Immunogenicity and protective efficacy of a rhesus adenoviral vaccine targeting conserved COVID-19 replication transcription complex

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The COVID-19 pandemic marks the third coronavirus pandemic this century (SARS-CoV-1, MERS, SARS-CoV-2), emphasizing the need to identify and evaluate conserved immunogens for a pan-sarbecovirus vaccine. Here we investigate the potential utility of a T-cell vaccine strategy targeting conserved regions of the sarbecovirus proteome. We identified the most conserved regions of the sarbecovirus proteome as portions of the RNA-dependent RNA polymerase (RdRp) and Helicase proteins, both of which are part of the coronavirus replication transcription complex (RTC). Fitness constraints suggest that as SARS-CoV-2 continues to evolve these regions may better preserve cross-reactive potential of T-cell responses than Spike, Nucleocapsid, or Membrane proteins. We sought to determine if vaccine-elicited T-cell responses to the highly conserved regions of the RTC would reduce viral loads following challenge with SARS-CoV-2 in mice using a rhesus adenovirus serotype 52 (RhAd52) vector. The RhAd52.CoV.Consv vaccine generated robust cellular immunity in mice and led to significant reductions in viral loads in the nasal turbinates following challenge with a mouse-adapted SARS-CoV-2. These data suggest the potential utility of T-cell targeting of conserved regions for a pan-sarbecovirus vaccine.

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INTRODUCTION

In December 2019, SARS-CoV-2 caused an outbreak of severe pneumonia in Wuhan, China marking the third coronavirus epidemic this century.1,2. Throughout the current pandemic multiple SARS-CoV-2 variants have emerged3. The continued development of new variants as well as the recurrent emergence of pathogenic coronaviruses highlights the need for more broadly applicable vaccines4,5.

The goal of our study was to focus on the most highly conserved region of the sarbecovirus proteome with the intention of generating broadly protective T-cell responses. Should an outbreak with a novel sarbecovirus occur, such a vaccine could prove useful either as a primary vaccine or as a supplement to optimal protection provided by current SARS-CoV-2 vaccines. The vaccine immunogen in the present study is a 1094 amino acid long stretch of ORF1ab comprising the last part of nsp13 (Helicase), spanning the most conserved contiguous section of the replication transcription complex (RTC)6-8 of the sarbecovirus proteome. We term this region CoV.Consv (coronavirus conserved).

Recent data have identified the RTC as a potential broadly protective coronavirus immunogen. Seronegative healthcare workers (SN-HCW) (determined by anti-spike-1 and anti-nucleoprotein neutralization assays) were found to have memory T cells targeting a wider variety of SARS-CoV-2 viral proteins than a pre-pandemic cohort, with the highest frequency of T cells targeting the RTC. Among the SN-HCWs, those with the highest abundance of RTC specific T cells also had an increase in IFI27, a marker used to detect early SARS-CoV-2 infection9, but no seroconversion was observed suggesting subclinical SARS-CoV-2 infection9. A different study identified that SARS-CoV-2 infected patients with mild symptoms had a higher abundance of CDB+ T cells specific for peptides from sequences sharing high similarity with other human CoVs, including two highly conserved epitopes within nsp1210. These studies provided further rationale to assess the protective efficacy of the CoV.Consv immunogen.

To test the immunogenicity and protective efficacy of this immunogen, the Wuhan SARS-CoV-2 variant of the CoV.Consv immunogen was expressed by a rhesus adenovirus serotype 52 vector (RhAd52) with the goal of inducing cellular immunity11,12. This vaccine was found to induce antigen specific CD8+ T-cell responses in BALB/c mice. After challenge with a mouse-adapted SARS-CoV-213,14, RhAd52.CoV.Consv led to significantly reduced viral loads in the upper respiratory tract and a trend towards reduced viral loads in the lower respiratory tract.

