An alpaca nanobody neutralizes SARS-CoV-2 by blocking receptor interaction

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Abstract

SARS-CoV-2 is the etiologic agent of COVID-19, currently causing a devastating pandemic for which pharmacological interventions are urgently needed. The virus enters host cells through an interaction between the spike glycoprotein and the angiotensin converting enzyme 2 (ACE2) receptor. Directly preventing this interaction presents an attractive possibility for suppressing SARS-CoV-2 replication. Here we report the isolation and characterization of an alpaca-derived single domain antibody fragment, Ty1, that specifically targets the receptor binding domain (RBD) of the SARS-CoV-2 spike, directly preventing ACE2 engagement. The nanobody binds with high affinity in the low nM range to the RBD, occluding ACE2. A cryo-electron microscopy structure of the bound complex at 2.9 Å resolution reveals that Ty1 binds to an epitope on the RBD accessible in both the ‘up’ and ‘down’ conformations and that Ty1 sterically hinders RBD-ACE2 binding. This 12.8 kDa nanobody does not need an Fc domain to neutralize SARS-CoV-2, and can be expressed in high quantities in bacteria, presenting opportunities for manufacturing at scale. Ty1 is therefore an excellent candidate as an intervention against COVID-19.
Main
SARS-CoV-2 emerged as the etiologic agent of COVID-19 in Wuhan, China in late 2019. In the comparatively short time since then, it has achieved pandemic status, causing more than 7.3 million cases, leading to at least 414,000 deaths and rising. Accordingly, the WHO declared the pandemic to be a public health emergency of international concern. A safe and effective vaccine is urgently needed, but requires time to develop. In the meantime, and indeed also in the post-vaccine era, highly specific and potent antiviral interventions are needed. Many generic or repurposed candidates are in trials, but so far results have been unremarkable. Since the virus is newly emerged, specifically designed drugs have not yet reached late phase trials. When available, specific antiviral drugs or antibody therapies will be used to protect individuals at risk and their widespread use will allow immunologically naïve populations to exit lockdowns more safely.

The virus is closely related to SARS-CoV-1, both being members of the lineage 2 betacoronaviruses. Cell entry of both viruses is achieved by first binding to the cell surface expressed receptor angiotensin-converting enzyme 2 (ACE2), followed by conformational changes in the viral spike glycoprotein trimer and subsequent membrane fusion. The affinity of SARS-CoV-2 receptor binding domain (RBD) for ACE2 is considerably higher than that for SARS-CoV-1\(^1,2\), supporting efficient cell entry and likely contributing to pathogenesis. The RBD is a globular domain situated on the distal surface of the spike protein. Two conformations have been observed in the stabilized trimer. Specifically, one conformation where one RBD is ACE2-accessible while two are not and one conformation where all three RBDs are down, i.e. receptor inaccessible\(^2,3\). As the receptor-engaging part of the spike, the RBD is an attractive target for coronavirus neutralization, and a number of conventional neutralizing monoclonal antibodies that target the RBD and block receptor binding have already been isolated from convalescent patients\(^4-6\).

Camelid-derived single domain antibody fragments, also called VHHs or nanobodies, offer several advantages over conventional antibodies as candidates for specific therapies. Despite being approximately one tenth of the size of a conventional antibody, they retain specificity and affinity similar to conventional antibodies while being far easier to clone, express and manipulate. They are readily expressed in bacteria in large quantities and show high thermal stability and solubility, making them easily scalable and extremely cost effective. Their modularity means that they can be oligomerized to increase avidity or to increase serum half-life\(^7\). Critical to their use as antivirals in humans, they can easily be humanized with existing protocols\(^8\). Importantly, they have proven to be highly potent inhibitors of viral infections in vivo, particularly respiratory infections\(^9,10\).
We immunized one alpaca with SARS-CoV-2 S1-Fc and RBD in a 60-day immunization schedule. We generated a phage display library and performed two consecutive rounds of phage display, followed by an ELISA-based binding screen (Fig. 1a). We isolated one nanobody, Ty1, that binds specifically to the RBD of the SARS-CoV-2 spike glycoprotein. In parallel we performed next generation sequencing (NGS) on the baseline and post-enrichment libraries, and quantified variant frequency before and after each enrichment step. Ty1 exhibited the greatest fold-change in frequency among all nanobody variants, increasing over 10,000-fold from baseline to after the second enrichment round (Fig.1b). We report the amino acid sequence of Ty1 in Figure 1c.

