High affinity nanobodies block SARS-CoV-2 spike receptor binding domain interaction with human angiotensin converting enzyme

Thomas J. Esparza1,2, Negin P. Martin3,4, George P. Anderson5, Ellen R. Goldman5 & David L. Brody1,6*

There are currently few approved effective treatments for SARS-CoV-2, the virus responsible for the COVID-19 pandemic. Nanobodies are 12–15 kDa single-domain antibody fragments that can be delivered by inhalation and are amenable to relatively inexpensive large scale production compared to other biologicals. We have isolated nanobodies that bind to the SARS-CoV-2 spike protein receptor binding domain and block spike protein interaction with the angiotensin converting enzyme 2 (ACE2) with 1–5 nM affinity. The lead nanobody candidate, NIH-CoVnb-112, blocks SARS-CoV-2 spike pseudotyped lentivirus infection of HEK293 cells expressing human ACE2 with an EC50 of 0.3 µg/mL. NIH-CoVnb-112 retains structural integrity and potency after nebulization. Furthermore, NIH-CoVnb-112 blocks interaction between ACE2 and several high affinity variant forms of the spike protein. These nanobodies and their derivatives have therapeutic, preventative, and diagnostic potential.

Coronaviruses are positive sense, single-stranded RNA viruses. There are seven types of coronaviruses known to infect humans, including the recent 2019 severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2)1,2. Patients infected with these viruses develop respiratory symptoms of various severity and outcomes. Since the beginning of the century, there have been three major world-wide health crises caused by coronaviruses: the 2003 SARS-CoV-1 outbreak, the 2012 MERS-CoV outbreak, and the 2019 SARS-CoV-2 outbreak3. To date, hundreds of thousands of people have succumbed to the virus during these outbreaks.

The SARS-CoV-2 virus gains entry to human cells via the angiotensin converting enzyme 2 (ACE2) receptor by the interaction of the receptor binding domain (RBD) of the spike protein on the viral surface4–8. This RBD-ACE2 interaction provides a clear therapeutic target for binding and prevention of infection (Fig. 1a).

Many of the antibodies considered for diagnostic or therapeutic applications have been conventional immunoglobulins (IgG). The use of IgGs as therapeutics, while successful in many diseases, is known to have potential pitfalls due to the risk of receptor-mediated immunological reactions9. Treatment or prophylaxis of a pulmonary virus can be delivered via aerosolization and inhalation; thus, the size and biophysical characteristics are paramount considerations.

The camelid family, which includes llamas, produce additional subclasses of IgGs which possess an unpaired heavy-chain variable domain10–12. This heavy-chain variable domain has demonstrated the ability to function as an independent antigen-binding domain with similar affinity as a conventional IgG. These heavy chain variable domains can be expressed as a single domain, known as a VHH or nanobody, with a molecular weight 10% of the full IgG. Nanobodies generally display superior solubility, solution stability, temperature stability, strong penetration into tissues, are easily manipulated with recombinant molecular biology methods, and possess robust environmental resilience to conditions detrimental to conventional IgGs13–15. In addition, nanobodies are weakly immunogenic which reduces the likelihood of adverse effects compared to other single domain antibodies such
as those derived from sharks or synthetic platforms. Successful framework modifications have been implemented to humanize VHH sequences without significant alteration of biophysical properties. Importantly, the recent approval of the first-in-class nanobody treatment for thrombotic thrombocytopenic purpura, Caplacizumab, bolsters the therapeutic potential of VHH derivatives. Therefore, nanobodies that bind to the SARS-CoV-2 RBD and block the ACE2 interaction are an attractive therapeutic for prevention and treatment of viral infection.

Results

Using standard methods to immunize llama, B-cell nanobody cDNA sequences were isolated and a phage display library with over $10^8$ clones was created (Fig. 1b). From this phage library, 13 unique lead candidate nanobodies that bind to the SARS-CoV-2 spike protein RBD were isolated, several of which block the spike protein-ACE2 interaction with high potency.

To isolate the candidate nanobodies, a novel screening strategy was designed and executed to specifically select for nanobodies that not only bind to the SARS-CoV-2 spike RBD, but also interfered with the interaction with the human ACE2 protein. The purity of in vitro binding of commercially available recombinant spike protein RBD and human ACE2 protein (Supplementary Fig. 1a–c) that were used for the screening strategy was validated. In this screening strategy (Fig. 1c), recombinant human ACE2 protein was immobilized and the surface blocked with non-specific protein. Biotinylated-RBD, was incubated with the nanobody phage library and then added to the immuno-tube and allowed to interact with the immobilized ACE2. Biotinylated-RBD with no associated nanobodies, or with nanobody associations which do not block the ACE2 binding domain, bound the immobilized ACE2. Biotinylated-RBD with associated nanobodies that blocked the ACE2 binding domain remained in solution and were recovered using streptavidin-coated magnetic particles that bind to the biotin. Nanobodies which did not bind to RBD were removed during washing of the magnetic beads. This method allowed for specific enrichment of nanobodies which both bind to the RBD and compete for the RBD-ACE2 binding surface.

Using this novel screening strategy, hundreds of phage clones were isolated, which when sequenced, revealed 13 unique full length nanobody DNA sequences, termed NIH-CoVnb-101 through NIH-CoVnb-113 (Fig. 2). These sequences were distinct from the previously published sequences that also bind SARS-CoV-2 spike protein. Additional reports demonstrate compelling binding to SARS-CoV-2 spike protein, yet have not disclosed nanobody sequences allowing for a direct comparison. The complementarity determining region (CDR) 3 domain
responsible for much of the specific binding of nanobodies to their targets, in NIH-CoVnb-112, highlighted in gray, set as the reference sequence for comparison. For comparison, seven previously reported nanobody sequences have clearly distinct sequences: Ty123, VHH724, H11-D425, MR326, Sb#1427, Sb2328, and W25UACh29 and possess shorter CDR3 domains (represented in NIH-CoVnb-112 by amino acids 99–120).