RESULTS

Development of RhAd52.CoV.Consv vaccine

Our first objective was to identify the most highly conserved contiguous region of the sarbecovirus proteome. We reasoned that highly conserved regions would be evolutionarily constrained, and thus would also be conserved within SARS-CoV-2 as it continues to evolve during the pandemic, while also eliciting responses that would likely cross-react with other sarbecoviruses. To define the most promising conserved regions, we first codon-aligned
sequences spanning all reading frames in the full-length proteome from 82 different sarbecoviruses.

We identified the regions in the proteome that were most conserved across the viruses in this alignment using two strategies, the entropy at each position in the alignment, and the coverage of Potential T-cell Epitopes (PTEs)\textsuperscript{15,16} in the sarbecovirus alignment by the SARS-CoV-2 pandemic Wuhan reference strain (accession number NC_045512) (Fig. 1). PTEs are defined as all possible linear peptides of 9 amino acids in length in the alignment. As expected, both strategies identified the same three conserved regions (Fig. 1). These included a short region in Spike, 111 amino acids long, S 944–1054 (region 3); a modestly conserved section of the ORF1ab polyprotein beginning in nsp6 and ending in nsp10, 555 amino acids long, ORF1ab 3848–4392 (region 2); and the most conserved and longest stretch, a region spanning part of the RdRp protein (nsp12) and part of the Helicase (nsp13), 1094 amino acids in length, ORF1ab 4692–5785 (region 1). We chose the most conserved region of 1094 amino acids for use in this study (region 1, CoV.Consv), which spanned the region of ORF1ab covering parts of nsp12 and nsp13 and located within the RTC (Fig. 1). Two amino acid mutations, D760A and D761A, were introduced into the RdRp protein for safety to abolish polymerase function\textsuperscript{17}.

We hypothesized that this highly conserved region would not be as susceptible as other regions to acquisition of non-synonymous changes within SARS-CoV-2 over time, and this has been observed as variants have emerged\textsuperscript{18,19}. For example, the Omicron BA.1 baseline sequence differs in only 1 of 1094 amino acids (0.09%) (nsp12 P323L) from the Wuhan reference sequence in this conserved region, and so 99.1% of the PTEs are perfectly matched between the Omicron BA.1 and the Wuhan reference strain\textsuperscript{18}. In contrast, the Omicron BA.1 Spike protein differs from the reference Spike in 39 of 1273 amino acids (3.0%), resulting in only 82.8% perfect PTE matches between the two Spike proteins. This reduction was consistent with the observed reduction in T-cell responses to Omicron BA.1 Spike in vaccinated or convalescent sera\textsuperscript{20,21}.

Alternative T-cell vaccine targets are Membrane (M) and Nucleocapsid (N) as both are highly targeted by T cells\textsuperscript{22,23} making them interesting targets for future vaccine design. Unfortunately, both of these proteins are highly variable among sarbecoviruses (Fig. 1). To assess the relative cross-reactive potential of vaccines targeting proteins of interest, we compared the PTE coverage by S, N, M, and CoV.Consv proteins to three different level of coronavirus diversity. First, we tested the immunogens’ capacity to provide theoretical coverage of SARS-CoV-2 pandemic diversity among SARS-CoV-2 variants of interest and variants of concern (Fig. 2). Second, Sarbecovirus diversity was assessed using an alignment of 162 sequences including 7 human SARS-CoV-1 related sequences isolated from human and civet as well as 155 additional diverse Asian, African, and European sequences sampled from bats and pangolins. A Wuhan reference-based CoV.Consv region antigen provided 93% PTE coverage in the CoV.Consv region, so T-cell responses generated to this region would be predicted to have high cross-reactive potential across sarbecoviruses. In contrast, only 33%, 48%, and 45% of the PTEs were matched for S, N, and M proteins respectively. Finally, other coronavirus data sets were assembled including 42 other coronaviruses known to infect people, and viruses related to these but sampled in non-human hosts (the beta coronaviruses, MERS, OC43, and HKU1, alphacoronaviruses, NL63, and 229E\textsuperscript{24} plus additional coronaviruses that have recently been associated with human infections\textsuperscript{25,26}). These viruses are so distant from sarbecoviruses that reliable alignments for phylogenetic
reconstructions are not feasible in the M, S, and N proteins, however the CoV.Consv region can be readily aligned and a phylogenetic tree uniting all three data sets, the SARS-CoV-2, sarbecovirus and these other coronaviruses is shown in Fig. 2. We found that almost no PTEs were shared between SARS-CoV-2 and these divergent coronaviruses except for a small number in the CoV.Consv region.

