ELISA titer (Log10)
   Weeks post immunization

e) SARS-CoV-2 Pseudovirus
   ID50 (Log10)
   Weeks post immunization

f) SARS-CoV-2 live virus
   PRNT50 (Log10)
   Weeks post infection

g) ChAdTS-GFP
   ID50 ChAdTS (Log10)
   Weeks post immunization

h, i) Cytokines in plasmas
   Weeks post immunization

Weeks post immunization

Legend:
- ChAdTS-S IN
- ChAdTS IN
- ChAdTS-S IM
- ChAdTS IM

Significance:
- ns
- ✱ ✱✱✱
- ✱✱✱✱
Fig. 4

(a) Lung Tissues
(b) Nasal swabs gRNA
(c) Nasal swabs sgRNA

Days post inoculation
Log_{10} copies/mL

Nasal swabs gRNA
Nasal swabs sgRNA

ChAdTS-S IN
ChAdTS IN
ChAdTS-S IM
ChAdTS IM

%CD8^{+} T cells

ChAdTS-S i.m.
ChAdTS-Empty i.m.
ChAdTS-S i.n.
ChAdTS-Empty i.n.

ns
✱✱✱
✱✱✱✱
ns
✱✱
✱✱✱

ns
✱
✱

ChAdTS
ChAdTS-IM
**Fig. 6**

**Mouse serums**

- ChAdTS-S IM
- ChAdTS-S IN

- ID50 (Log10)

- Wuhan-Hu-1
- UK501Y.V1
- SA501Y.V2
- BR501Y.V3

**Rhesus macaque serums**

- ChAdTS-S IM
- ChAdTS-S IN

- ID50 (Log10)

- P=0.14
- P=0.55
- P=0.28
- P=0.24
- P=0.05
- P=0.39
- P=0.42
- P=0.54