Figure 2. Protein sequences for novel nanobodies that bind to the SARS-CoV-2 spike protein receptor binding domain. Single letter amino acid codes. Clustal Omega algorithm used for alignment. Blue highlights indicate sequence diversity with NIH-CoVnb-112, highlighted in gray, set as the reference sequence for comparison. For comparison, seven previously reported nanobody sequences have clearly distinct sequences: Ty123, VHH724, H11-D425, MR326, Sb#1427, Sb2328, and W25UACh29 and possess shorter CDR3 domains (represented in NIH-CoVnb-112 by amino acids 99–120).

Figure 3. Affinity binding curves of isolated anti-SARS-CoV-2 RBD nanobodies. Using Biolayer Interferometry on a BioForte Octet Red96 system, association and dissociation rates were determined by immobilizing biotinylated-RBD onto streptavidin coated optical sensors (a–e). The RBD-bound sensors were incubated with specific concentrations of purified candidate nanobodies for a set time interval to allow association. The sensors were then moved to nanobody-free solution and allowed to dissociate over a time interval. Curve fitting using a 1:1 interaction model allows for the affinity constant (Kᵦ) to be measured for each nanobody as detailed in (f).

Representatives from each of these unique nanobody sequences were produced in bacteria, purified, and tested for binding to SARS-CoV-2 spike RBD (Fig. 3). All of the nanobodies bound to recombinant SARS-CoV-2 spike RBD with high affinity as measured by Octet biolayer interferometry. The strongest binding nanobody was NIH-CoVnb-112 (Fig. 3e) with an affinity of 4.9 nM. NIH-CoVnb-112 had both a fast on rate (1.3e5/M/sec.) and a slow off rate (6.54e-4/sec.), with kinetics compatible with 1:1 binding. In a complementary measurement, ProteOn XPR36 surface plasmon resonance provided good agreement (Supplementary Fig. 2) with an affinity of 2.1 nM. These results indicate that the novel nanobodies bind to the SARS-CoV-2 spike RBD with very high affinity. The long CDR3 in NIH-CoVnb-112 and the other new nanobodies were generally longer than those currently reported. Twelve of the 13 nanobodies are relatively similar to each other, whereas one nanobody (NIH-CoVnb-110) was distinct from the other 12. These findings indicated that novel nanobody DNA sequences were isolated.

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In an important validation of the in vitro efficacy of the candidate nanobodies, the nanobodies were able to interfere with SARS-CoV-2 spike RBD binding to human ACE2 protein (Fig. 4a). Specifically, a competitive inhibition assay was designed and implemented, in which recombinant RBD was coated onto Enzyme Linked Immuno-Sorbent Assay (ELISA) plates and soluble ACE2 binding was assessed. Without any interference, soluble ACE2 binding was indicated by high colorimetric absorbance. At increasing concentrations, each of the new nanobodies, showed progressively less ACE2 binding. For the most potent anti-SARS-CoV-2 RBD nanobody, NIH-CoVnb-112, the concentration at which 50% of the ACE2 binding was blocked (half maximal effective concentration; termed EC\textsubscript{50}) was found to be 0.02 µg/mL (1.11 nM). The rank order of ACE2 competition EC\textsubscript{50} matched the rank order of RBD binding affinities for the novel nanobodies assessed. As an additional confirmation, the experiment was repeated using a commercially available SARS-CoV-2 spike RBD-human ACE2 protein competitive inhibition assay (GenScript). The results were very similar to those obtained using the initial assay (Fig. 4b). These results indicate that the novel nanobodies block SARS-CoV-2 spike protein RBD binding to ACE2, an essential human receptor responsible for viral infection.

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There have been many variants of the spike protein RBD described recently that increase the binding affinity to the human ACE2 receptor\textsuperscript{31}. Several of these variants including N354D D364Y, V367F, and W436R have been reported to have up to 100 fold higher affinity for ACE2 than the prototype RBD in vitro. NIH-CoVnb-112 blocked interaction between human ACE2 and three of these variants with similar EC\textsubscript{50} compared to its blocking effects on the prototype sequence spike protein RBD (Fig. 5b). Binding affinity of NIH-CoVnb-112 to the variants was also similar to that of the prototype sequence (Fig. 5a). Thus, NIH-CoVnb-112 may provide robust blocking of SARS-CoV-2 spike protein RBD binding to ACE2.
We then assessed whether NIH-CoVnb-112 binds to the same spike protein RBD epitope as the previously reported nanobody VHH72\cite{22}. We reasoned that if both nanobodies bind to the same epitope, their signals would occlude each other when applied at saturating concentrations on bilayer interferometry. In contrast, if they bind to different epitopes, their signals would be additive on bilayer interferometry. We found that when NIH-CoVnb-112 was applied at 500 nM (\(> 100 \times \) greater than the \(K_c\) for long enough to reach steady state, and then VHH72 was applied, the signals were clearly additive (Supplementary Fig. 5a). This result clearly indicates that NIH-CoVnb-112 binds to a SARS-CoV-2 spike protein RBD epitope that is distinct from that of VHH72. This result is consistent with the reported findings that VHH72 recognizes a non-ACE2 binding motif on the spike protein\cite{24}, whereas NIH-CoVnb-112 has RBD-ACE2 interaction disruption potency that is quantitatively similar to its affinity - a result that is most consistent with NIH-CoVnb-112 binding directly to the ACE2 interaction domain.