To evaluate the protective efficacy of CoV.Consv, we generated a rhesus adenovirus serotype 52 vector expressing this immunogen (RhAd52.CoV.Consv). Our goal was to identify a suboptimal dose of RhAd52.S.PP to use in combination with the CoV.Consv vaccine in order to model, via a heterologous boost strategy, prior exposure to a spike targeted SARS-CoV-2 vaccine with suboptimal protection. We first performed a dose titration study with adenoviral vectors expressing SARS-CoV-2 Wuhan spike antigens containing a modified furin cleavage site and diproline substitution to stabilize the Spike antigen in a prefusion conformation (RhAd52.S.PP).

To identify an appropriate suboptimal dose of RhAd52.S.PP, we vaccinated mice with a single shot of RhAd52.S.PP at $10^7$, $10^8$, and $10^9$ viral particles (vp) per dose. We measured Spike-specific binding antibody titers biweekly for 8 weeks post injection. Interestingly, we found that Spike-specific binding antibodies continued to increase after week 4 with the highest antibody titers at week 8 (Fig. 3a), consistent with recently published work using Ad2627. We further investigated this phenomenon by measuring SARS-CoV-2 neutralizing antibodies against the original Wuhan strain. We observed a clear increase in neutralizing antibody titers from week 4 to week 8 after injection of RhAd52.S.PP.
To assess cellular immune responses, BALB/c mice were immunized intramuscularly with RhAd52.S.PP at 10^7 vp RhAd52.S.PP as a suboptimal vaccine dose for future experiments.

**Cellular immunity**

To assess cellular immune responses, BALB/c mice were immunized at week 0 and week 4 with RhAd52 vaccines, and spleen and lungs were collected to evaluate Spike and CoV.Consv specific cellular immunity at week 8. Cellular immunity was evaluated using IFN-γ enzyme-linked immunospot (ELISPOT) assays and intracellular cytokine staining (ICS) assays. Spike-specific cellular immune responses were evaluated by stimulation with S1 and S2 peptide subpools. CoV.Consv responses were studied by stimulation with nsp12 and nsp13 peptide subpools. Using ELISPOT assays, we identified that RhAd52.S.PP vaccination induced cellular responses favoring nsp13 over nsp12 in BALB/c mice (Fig. 4a). ICS assays in the lung showed higher IFN-γ positive CD8+ T cells than in the spleen for RhAd52.S.PP vaccination but not for RhAd52.S.PP (Fig. 4b, c, Supplementary Figs. 1–4). This increase in IFN-γ positive CD8+ T cells in the lungs compared to spleen was also observed after combination CoV.Consv/S.PP vaccination (Group 6 and 7) in response to both CoV.Consv peptides and spike peptides (Fig. 4b, c, Supplementary Figs. 1–4).

**DISCUSSION**

The three coronavirus epidemics over the past 20 years highlight the need for pan-sarbecovirus vaccines. In this study, we identified and evaluated the most highly conserved region in the sarbecovirus proteome as a candidate vaccine immunogen. We identified a region termed CoV.Consv, which spanned the junction between the SARS-CoV-2 nsp12 and nsp13 proteins as highly conserved among human sarbecoviruses, and more conserved than Membrane, Nucleocapsid, and Spike. RhAd52 vectors expressing CoV.Consv elicited cellular immune responses and led to reduced virus in the upper respiratory tract following mouse-adapted SARS-CoV-2 challenge of BALB/c mice. These data are consistent with prior studies from our laboratory that showed that depletion of CD8+ T cells from convalescent rhesus macaques
prior to SARS-CoV-2 rechallenge led to increased viral replication in the upper respiratory tract\textsuperscript{29}. Future studies could include T-cell depletion experiments following RhAd52.CoV.Consv vaccination.