To determine whether Ty1 neutralized SARS-CoV-2 we employed an in vitro neutralization assay using lentiviral particles pseudotyped with the SARS-CoV-2 spike protein. Ty1 neutralized SARS-CoV-2 pseudotyped viruses at an IC$_{50}$ of 0.77 µg/ml (54 nM) (Fig. 2a). No neutralization of a lentivirus pseudotyped with VSV-G by Ty1 was evident, and a control nanobody produced and purified in the same way, but specific for influenza A virus nucleoprotein showed no evidence of neutralization of SARS-CoV-2 pseudotyped viruses. When Ty1 was expressed in mammalian cells as a Fc-fusion protein the potent neutralization could be further increased to ~5 ng/ml (Fig. S1).

To confirm that Ty1 is directed specifically against the SARS-CoV-2 spike protein, we characterized the specificity of Ty1 by flow cytometry. We site-specifically conjugated a fluorophore to the C-terminus of Ty1 by means of a Sortase A reaction and copper-free click chemistry (Ty1-AS635P) and stained untransfected cells and cells transiently transfected with SARS-CoV-2 spike under permeabilizing conditions (Fig. 2b). While untransfected and unstained cells displayed similar signals, cells expressing the viral spike protein showed a strong shift in fluorescence intensity when stained with Ty1-AS635P. The apparent double peak likely reflected the varying efficiency of this transient transfection. To determine if the same probe can be exploited to recognize the viral spike protein in immunofluorescence, we infected Vero E6 cells with infectious SARS-CoV-2 at MOI 1 for 24 hours and stained the fixed and permeabilized cells with Ty1-AS635P and anti-dsRNA antibody (Fig. 2c). While uninfected cells showed no signal, infected cells were strongly labelled with both dsRNA antibody and Ty1-AS635P. Thus, Ty1 recognized the viral spike glycoprotein with high specificity in its native conformation in SARS-CoV-2-infected cells. Importantly, the low background in both experiments also suggested that Ty1 is a highly specific and suitable tool for research, diagnostics and therapy.

To understand the mechanism of neutralization, we evaluated the effect of Ty1 on RBD binding to ACE2. We site-specifically conjugated a fluorophore to the C-terminus of the RBD (RBD-AS635P) and used this probe to stain ACE2 expressing HEK293T cells (Fig. 2d). Preincubation of RBD-AS635P with unlabeled
Ty1 resulted in a significant reduction of ACE2 staining, while preincubation with the control nanobody NP-VHH1 had no such effect. This result indicated that Ty1 directly prevents binding of SARS-CoV-2 RBD to its host cell receptor ACE2.