We then performed additional tests to determine if NIH-CoVnb-112 possesses the ability to also bind to the SARS-CoV-1 spike protein RBD domain. We prepared a biotinylated version of NIH-CoVnb-112 which could be used in parallel to test binding to several targets. Biotinylated NIH-CoVnb-112 was immobilized onto streptavidin Octet sensors and then allowed to associate with either SARS-CoV-1 RBD or SARS-CoV-2 RBD at 500–125 nM concentrations. As previously demonstrated, NIH-CoVnb-112 binds robustly to SARS-CoV-2 RBD, yet we found that there was essentially no binding to SARS-CoV-1 RBD. In an orthogonal confirmation, we performed an ELISA in which each RBD was coated onto a standard ELISA plate and allowed to incubate with NIH-CoVnb-112. The ELISA results were similar to the Octet results: (Supplementary Fig. 5c) NIH-CoVnb-112 binds strongly to SARS-CoV-2 RBD and does not associate with SARS-CoV-1 RBD. These results clearly indicate that NIH-CoVnb-112 does not cross react and bind to SARS-CoV-1 spike protein RBD.

The presence of bacterial endotoxins introduces an unwanted liability during the production of NIH-CoVnb-112 due to expression in E. coli. To remove this risk, we cloned the coding sequence for the nanobody into an expression vector for the methlylotrophic yeast Pichia pastoris. The expression vector, pICZalpha, contains an alpha-factor secretion signal sequence which directs the expressed protein into the media supernatant. Following selection of a positive recombinant clone we found an expression yield of 40 mg per liter following 96 h of methanol batch-fed culture.

We envision therapeutic utility for NIH-CoVnb-112 as a nebulized deliverable treatment to the respiratory tract of patients. Owing to the small size and general stability of the nanobody, we sought to test if we could successfully nebulize, recover, and confirm the stability of NIH-CoVnb-112 relative to non-nebulized nanobody. Using the commercially available Aerogen Solo vibrating mesh nebulizer, we nebulized Pichia expressed NIH-CoVnb-112 into an in line custom bead condenser (Fig. 6a) to collect the post-nebulized nanobody. We recovered greater than 90% of the total input protein mass. Following nebulization recovery, we incubated the pre-nebulization and post-nebulization samples at 37 °C for 24 hr to provide physiologic temperature exposure. When assessed by SDS-PAGE gel (Fig. 6b) we found that the pre-nebulization and post-nebulization NIH-CoVnb-112 appear identical with no evidence of degradation or aggregation products. Similarly, analysis by size-exclusion chromatography (Fig. 6c) on a Superdex 75 column yielded similar elution profiles with no evidence of degradation or aggregation relative to the pre-nebulization sample. We conclude that NIH-CoVnb-112 is resilient to degradation or aggregation during nebulization.

To further assess the stability of NIH-CoVnb-112, we performed incubation in pooled normal human plasma and recombinant human albumin followed by affinity measurement to assess preservation of SARS-CoV-2 RBD binding potential. NIH-CoVnb-112, at 5 µM, was incubated for 24 or 48hrs in the presence of human plasma, which contains proteases, or recombinant human albumin alone at 37 °C with gentle mixing. In addition, for each condition a time zero control was prepared to account for potential matrix effects. The samples were diluted to 500 nM based on the NIH-CoVnb-112 starting concentration and assessed by biolayer interferometry for binding to SARS-CoV-2 RBD. NIH-CoVnb-112 binding (Supplementary Fig. 4a) SARS-CoV-2 RBD following treatment in pooled human plasma at time zero, 24 hr, and 48 hr had negligible impact on binding. Similarly, NIH-CoVnb-112 binding (Supplementary Fig. 4b) SARS-CoV-2 RBD following treatment in recombinant human albumin at all time points had no apparent effect on binding. While further in vivo characterization is necessary, these data support the interpretation that NIH-CoVnb-112 is satisfactorily stable in the presence of plasma.

Finally, while in a plate-based protein inhibition assay NIH-CoVnb-112 robustly inhibits the SARS-CoV-2 RBD and ACE2 interaction, we sought to demonstrate if a similar inhibition is achieved in a viral transduction assay. We used pseudotyped SARS-CoV-2 spike protein bearing lentivirus with an RFP reporter system to test if NIH-CoVnb-112 could inhibit the transduction of human embryonic kidney cells (HEK293T/17) overexpressing human ACE2 (HEK293-ACE2). Transduction of HEK293-ACE2 cells results in RFP expression (Fig. 6d) in the absence of an inhibitory agent. The pseudotyped virus were produced using a human codon optimized SARS-CoV-2 spike protein sequence with the endoplasmic reticulum retention signal removed. Pseudotyped virus were purified over a sucrose cushion and viral titre determined using digital droplet PCR. A dilution of Pichia expressed, pre-nebulization and post-nebulization NIH-CoVnb-112 samples were incubated with a Multiplicity of Infection (MOI) of 0.5 followed by incubation on HEK293-ACE2 cells for 48hrs. Following transduction, brightfield and epifluorescence images were acquired for both pre-nebulization (Supplementary Fig. 6a) and post-nebulization (Supplementary Fig. 6b) conditions. Incubation at or above 0.3 µg/mL NIH-CoVnb-112 results in robust inhibition for both conditions. Following trypsinization and fixation, the individual conditions were analyzed by flow cytometry to determine the percentage of RFP expressing cells. The population of single events were recorded for both pre-nebulization (Supplementary Fig. 7a) and post-nebulization (Supplementary Fig. 7b) conditions. The number of RFP positive events for each condition were normalized relative to the virus-alone control to yield a percent inhibition of the SARS-CoV-2 pseudotyped lentivirus (Fig. 6e). The pre-nebulization NIH-CoVnb-112 has an EC\(_{50}\) of 0.323 µg/mL (23.1 nM) while the post-nebulization NIH-CoVnb-112 has an EC\(_{50}\)
of 0.116 µg/mL (8.3 nM). The difference in EC50 values between the two conditions is most likely an effect of assay variance and we do not speculate that nebulization produced an increase in potency. Thus, we find that NIH-CoVnb-112 potently inhibits viral transduction in an infection relevant pseudotyped SARS-CoV-2 virus model.