Previous publications identified the RTC as a highly conserved coronavirus region\textsuperscript{8,10}. In one study of seronegative healthcare workers, those with detectable RTC-specific T-cell responses were more likely to demonstrate features of abortive SARS-CoV-2 infection\textsuperscript{8}. In a separate report, CD8\textsuperscript{+} T cells targeting the RTC were associated with milder COVID-19 disease\textsuperscript{10}. Both reports emphasized the need to investigate the RTC as a potential pan-coronavirus immunogen. Our results suggest that these regions have potential as vaccine immunogens for SARS-CoV-2 as well as other sarbecoviruses.

To further investigate T-cell based coronavirus vaccines, use of an animal model with longer disease duration would be desirable. In the mouse-adapted SARS-CoV-2 challenge model, disease naturally begins to resolve after day 4 with virus clearance by day 6 or 7, while human disease extends to about

\begin{figure}
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\includegraphics[width=\textwidth]{fig4.png}
\caption{Antiviral cellular immunity elicited by multiple RhAd52.CoV.Consv prime / boost regimens in BALB/c mice. a–c Antiviral cellular immunity as measured by IFN\textsubscript{\gamma} ELISpot (A) and IFN\textsubscript{\gamma}-secreting CD8\textsuperscript{+} CD44\textsuperscript{+} T cells in isolated lung (b) and spleen (c) tissue from BALB/c mice immunized with the designated prime RhAd52 vaccine, boosted with the corresponding RhAd52 vaccine 4 weeks later, and sampled 4 weeks following the boost (8 weeks following the prime immunization). Individual mice are denoted by single dots with bars displaying the mean ± the SEM. Gating strategy and representative flow plots are found in Supplementary Figs. 1–4.}
\end{figure}
2 weeks\textsuperscript{13,14}. The short timescale of the mouse model is sufficient for observing sterilizing immunity, but reduction in disease severity may be more difficult to observe. Thus, apparent benefits conferred by T-cell responses to the highly conserved regions within the RTC in humans may not be apparent using this animal model system. In the future, alternative models could be explored such as hamsters where disease typically peaks on the order of 7 days post infection\textsuperscript{10}. In addition, it may be useful to combine this T-cell focused vaccine with an antibody-based B-cell vaccine.
In conclusion, this CoV.Consv vaccine-elicited T-cell responses in mice and led to a modest but significant reduction of virus in the upper respiratory tract in a mouse-adapted acute model of SARS-CoV-2 infection. A single CoV.Consv immunogen would likely suffice for broad pan-sarbecovirus coverage of this region, but for broader alpha and betacoronavirus coverage, polyvalent designs would likely be essential even for the CoV.Consv region. Combining multivalent Spike immunogens with CoV.Consv may prove useful for candidate pan-sarbecovirus vaccines.