Specific and high-affinity binding of Ty1 to the RBD was also demonstrated in kinetic bio layer interferometry (BLI) experiments. Dipping of surface-immobilized nanobodies into monomeric RBD solutions at 550 nM yielded binding responses with fast association kinetics and amplitudes reaching 1.5 nm only for Ty1 but not NP-VHH1 (red and blue curves, respectively, in Figure 3a). Titration experiments performed under normal (280 mM) and high salt (680 mM) conditions revealed concentration-dependent kinetic response curves for binding of RBD to Ty1 (Figs. 3b and S2a, respectively). The derived semi-log concentration-response curves revealed sigmoidal line-shapes with fitted apparent K_D-values of 8 ± 1.5 and 13 ± 1.5 nm for binding at normal and high salt conditions, respectively. Local fits to individual sensorgrams applying the standard 1:1 binding model appeared reasonable for the association phases at lower to intermediate RBD concentrations, as well as for all dissociation curves when fits were allowed to stay above zero (grey lines Figs. 3b and S2a 1st panel). However, the model deviated from the observed data at higher RBD concentrations. Instead almost perfect fits were obtained when the same data were analyzed in terms of a Bayesian two-dimensional distribution of K_D and k_off-rate constants to address heterogeneous ligand site populations on the sensor surface\textsuperscript{12-14}. For the two titrations at low and high salt conditions, distinct peaks at K_D- and k_off-rate values of 5-9 nM and 4-5*10\textsuperscript{-3} s\textsuperscript{-1} were obtained (Fig. 3b and S2a 4th panel). In both conditions, a second elevated plateau with K_D- and k_off-rate values of about 7 nM and 1-4*10\textsuperscript{-7} contributed significantly to the observed sensorgrams. Since most high-affinity protein-protein interactions in the nM-range have dissociation rates in the 1*10\textsuperscript{-3} s\textsuperscript{-1} range\textsuperscript{15}, we attribute the first defined peak as the relevant Ty1:RBD interaction. The second broad plateau is likely caused by RBD competition and rebinding effects on the sensor surface, as well as heterogeneous ligand populations\textsuperscript{14,16,17}. The orthogonal biophysical method isothermal titration calorimetry (ITC)\textsuperscript{18} confirmed the high affinity binding of Ty1 to RBD with a K_D of 9 nM (with estimated bounds of 1 and 70 nM) characterized by an exothermic enthalpy of about -10 ± 0.5 kcal/mole (Fig. 3c, left panel). Exothermic binding was already evident from the three initial relatively constant negative spikes that were caused by the injection of Ty1 to RBD (Fig. 3c, right panel). The amplitude of the following three to four spikes returned to baseline demonstrating saturation of the available RBD sites by Ty1 binding. Notably, return to baseline was accompanied by the appearance of preceding positive spikes (Fig. 3c, left panel and Fig. S2b). These spikes were also detected when Ty1 was injected into the buffer (HBS) and thus treated as Ty1 dilution effects during data analysis. Injection of NP-VHH1 into RBD did not cause any binding or dilution heat changes above background noise. It
should be noted that the ITC measurements were performed at the lowest possible protein concentrations
to derive $K_D$-values in the low nM range, while still being able to detect interaction heat above
background noise signals that were at about -0.17 ucal/s (maximum spike amplitude) and ± 0.05 ucal/s,
respectively. Altogether, we concluded from these results that RBD bound to surface-immobilized Ty1
with high affinity of about 5-10 nM.

To understand the structural basis underlying the potent neutralization of SARS-CoV-2 we performed a
cryo-EM structure determination of the prefusion-stabilized spike ectodomain in complex with Ty1. The
cryo-EM reconstruction reaches an overall resolution of 2.9 Å (Fig. 4a; 0.143 FSC) with strong variation
of estimated local resolution from high resolution in the core of the spike trimer to relatively low
resolution in the top of the spike. Nevertheless, the current reconstruction clearly shows that the spike
retains only one main conformation with one RBD ‘up’ and two RBDs ‘down’. Importantly, all three
RBDs are decorated in their upper parts with a Ty1 nanobody. The nanobodies retain a similar binding
orientation to the RBD whether the RBD is found in the ‘up’ or ‘down’ conformation (Fig. 4a,b). Primary
interactions with the RBD are through the CDRs. Specifically, CDR1 interacts with RBD T470 and
V483-E484 and CDR3 interacts with RBD Y449, F490 and Q493. Interestingly, CDR2 does not form any
interactions with the RBD, instead it stabilizes the conformation of CDR1 in the RBD bound mode and
thereby acts indirectly to potentiate the Ty1-RBD binding.

Since ACE2 can only be bound by an RBD in the ‘up’ conformation, the current cryo-EM reconstruction
clearly shows that ACE2 binding is sterically hindered from two sides (Fig. 4c). Specifically, ACE2
binding is blocked both by the Ty1 nanobody bound to the RBD in the ‘up’ conformation and the
neighboring RBD in the ‘down’ conformation. Hence, ACE2 binding is sufficiently hindered with any
two of the available three binding RBD sites in the spike trimer.