**Discussion**

Here, we report several nanobodies that bind to the SARS-CoV-2 spike protein RBD and block spike protein interaction with the ACE2 receptor. The affinity of NIH-CoVnb-112 in monomeric form is substantially better than the monomeric form of previously reported candidate nanobody therapeutics for SARS-CoV-2: 39 nM for monomeric VHH7224, ~ 50 nM for monomeric Ty135, 30.7 nM for monomeric Sb#14315, 10 nM for monomeric Sb2330, and incompletely reported but likely lower than 1B and 3F27. The affinity and blocking potency of NIH-CoVnb-112 will likely be even higher when formulated into dimeric, trimeric, or other protein engineered constructs27,30,32.

Importantly, these nanobody therapeutics may be delivered via inhalation34. Inhalation has major advantages over other routes of administration and could be one of the most important potential uses for a nanobody therapeutic for SARS-CoV-2. The most directly analogous study was performed by Larios Mora et al., who showed that nebulized treatment of newborn lambs with ALX-0171 reduced clinical, virological, and pathological manifestations of respiratory syncytial virus35. ALX-0171 is a trimeric nanobody therapy produced in *Pichia pastoris* that binds to the respiratory syncytial virus fusion protein. Nebulization was performed using a commercially available nebulizer system.

Figure 6. NIH-CoVnb-112 stability and potent inhibition of SARS-CoV-2 pseudovirus following nebulization. (a) An Aerogen Solo High-Performance Vibrating Mesh nebulizer was placed in line with a custom glass bead condenser to allow for collection of the nanobody following nebulization. (Image element courtesy of Aerogen.) (b) A pre-nebulization and post-nebulization sample of purified NIH-CoVnb-112 was analyzed by SDS-PAGE gel. The dominant band for each sample remains at approximately 14 kDa indicating no detectable degradation or aggregation of NIH-CoVnb-112 following nebulization. (c) Size exclusion chromatography demonstrated a prominent peak eluting at 13.5 mL elution volume in both pre and post-nebulization samples. (d) A fluorescence reporter assay utilizing SARS-CoV-2 spike protein pseudotyped lentivirus was used to demonstrate potent inhibition of the RBD-ACE2 interaction. HEK293 cells overexpressing human ACE2 were cultured for 24 h with pseudotyped virus which was pretreated with NIH-CoVnb-112 at different concentrations. Inhibition of the spike RBD occurs when the virus is not able to transduce the HEK293-ACE2 cells and subsequently produce RFP reporter protein. (e) Following 48 hr incubation, HEK293-ACE2 cells were analyzed by flow cytometry to quantify the fluorescence level. NIH-CoVnb-112 potently inhibited viral transduction both pre and post-nebulization with an EC50 of 0.323 µg/mL (23.1 nM) and 0.116 µg/mL (8.3 nM) respectively. (Figure elements generated using BioRender.com).
available human-use Aeroneb Solo system (https://www.aerogen.com/aerogen-solo-3/), interfaced with a nose mask. The Aeroneb Solo nebulizer produced ~3.3 micron particles, appropriate for deep lung delivery. The authors demonstrated that once daily nebulization resulted in concentrations of ALX-0171 in lung epithelial lining fluid that were ~10× higher than the in vitro EC$_{50}$ for respiratory syncytial virus. All treated lambs had undetectable infectious virus in lung epithelial lining fluid, which supports that the proposed route of administration could reduce infectivity. After nebulization, blood concentrations were ~1000-fold lower than lung epithelial lining fluid concentrations, which is important because low blood concentrations likely reduce systemic toxicity and risk of host antibody formation. Of note, the parent monomeric nanobody to the respiratory syncytial virus fusion protein had an affinity of 17 nM, whereas the nanobody trimer ALX-0171 had an affinity of 0.113 nM. The trimeric ALX-0171 was produced by fermentation in _Pischia pastoris_ with a yield of ~7.5 g per liter. There is great potential for an inhaled therapeutic for SARS-CoV-2 derived from the NIH-CoVnb-112 nanobody, and other similar nanobodies. There are many other potential therapeutic uses for the NIH-CoVnb-112 nanobody. These include intravenous, intramuscular, or subcutaneous treatment formulations for early stage disease, as well as prevention in high risk individuals.

It appears that late-stage disease is mediated more by immunological responses and less by virological pathogenesis, making it important to initiate virologically-targeted therapy early. Importantly, it is likely that nanobody treatments would be substantially less expensive to produce than conventional monoclonal antibodies such as those currently in clinical development pipelines (e.g. REGN-CoV-2 “antibody cocktail” https://www.regeneron.com/covid19 and LY-CoV555: https://investor.lilly.com/news-releases/news-release-details/lilly-begins-worlds-first-study-potential-covid-19-antibody). Thus, it would be reasonable to initiate treatment with an inexpensive and widely available therapeutic early, even before it is clear whether or not an infection will become severe. Furthermore, even as vaccines are successfully developed, preventative treatments for acute illness would be beneficial, as vaccines may not be 100% effective, especially in those who do not mount adequate immune responses. By analogy, oseltamivir (Tamiflu) is an essential medicine even in the context of successful influenza vaccines. Thus, there is a substantial need for inexpensive and safe preventative and acute candidate treatments such as the NIH-CoVnb-112 nanobody.

In addition to therapeutic applications, an inexpensive binding reagent such as NIH-CoVnb-112 nanobody could be used diagnostically. Antibody-based tests could be used to assess for SARS-CoV-2 spike protein in body fluids and the environment. The low cost and temperature stability of nanobodies would be advantageous in some settings, though it is unlikely that antibody-based tests will replace current PCR-based testing for routine clinical use.

Substantial effort will be required to fully characterize and develop NIH-CoVnb-112. We have demonstrated successful neutralization assays involving pseudotyped lentivirus and authentic SARS-CoV-2 isolate neutralization assays are currently underway. Future activities include performing detailed structural and biophysical characterization involving Cryo-EM, X-ray crystallography, in vivo pharmacokinetics and stability assessment, immunogenicity determination, framework humanization, assessing multimerized formats, and further storage and delivery stability assessments. It is encouraging that NIH-CoVnb-112 binds to and blocks three recently reported RBD variants, but it will be of great interest to determine whether the candidate nanobodies provide broad spectrum blocking against other variant forms of the SARS-CoV-2 spike proteins. Additional assays will be required to determine whether NIH-CoVnb-112 protects against mutation-based viral escape alone or as part of a therapeutic cocktail.