METHODS

Defining regions of conservation in the sarbecovirus proteome

Our original alignment of sarbecoviruses included 14 SARS-CoV-2 variants, 14 SARS-CoV variants, 48 diverse sarbecovirus isolates from bats, and 6 pangolin sequences, a data set assembled in 2020, that started from the sarbecovirus reference alignment used in Li et al.\(^{31}\). We identified the regions of the proteome that were most conserved across the viruses in this alignment using two strategies (Fig. 1). First, we used Shannon entropy as a measure of relative diversity in each position in the full proteome alignment, using the Entropy tool at the Los Alamos HIV database (www.hiv.lanl.gov)\(^{32,33}\). Next, we used the Positional Epitope Coverage Assessment Tool to identify the regions of the proteome that enabled the best coverage of Potential T-cell Epitopes PTEs\(^{15}\) in the sarbecovirus alignment by the Wuhan reference variant of the SARS-CoV-2 pandemic (GenBank reference strain NC_045512, or GISAID variant WIV04, EPI_ISL_402124, which are equivalent). PTEs are defined as all possible linear peptides of 9 amino acids in length in the alignment. Nine amino acids was selected as the peptide length for PTE coverage assessment as it is the most common length of CD8+ T-cell epitopes; and is also the typical length of the core region of CD4+ T-cell epitopes. As expected, both strategies identified precisely the same three highly conserved regions within the alignment, as shown in Fig. 1. The alignment set included some subsets of highly related viruses, like the human SARS-CoV-2 subset, and pangolin virus subset; the proportion of these groups will of course impact the values of the entropy and PTE coverage scores. However, as a codon-aligned full-length genome alignment was used, the composition of variant strains was identical across all protein coding regions, thus the relative scores in the different protein regions enables a meaningful comparison, and it was these relative values across the proteome that were used for defining conserved regions. PTE scores approaching 1 dominate in the conserved regions we defined, so these regions were extremely highly conserved, including among the diverse variants isolated from bats. In contrast, PTE scores of ≤0.17 reflect 9-mers that match the Wuhan reference sequence only among the 14 SARS-CoV-2 viruses included in the alignment.

An extended alignment of the CoV.Consv region generated in 2022 was used to create the tree in Fig. 2, including a representative sequence of each WHO variant of interest or concern (VOI and VOC) and Omicron variant noted to be important for continued monitoring through July 2022 (https://www.who.int/activities/tracking-SARS-CoV-2-variants). The SARS-CoV-2 alignments are available through the Los Alamos National Laboratory download page at GISAID. An additional 162 sarbecovirus sequences were added based on literature\(^{34,40}\) and GenBank searches, selected to be representative of diversity, and an additional 44 alpha- and betacoronavirus sequences were added based on BLAST searches of coronavirus from human infections that were down selected to have complete sequences spanning CoV.Consv, S, M and N proteins and were representative of sampled diversity among similar sequences. This alignment is available, and the tree shown in Fig. 2 was generated using IQ-tree 1.6.11\(^{41}\) based on the CoV.Consv region using protein sequences, with a best fit model of LG + F + R3 and midpoint rooting, run via the Los Alamos HIV database IQ-tree web interface and visualized using their Rainbow tree tool.

Rhad52 vectors

Rhad52.Cov.Consv vaccines were generated from region 1 of the SARS-CoV-2 genome (Fig. 1). The sequences were codon optimized and synthesized through Thermo Fisher Scientific Geneart. Replication incompetent, Rhad52 vectors with E1/E3 regions were deleted were grown in HEK 293B-S5K.TetT cells\(^{42}\). The E1 region was replaced by the CoV.Consv immunogen. All vectors were sequenced and tested for infectivity before use.

Animals and study design

Female BALB/c mice (The Jackson Laboratory) were randomly allocated to groups. Mice received Rhad52 vectors expressing the SARS-CoV-2 prefusion spike protein with diproline mutation, CoV.Consv immunogen, a combination of the two, or sham controls (N = 5 per group). Animals received a prime and boost immunization of either 10\(^7\) or 10\(^9\) viral particles (VPs) of Rhad52 vectors by the intramuscular route without adjuvant. All Adenoviral vector injections were 10\(^9\) VPs unless otherwise stated. Peripheral blood was collected via the submandibular route to isolate serum for immunologic assays. Spleen and lungs were collected and processed for cellular assays. For viral challenge, mice were administered 1 x 10\(^4\) PFU MA10 SARS-CoV-2 in a volume of 50 μL via the intranasal route\(^{15,14}\). Following challenge, body weights were assessed daily. Animals were euthanized on day 4 post-challenge for viral outgrowth assays. All animal studies were conducted in compliance with all relevant local, state and federal regulations and were approved by the Beth Israel Deaconess Medical Center and University of North Carolina at Chapel Hill Institutional Animal Care and Use Committees.