The current coronavirus pandemic has drastic consequences for the world’s population, and vaccines,
antibodies or antivirals are urgently needed. Neutralizing antibodies can block virus entry at an early step
of infection and potentially protect individuals that are at high risk of developing severe disease. We
report the identification and characterization of a SARS-CoV-2 RBD-specific single domain antibody
fragment (nanobody) termed Ty1 that potently neutralizes the virus. We identified Ty1 by binding assay
after two consecutive rounds of phage display, simultaneously monitoring sequence enrichment by NGS.
Although Ty1 exhibited the greatest fold-enrichment in the NGS analysis, multiple additional nanobodies
exhibited enrichment of varying extent across both rounds. As the correlation between phage display
enrichment and neutralization is likely imperfect, further analyses of our libraries will likely yield other
potent SARS-CoV-2 neutralizing nanobodies. In addition to neutralization activity, we also show that Ty1 can be used as a detection reagent in flow cytometry and immunofluorescence demonstrating its suitability as a research tool and for diagnostics.

It should be noted that the nanobody Ty1 can be readily produced in bacteria at very high yield (in excess of 30 mg/L culture), making it an excellent candidate for a low-cost, scalable antiviral agent against SARS-CoV-2, and we provide the amino acid sequence, encouraging direct exploitation as such. While nanobodies capable of binding SARS-CoV-2 spike have recently been isolated, these were generated after SARS-CoV-1 spike immunization\(^\text{19}\), or PCR maturation\(^\text{20}\). Also, in both cases a fusion to human Fc domain is required for neutralization of SARS-CoV-2, precluding expression in bacterial culture. Naive libraries of human single-domain antibodies (sdAbs) have also been screened to identify SARS-CoV-2 spike-specific nanobodies\(^\text{21,22}\), but they lack detailed structural information. Ty1 represents the first single-domain antibody isolated from an animal specifically immunized with a SARS-CoV-2 protein.

Future work will aim to improve the potency and potential efficacy of Ty1 through various strategies. For example, mutational scanning may yield potency improvements to Ty1. Also, since Ty1 already neutralizes as a monomeric protein, the generation of homodimeric or trimeric fusion constructs is expected to further increase its neutralization activity. Indeed, fusion of Ty1 to a human IgG1-Fc dramatically improved the IC\(_{50}\) of this molecule, to approximately 5 ng/ml (Fig. S2). Additional strategies will explore linker-based constructs that chain multiple copies of Ty1 together, which may provide similar improvements in potency while retaining the possibility of being expressed in bacteria. Ty1 may additionally be a useful component of a bi-specific or tri-specific antibody, which could combine epitope specificities to increase the mutational barrier to viral escape. Based on our work, we hope that Ty1 will be investigated as a candidate for antiviral therapy.

**Methods**

**Cells and virus.** Vero E6 cells (ATCC-CRL-1586) and HEK293T cells (ATCC-CRL-3216) were maintained in Dulbecco’s Modified Eagle Medium (Gibco) supplemented with 10% fetal calf serum and 1% Penicillin-Streptomycin and cultured at 37°C in a humidified incubator with 5% CO\(_2\). A HEK293T cell line engineered to overexpress human ACE2 (HEK293T-ACE2) was generated by the lentiviral transduction of HEK293T cells. Briefly, lentiviruses were produced by co-transfecting HEK293T cells with a plasmid encoding VSV-G (Addgene cat#12259), a lentiviral Gag-Pol packaging plasmid (Addgene cat#8455), and a human ACE2 transfer plasmid. Virions were harvested from the supernatant, filtered through 0.45 µm filters, and used to transduce HEK293T cells. All cell lines used for experiments were
negative for *Mycoplasma* as determined by PCR. Infectious SARS-CoV-2\(^2\) was propagated in Vero E6 cells and titrated by plaque assay.

**Proteins and probes.** The plasmid for expression of the SARS-CoV-2 prefusion-stabilized spike ectodomain with a C-terminal T4 fibritin trimerization motif was obtained from\(^2\). The plasmid was used to transiently transfect FreeStyle 293F cells using FreeStyle MAX reagent (Thermo Fisher Scientific). The S ectodomain was purified from filtered supernatant on Streptactin XT resin (IBA Lifesciences), followed by size-exclusion chromatography on a Superdex 200 in 5 mM Tris pH 8, 200 mM NaCl.