Animal testing will be critically important. The optimized nanobody construct will be tested for in vivo safety and efficacy in an established small animal model of SARS-CoV-2 infection such as human ACE2 transgenic mouse or Syrian hamster. Subsequently, testing for in vivo safety and efficacy in an established large animal model of SARS-CoV-2 infection such as rhesus macaque or baboon will be conducted to further support an investigational new drug application. Although in vitro stability and efficacy appears to be excellent, there is no guarantee that NIH-CoVnb-112 will have appropriate in vivo stability and pharmacokinetics. If NIH-CoVnb-112 proves sub-optimal in vivo, we plan to further characterize other candidate nanobodies from the initial screen, as well as performing additional screening and protein engineering. While other nanobodies have been produced in large scale current Good Manufacturing Practice facilities by fermentation, there is no guarantee that NIH-CoVnb-112 constructs can be produced in sufficient quantities at reasonable cost. Exploration of the manufacturing parameters will be critical for product development.

There are several limitations of the current results. First, the characterization of NIH-CoVnb-112 and the other nanobodies is not yet complete. Under the current circumstances, we have opted to move forward with dissemination of our findings earlier than might otherwise be typical. For example, we have not fully characterized all of the candidate nanobodies, nor nanobodies that do not block the interaction of the RBD with ACE2. Likewise, we have not exhaustively tested epitope binding overlap between NIH-CoVnb-112 and all of the other SARS-CoV-2 spike nanobodies developed to date. Second, a potential new therapeutic such as this nanobody would, even under the best circumstances, come to market relatively slowly compared to a repurposed approved drug. Third, it is possible that NIH-CoVnb-112 and other nanobodies may provoke host immune responses. NIH-CoVnb-112 is a llama protein fragment that is similar to human proteins yet cannot be fully humanized and maintain desirable biophysical characteristics. Fourth, NIH-CoVnb-112 and other nanobodies may have unexpected toxicity. There is only one United States Food and Drug Administration-approved nanobody therapeutic, Caplacizumab, and there has not been exposure to a sufficient number of human patients to fully characterize the safety of this class of therapeutics.

In conclusion, while a substantial amount of additional characterization will be required, the novel nanobodies that bind to the spike protein RBD have potential widespread utility in the battle against the SARS-CoV-2 pandemic. We are especially optimistic about the possibility that low-cost, stable, and safe nanobody-based therapeutics will be developed for inhaled use in the home and outside of formal healthcare environments.
Generation of immune single-domain antibody phage display library. The general synthesis methods utilized are adapted from work by Pardon et al. Peripheral blood mononuclear cells (PBMCs) were isolated from llama whole blood using Uni-Sep Maxi density gradient tubes (#U-10, Novamed) according to the manufacturer’s directions. The platelet-rich plasma was further processed and stored at −80 °C for use in measuring antibody titer. The PBMCs were rinsed with 1× phosphate buffered saline (PBS), aliquoted, and frozen at −80 °C prior to use. For total RNA isolation a minimum of 1×107 PBMCs were processed using a RNeasy Mini Kit (Qiagen). First-strand synthesis of mRNA from the resulting total RNA was performed using the SuperScript IV First-Strand Synthesis kit (#18091050, Invitrogen) with random hexamer primed conditions and 2 µg total RNA. Synthesized first-strand complimentary DNA was used to amplify the heavy-chain variable domains using Q5 high-fidelity DNA polymerase (New England Biolabs) and the described primers (CALL001: 5′-GTCTGGCTGCTCTTCTAAAGG-3′ and CALL002: 5′-GTTACGGTGTTGAACCTGTTCC-3′). The resulting amplified variable domains were sequenced on a 1.2% (w/v) low melting point agarose gel and the approximately 700 base pair band corresponding to the heavy chain only immunoglobulin was extracted and purified using QIAquick Gel Extraction Kit (Qiagen). A secondary amplification of the VHH domain was performed using modified primers (VHH-Esp-For: 5′-CCGGCCATGGCTATGTTGACAGCAGGAGTCT GRRGGAGG-3′ and VHH-Esp-Rev: 5′-GTGCGCCGGCTGAGAGACGGTAGTACCTGTT-3′) based on those used by Pardon et al. to introduce cloning compatible sequence for the phagemid pHEN2. The pHEN2 phagemid allows for in-frame cloning of the VH and VL variable domains using Q5 high-fidelity DNA polymerase (New England Biolabs) and the described primers (CALL001: 5′-GGG GGAGG-3′ and CALL002: 5′-GGT ACG TGC TGT TGA ACT GTTCC-3′). The amplified VH domains were purified using QIAquick Gel Extraction Kit (Qiagen) and finally cloned into NcoI/NotI digested pHEN2 phagemid at a 3:1 (insert:phagemid) ratio overnight at 16 °C followed by purification and then electroproporation into phage-display competent TG-1 cells (#60502-1, Lucigen). The library was plated onto 2xYT agar plates containing 100 µg/mL carbenicillin and 2% glucose at 37 °C overnight. The resulting library contained > 108 independent clones. The library was pooled and archived as glycerol stocks. Standard phage amplification of the representative library was performed using M13KO7 helper phage (#18311019, Invitrogen) followed by precipitation by addition of one-fifth volume 20% polyethylene glycol 6000 /2.5 M sodium chloride on ice to purify the phage particles for downstream immunopanning. All post-reaction purifications utilized the MinElute PCR Purification Kit (Qiagen).