SARS-CoV-2 Spike ELISA

S binding antibodies were assessed as follows: 96-well plates (Thermo Fisher Scientific) were coated with 1 μg/ml of SARS-CoV-2 spike protein (Sino Biological) in 1X DPBS and incubated at 4 °C overnight. After incubation, plates were washed once with wash buffer (0.05% Tween 20 in 1X DPBS) and blocked with 350 μL of Casein per well for 2–3 h at room temperature after which block solution was discarded and plates were blotted dry. Three-fold serial dilutions of mouse serum in casein block were added to wells and plates were incubated for 1 h at room temperature.
Intracellular cytokine staining assay

Lungs were placed into Miltenyi gentleMACS C-Tubes (130-093-237) containing 4 mL R10 and 1 mL 5x digestion buffer (16.5 mL R10, 100 mg Type IV Collagenase). Lungs were processed using the Miltenyi gentleMACS Dissociator with heating (130-093-235) on “m_lung_01_02” program. Lung cells were then passed through a 70 µm filter. Cells were spun down for 5 min at 1400 rpm. Cells were resuspended in 3 mL 1x ACK Lysing Buffer (gibco A10492-01) for 3 min to lyse red blood cells. ACK was neutralized with 20 mL R10. Cells were spun down for 5 min at 1400 rpm. Lung cells were then resuspended in 10 mL R10 and run through 30 µm filter and were then treated as splenocytes for ICS. Splenocytes were resuspended in 100 µL of R10 media. Each sample was incubated with mock (100 µL of R10; background control), Spike 1, Spike 2, nsp12, nsp13 peptide pools (2 µg/mL), or Leukocyte Activation Cocktail with BD GolgiPlug from BD Biosciences (550583) (positive control) and incubated at 37 °C for 1.5 h. After incubation, 0.2 µL of GolgiStop and 0.2 µL of GolgiPlug in 50 µL of R10 was added to each well and incubated at 37 °C for 5 h and then held at 4 °C overnight. The following day, cells were washed twice with MACS Buffer, stained with Aqua live/dead dye and then stained with predetermined titers of mAbs against CD8a (clone 53-6.7; BUV395), CD3 (clone 17A2; BUV737), CD103 (clone 2E7; BUV605), CD44 (clone IM7; BUV711), CD62L (clone MEL-14; Alexa Fluor 488), CD4 (clone RM4-5; PE/Dazzle 594), and CD69 (clone H1.2F3; PE-Cy7) (all antibodies diluted 1:100) for 1 h. Cells were then washed twice with MACS buffer and incubated with 100 µL of BD Cytofix/Cytoperm Fixation/Permeabilization solution for 15 min. Cells were then washed twice with 1x Perm Wash buffer (BD Perm/WashTM Buffer 10x in CytoFix/CytoPerm Fixation/Permeabilization kit diluted with MilliliQ water and passed through 0.22 μm filter) and stained intracellularly with mAbs against IFN-γ (clone XMG1.2; BV421), IL-2 (clone JES5-5H4; PE), IL-4 (clone 11B11; APC), and TNFα (clone MP6-XT22; Alexa Fluor 700) (all antibodies diluted 1:100) for 1 h. Cells were then washed twice with 1X Perm Wash buffer and washed once with MACS buffer then fixed with 250 µL of freshly prepared 2% formaldehyde. Fixed cells were analyzed by BD LSRRII system.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

DATA AVAILABILITY

Authors can confirm that all relevant data are included in the paper.

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AUTHOR CONTRIBUTIONS

G.D., J.D.V., K.W., B.K. and D.H.B. designed and planned experiments. G.D., J.D.V., T.A., B.S.C, M.S., J.B., J.M., J.Y., and L.H.T. generated the vectors and performed the mouse immunogen test studies. D.R.M., A.S., and R.B. performed the mouse challenge studies. G.D. and D.H.B wrote the paper together with all co-authors. G.D and J.D.V are listed as co-first authors.

COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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