The RBD domain (RVQ – VNF) of SARS-CoV-2 was cloned upstream of an enterokinase cleavage site and a human Fc. This plasmid was used to transiently transfect FreeStyle 293F cells using the FreeStyle MAX reagent. The RBD-Fc fusion was purified from filtered supernatant on Protein G Sepharose (GE Healthcare). The protein was cleaved using bovine enterokinase (GenScript) leaving a FLAG-tag at the C-terminus of the RBD. Enzyme and Fc-portions was removed on HIS-Pur Ni-NTA resin (Thermo Fisher Scientific) and Protein G sepharose (GE Healthcare) respectively, and the RBD was purified by size-exclusion chromatography on a Superdex 200 in 50 mM Tris pH 8, 200 mM NaCl.

In addition, the RBD domain (RVQ – VNF) was cloned upstream of a Sortase A recognition site (LPETG) and a 6xHIS tag and expressed in FreeStyle 293F cells as described above. RBD-HIS was purified from filtered supernatant on His-Pur Ni-NTA resin, followed by size-exclusion chromatography on a Superdex 200.

The nanobodies were cloned for expression in the pHEN plasmid with a C-terminal Sortase recognition site (LPETG) and a 6xHIS tag. This plasmid was used to transform BL21 cells for periplasmic expression. Expression was induced with 1 mM IPTG at OD600 = 0.6; cells were grown overnight at 30°C. Nanobodies were retrieved from the periplasm by osmotic shock and purified by Ni-NTA affinity purification and size-exclusion chromatography.

For mammalian expression, the sequence encoding the nanobody Ty1 was cloned upstream of an human IgG1. This plasmid was used to transiently transfect FreeStyle 293F cells using the FreeStyle MAX reagent. The Ty1-Fc fusion was purified from filtered supernatant on Protein G Sepharose followed by size-exclusion chromatography.

Biotinylated and fluorescent probes were generated using Sortase A as described here\(^24\) and here\(^25\). In brief, nanobodies were site-specifically biotinylated on the C-terminus using Sortase A 5M. Nanobody at a concentration of 50 μM was incubated with sortase A 5M (5 μM), GGGK-biotin (200 μM) in 50 mM...
Tris, pH 7.5, 150 mM NaCl, 10 mM CaCl$_2$, for 2 hours at 25°C. Unreacted nanobody and sortase was removed with Ni-NTA resin and excess GGGK-biotin was removed using Zeba spin desalting columns (0.5 mL, 7k MWCO, Thermo Fisher Scientific).

To generate the fluorescently labeled probes, first a dibenzocyclooctyne-amine (DBCO-amine, Sigma Aldrich) was attached via sortase A to the nanobody or the RBD. (Reaction conditions: 50 μM RBD or nanobody, 50 μM Sortase A 5M, 8 mM DBCO-amine in 50 mM Tris pH 7.5, 150 mM NaCl, 10 mM CaCl$_2$, 2 hours, 25°C). Unreacted probe, sortase and excess DBCO-amine was removed using Ni-NTA resin and PD-10 columns (GE Healthcare) respectively. Abberior Star 635P-azide (Abberior GMBH) was attached to the DBCO-labeled proteins in a copper-free click chemistry reaction. Unreacted fluorophore was removed on PD-10 column (RBD) or size-exclusion chromatography (nanobody).

**Alpaca immunization.** Alpaca immunization and phage display was performed similarly as described here$^{26}$ and here$^{27}$. In brief, the adult male alpaca Tyson at PreClinics, Germany, was immunized 4 times in a 60-day immunization schedule. SARS-CoV-2 S1-sheep-Fc (Native Antigen Company, SKU: REC31806) was used for the first two immunization, and SARS-CoV-2 RBD produced in FreeStyle 293F cells was used for the last two immunizations.