Phage display clone screening. Individual colonies were selected from the phage display enrichment plates and cultured in 2xYT-carbenicillin containing microbial media in a 96-deep well plate at 37 °C with 300 rpm shaking for 4–6 h. Expression of nanobody was induced by addition of isopropyl-beta-D-thiogalactoside (GoldBio) to a final concentration of 1 mM and continued incubation overnight. The cells were pelleted by centrifugation at 1000g for 20 min. The media supernatant was removed, and the block placed at −80 °C for 1 h. The block was then allowed to equilibrate to room temperature for 15 min, 500 µL 1xPBS was added, and the block agitated at 1500 rpm for 1 h to allow for release of nanobody from the periplasmic space. The block was then centrifuged for 20 min at 2000×g. Nunc Maxisorp plates were coated with SARS-CoV-2 RBD (SPD-C52H3, ACRO Biosystems) as described above and blocked with 2% bovine serum albumin (BSA). Nanobody containing supernatant was transferred to the blocked plate and incubated for 1 h at room temperature. The assay plate was washed and 100 µL peroxidase conjugated goat anti-alpaca VH domain specific antibody (#128-035-232, Jackson ImmunoResearch) transferred into each well and incubated for 1 h at room temperature. After a final
wash the plate was developed by addition of 100 µL tetramethylbenzidine (TMB) (#T5669, Sigma-Aldrich). Assay development was stopped by addition of 50µL 1 M sulfuric acid and the assay absorbance measured at 450 nm on a Biotek Synergy 2 plate reader. The assay plate was washed with 1xPBS 5 times between each step of the assay. Clones were considered positive if the optical density 450 nm was greater that two standard deviations above the background.

Bio-layer interferometry selection of receptor binding domain binding VHH clones. ELISA-positive clones were further assessed by bio-layer interferometry binding to RBD. Assay buffer was defined as 0.1% BSA (w/v) in 1xPBS. All assay conditions were prepared in a Greiner 96-well plate (#655209) in a volume of 300 µL. Biotinylated SARS-CoV-2 S protein RBD (SPD-C82E9, ACRO Biosystems) with a C-terminal AviTag was used to ensure uniform directionality of the protein. Biotinylated-RBD was diluted into assay buffer at 1 µg/mL and immobilized onto streptavidin coated biosensors (#18-5019, BioForté) to a minimum response value of 1 nm on the Octet Red96 System (ForteBio). A baseline response was established in assay buffer prior to each association. Candidate clone supernatant prepared previously was diluted 1:1 with assay buffer and allowed to associate for 60 s and dissociate for 60 s, which provided a sufficient interval to detect positive binding. Biosensors were regenerated in 250 mM imidazole in assay buffer for 45 s which removed residual bound candidate nanobody and returned to the baseline well. This method allowed for detection of clones that bind RBD in both ELISA and a qualitative estimate of binding based on response curve dynamics. Clones positive on both the RBD ELISA and bio-layer interferometry measurement were selected for sequencing and further characterization.

ProteOn XPR36 surface plasmon resonance affinity measurement of NIH-CoVnb-112 RBD binding. Surface plasmon resonance affinity and kinetic measurements were performed using the ProteOn XPR36 (Bio-Rad). Lanes of a general layer compact (GLC) chip were individually coated with 2 µg/mL SARS-CoV-2 Receptor Binding Domain (RBD) (#SPD-C52H3, ACRO Biosystems) in 10 mM sodium phosphate pH 6.0 and attached to the chip following the standard 1-ethyl-3-(3-dimethylaminopropl)carbodiimide hydrochloride (EDC)/N-hydroxysulfosuccinimide (sulfo-NHS) coupling chemistry available from the manufacturer resulting – 1000RU of protein deposited. Binding kinetics of NIH-CoVnb-112 were tested at 25 °C by flowing six concentrations varying from 300 to 0 nM at 100 µL/min for 90 s or more then dissociation was monitored for at least 600 s. Following each run, the chip was regenerated by flowing 0.85% phosphoric acid (~ pH 3.0) across the surface. Data analysis was performed with ProteOn Manager 2.1 software, corrected by subtraction of an uncoated column as well as interspot correction. The standard error of the fits was less than 10%. Binding constants were determined using the Langmuir model built into the analysis software.

Sanger dideoxy sequencing. Clones enriched by phage display and positively selected by ELISA assay and bio-layer interferometry measurement were sequenced using a universal Lac-promoter primer (Lac-fwd: 5’-CGTATGTTGTGGGAATTGTCG-3’) with standard Sanger dideoxy sequencing at Genewiz. The resulting sequences with Quality Scores above 40 and with Contiguous Read Lengths of greater than 500 were considered reliable. Sequences were trimmed to include only the VHH coding region and the protein coding sequences aligned using the Clustal Omega algorithm (60) included in SnapGene software (GSL Biotech LLC).

Nanobody expression and purification. Lead candidates were expressed by transferring the phagemid from TG-1 cells into the BL21 (DE3) competent E. coli strain (C25271, New England BioLabs). Cultures were grown in 250 mL 4xYT-carbenicillin in a 1-L baffled flask at 37 °C and 300 rpm until the OD600 reached between 0.7–0.9. Protein expression was induced by addition of isopropyl-beta-D-thiogalactoside to a final concentration of 1 mM and the culture conditions reduced to 30 °C and 180 rpm for overnight growth. The expression cultures were pelleted at 8000xG for 10 min at 4 °C; the resulting pellets maintained on ice. Osmotic shock was achieved by resuspension of the cell pellet in TES buffer (0.2 M Tris, 65 µM EDTA, 0.5 M sucrose, pH 8.0) at 1/5th the original culture volume (e.g. 20 mL TES for 100 mL culture volume) and incubated on ice for 1 h with occasional agitation. The suspension was pelleted again as described above and then resuspended in an equal volume of ice-cold 10 mM magnesium chloride and incubated on ice for 30 min with occasional agitation. A final centrifugation was performed at 17,000xg for 10 min at 4 °C to pellet all cellular debris. The resulting supernatant was filtered through a 0.22 µm syringe filter to remove any residual particulates.

Purification of the nanobody was accomplished by injection of the clarified osmotic shock supernatant onto a 1 mL HisTrap FF column (H7531901, Cytiva Lifesciences) coupled to an AKTA Pure FPLC system. Unbound protein was washed with 10 column volumes 1xPBS followed by 10 column volumes 20 mM imidazole in 1xPBS. The enriched nanobody was eluted with 250 mM imidazole in 1xPBS. An automated collection of the unbound flow-through and fractionation of the wash and elution steps was achieved with the fraction collector F9-C. The nanobody typically eluted into two 1.5 mL fractions. The column was then equilibrated with 1xPBS and fractions were collected. The protein concentration was estimated using absorbance at 280 nm and quantified using a standard Bicinchoninic acid protein assay (#23225, ThermoScientific) with BSA as the concentration standard.

Bio-layer interferometry affinity measurement of RBD binding VHH clones. Bio-layer interferometry was used to measure the affinity binding constants of purified VHH clones. Assay buffer is defined as 0.1% BSA (w/v) in 1xPBS. All assay conditions are prepared in a Greiner 96-well plate (#655209) in a volume of
300 µL. Biotinylated SARS-CoV-2 S protein RBD (SPD-C82E9, ACRO Biosystems) with a C-terminal AviTag was used to ensure uniform directionality of the protein. Biotinylated-RBD was diluted into assay buffer at 1 µg/µL and immobilized onto streptavidin coated biosensors (#18-5019, BioForte) to a minimum response value of 1 nm on the Octet Red96 System (ForteBio). A baseline response was established in assay buffer prior to each association. The purified VHH clones were diluted into assay buffer at the specified concentrations (typically 1000–0 nM). The VHH clones were allowed to associate for 180–240 s followed by dissociation for 300–600 s in the same baseline wells. The assay included one biosensor with only assay buffer which was used as the background normalization control. Using the ForteBio Data Analysis suite, the data was normalized to the association curves following background normalization and Savitzky-Golay filtering. Curve fitting was applied using global fitting of the sensor data and a steady state analysis calculated to determine the association and dissociation constants.

Receptor binding domain competition assay. SARS-CoV-2 RBD (#SPD-C52H3, ACRO Biosystems) was coated at 0.5 µg/mL in sodium carbonate buffer (pH 9.6), 100 µL per well, onto Nunc Maxisorp plates overnight at 4 °C. Coating solution was removed, and plate blocked with 300 µL 2% BSA (#5217, Tocris) in 1xPBS for 1 h at room temperature. Purified nanobodies were diluted at specified concentrations in 0.2% BSA in 1xPBS and transferred to the blocked plate in triplicate. The nanobodies were incubated for 45 min to allow for association with RBD. Biotinylated ACE2 (AC2-H82E6, ACRO Biosystems) was prepared at 0.2 µg/µL in 0.2% BSA solution. 10 µL of the ACE2-biotin solution was transferred into each well of the assay and allowed to incubate for 15 min with at 600 rpm. The assay plate was washed and 100 µL of poly-streptavidin (#85R-200, Fitzgerald Industries International) was diluted 1:2000 in 2% BSA solution. The assay plate was transferred to the blocked plate in triplicate. The assay plate was washed and 100 µL of poly-streptavidin (#85R-200, Fitzgerald Industries International) diluted 1:2000 in 2% BSA solution was transferred to each well and incubated with 600 rpm for 30 min. After a final wash, the plate was developed by addition of 100 µL tetramethylbenzidine (#T5656, Sigma-Aldrich). Assay development was stopped by addition of 50 µL 1 M sulfuric acid and the assay absorbance measured at 450 nm on a Biotek Synergy 2 plate reader. The assay plate was washed with 1xPBS 5 times between each step of the assay.

Receptor binding domain variant competition assay. SARS-CoV-2 RBD variants have been noted which increase the affinity for human ACE2 binding. Competition between the RBD mutants was performed as described above with the novel mutation RBD proteins replacing the canonical RBD sequence. SARS-CoV-2 RBD mutants W436R, N354D/D364Y, and V367F (#SPD-S52H7, SPD-S52H3, and SPD-S52H4 respectively, ACRO Biosystems) were coated at 0.5µg/mL in sodium carbonate buffer (pH9.6), 100µL per well, onto Nunc Maxisorp plates overnight at 4 °C. The assay was completed as described above.

Genscript SARS-CoV-2 neutralization antibody detection Kit. Secondary assessment of the RBD-ACE2 interaction blocking potential of the isolated nanobodies was performed using the Genscript SARS-CoV-2 Neutralization Antibody Detection Kit (#L00847, Genscript) according to the manufacturer’s instructions. Briefly, horse radish peroxidase conjugated RBD was diluted using the supplied assay buffer and then incubated with specified concentrations of nanobody for 30 min at 37 °C. The samples were then transferred onto the supplied human ACE2-coated assay plate and incubated for 15 min at 37 °C. The plate was washed using the supplied solution and the assay developed using a supplied 3,3′,5,5′-Tetramethylbenzidine reagent. The assay was stopped with 50 µL of the supplied Stop Solution. The assay was measured at 450 nm on a Biotek Synergy 2 plate reader.

SARS-CoV-2 pseudotyped lentivirus-based transduction fluorescence inhibition assay. All lentiviruses were propagated in HEK293T/17 cells (ATCC # CRL-11268) according to published Current Protocols in Neuroscience. Briefly, 293 T cells were transiently transfected with plasmids expressing SARS-CoV-2 spike protein (GenScript MC_0101081, human codon optimized, ER retention signal removed), psPAX2 (Addgene #12260), and a lentiviral transfer vector CD512-EF1a-RFP (System Biosciences CD512B-1) using Lipofectamine 2000. Supernatant was collected 48 h post transfection and concentrated by centrifugation at 600 rpm for 30 min. After a final wash, the plate was developed by addition of 100 µL tetramethylbenzidine (#T5656, Sigma-Aldrich). Assay development was stopped by addition of 50 µL 1 M sulfuric acid and the assay absorbance measured at 450 nm on a Biotek Synergy 2 plate reader. The assay was washed with 1xPBS 5 times between each step of the assay.

Expression of NIH-CoVnb-112 in Pichia pastoris. Increased expression yield and elimination of potential endotoxin from the bioprocess was achieved by cloning and expression of the nanobody in the methylotrophic yeast Pichia pastoris. The EasySelect Pichia Expression Kit (#K1740-01, Invitrogen) was used for cloning of the VHH sequence into expression vector pICZa by amplification of the pHEN2 containing phagemid with reformatting primers (Pichia-nominal-fwd: 5’TATCTCTCGAGAAGAAGATGTGCAGCTGGAG AGTCTG-3’ and Pichia-nominal-rev: 5’TGGTCTAGATTTAATGATTGATTGATGTGCGGCGC-3’). The reformatting primers introduce Xhol and XbaI (New England Biolabs) restriction endonuclease sites which allow for in-frame cloning of the VHH sequence with the secretion signal and includes a vector independent 6xHis tag at the C-terminus. The resulting expression vector was linearized using Scal-HF (New England Biolabs).
England Biolabs) and transformed into competent *Pichia* *pastoris* strain X-33 followed by selection on YPD agar with 1 M sorbitol and 100 μg/mL Zeocin. Resulting clones were confirmed for recombination and for phenotype by small scale expression. For scaled expression, selected clones were grown in buffered glycerol complex medium to an OD600 of 1 prior to addition of methanol to a final concentration of 0.5%. The culture was batch-fed every 24 h to a final methanol concentration of 0.5% until harvest of the culture supernatant. Spent cells were removed by centrifugation at 3000xG for 10 min at room temperature and the supernatant clarified by filtration through a 0.45 μm filtration unit. The media supernatant was concentrated using a Minimate tangential flow filtration system (#OAPMPUNV, Pall) and buffer exchanged using 1xPBS containing 10 mM imidazole. Purification was performed as described above using Ni-NTA affinity chromatography followed by polishing on a Superdex 75 Increase 10/300 GL size exclusion column.

### Nebulization stability assessment of NIH-CoVnb-112

Stability following nebulization of *Pichia pastoris* expressed NIH-CoVnb-112 was performed using an Aerogen Solo High-Performance Vibrating Mesh nebulizer placed in line with a custom glass bead condenser. A plastic culture tube was fitted with a glass-pore frit and filled with sterilized 5 mm borosilicate glass beads. A three-way stopcock was positioned distal to the frit to prevent pressurization during nebulization. A 2 mg/mL SEC polished NIH-CoVnb-112 solution was prepared in 0.9% normal saline to model potential patient delivery. The nanobody was nebulized and the resulting condensate incubated at 37 °C for 24 hr to mimic exposure to body temperature. The nebulized, 37 °C treated nanobody was then collected for stability assessments and protein concentration measurements by BCA assay. Equal masses of pre and post-nebulization samples were denatured in LDS sample buffer (Invitrogen) and run on a NuPAGE 12% Bis–Tris precast polyacrylamide gel with SeeBlue Plue 2 protein standards. Additional precondensate incubated at 37 °C for 24 hr to model exposure to body temperature. The nebulized, 37 °C treated frit to prevent pressurization during nebulization. A 2 mg/mL SEC polished NIH-CoVnb-112 solution was pre-frit and filled with sterilized 5 mm borosilicate glass beads. A three-way stopcock was positioned distal to the nebulizer placed in line with a custom glass bead condenser. A plastic culture tube was fitted with a glass-pore pastoris

### Stability of NIH-CoVnb-112 in human plasma

NIH-CoVnb-112 expressed in *Pichia pastoris* was diluted from a concentrated stock solution into apheresis derived pooled human plasma (#IPLA-N, Innovative Research) to a final concentration of 5 μM and incubated at 37 °C for either 24 hr or 48 hr with gentle rotation. An identical sample set was prepared at 5 μM in a solution containing 35 mg/mL recombinant human albumin (#A9731, Sigma-Aldrich) and incubated at 37 °C for either 24 hr or 48 hr with gentle rotation. A no-incubation control for each the plasma and recombinant human albumin conditions was prepared at 5 μM. The samples were prepared in a manner providing all conditions complete at the same time. The samples were diluted 1:10 with 1xPBS to yield a final nanobody concentration of 500 nM and Bio-layer Interferometry was performed using immobilized biotinylated SARS-CoV-2 S protein RBD to determine retention of binding potential.

### Determining melting temperature and refolding by circular dichroism

Circular Dichroism (CD) was performed using a Jasco J-815 Spectropolarimeter. For thermal stability measurements NIH-CoVnb-112 was diluted to 10 μg/mL in deionized water and placed in a quartz cuvette with 1 cm path length and CD was measured at an ultraviolet wavelength of 205 nm. NIH-CoVnb-112 heated from 25 °C to 85 °C at a rate of 2.5 °C/min while stirring and then cooled back to 25 °C at the same rate.

### Ethics approval

All experimental protocols were approved by the NIH Intramural Research Program and performed according to the relevant regulations.

The views expressed in this manuscript reflect those of the authors, and not those of the Uniformed Services University of the Health Sciences or the Department of Defense.

Note added in press: After the submission of this manuscript, the repurposed antiviral Remdesivir was approved by the US FDA and the novel monoclonal antibody Bamlanivimab received US FDA emergency use authorization to treat SARS-CoV-2.

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Author contributions
D.L.B. and T.J.E. conceived the experiments. T.J.E., N.P.M., G.P.A., and E.R.G. performed experiments. D.L.B. and T.J.E. wrote the manuscript.

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Correspondence and requests for materials should be addressed to D.L.B.

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