**Library generation and nanobody isolation.** After the final boost, RNA was isolated from PBMCs (RNA Plus mini kit, Qiagen). For cDNA synthesis, SuperScript III RT (Thermo Fisher Scientific) was used with a combination of oligo(dT), random hexamers, or gene specific primers (AL.CH2, ATGGAGAGGACGTCCTTGGGT and AL.CH2.2 TTCGGGGGGAAGAYRAAGAC)$^{27}$. Nanobody sequences were PCR amplified and cloned into a phagemid vector for expression as pIII fusion. TG1 cells (Lucigen) were transformed with this library by electroporation.

Cells were inoculated with VCSM13 helper phage, and the resulting phage was enriched in two consecutive rounds of phage display on RBD immobilized on magnetic beads. After the second round of phage display, individual bacterial colonies were picked in a 96 well format, grown until OD = 0.6 and nanobody expression was induced by addition of 1 mM IPTG. After 16 hours incubation at 30°C, bacterial supernatant was used as primary detection reagent in an ELISA coated with RBD or S ectodomain. Bound nanobodies were detected with anti-E tag (Bethyl laboratories) secondary antibody. Positive clones were sequenced and cloned into the pHEN expression vector for further characterization.

**Amino acid sequence of Ty1**
Next generation sequencing (NGS) and analysis of nanobody libraries. Plasmids from nanobody libraries before enrichment, and after each enrichment step, were amplified for 13 cycles using Q5 High-Fidelity 2X Master Mix (NEB) according to manufacturer's instructions, using primers:

MAQVQLVETGGGLVQPGGSLRLSCAASGFTFSSVYMNWVRQAPGKGPEWVSRISPNSGNIGYT

DSVKGRFTISRDNAKNTLYLQMNNLKPEDTALYYCAIGLNLSSSSVRGQGTQVTVSS

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MAQVQLVETGGGLVQPGGSLRLSCAASGFTFSSVYMNWVRQAPGKGPEWVSRISPNSGNIGYT

DSVKGRFTISRDNAKNTLYLQMNNLKPEDTALYYCAIGLNLSSSSVRGQGTQVTVSS

CACTCTTTCCCTACACGACGCTCTTCCGATCT

CTCGCGGCCCAGCCGGCCATGG and

GGAGTTCAGACGTGTGCTCTTCCGATCTACCGGCGCACCACTAGTGCA, annealing at 72°C.

Illumina indexing primers were added using an additional 9 cycles, with Kapa HiFi. Amplicons were size selected using Agencourt AMPure XP beads (bead ratio: 1:1), and were pooled at ratios of 6:2:1 for pre:post-1:post-2 libraries, to account for the reduction in diversity expected during enrichment, and sequenced on an Illumina MiSeq using the MiSeq Reagent Kit v3 (2x300) MS-102-3003.

Paired-end reads were merged using USEARCH11, and then processed in the Julia language, primarily using the NextGenSeqUtils.jl package. Briefly, reads are trimmed of primer sequences, and deduplicated, maintaining read frequencies. Variant frequencies are calculated as combined frequency of any reads matching a variant within 3% nucleotide divergence, using a kmer-based distance approximation for rapid database search. Any reads with counts >3 from the second enrichment library are searched for their variant frequencies across all databases. When calculating enrichment, to avoid zeros due to sampling and to regularize against over-sensitivity to low-frequency baseline variants, all frequencies are increased by the reciprocal of the size of the pre-enrichment database.

Neutralization Assay. Pseudotyped viruses were generated by the co-transfection of HEK293T cells with plasmids encoding the SARS-CoV-2 spike protein harbouring an 18 amino acid truncation of the cytoplasmic tail, a plasmid encoding firefly luciferase, and a lentiviral packaging plasmid (Addgene cat#8455) using Lipofectamine 3000 (Invitrogen). Media was changed 12-16 hours after transfection, and pseudotyped viruses were harvested at 48- and 72-hours post transfection, filtered through a 0.45 µm filter, and stored at -80°C until use. Pseudotyped virus neutralization assays were adapted from protocols previously validated to characterize the neutralization of HIV, but with the use of HEK293T-ACE2 cells. Briefly, pseudotyped viruses sufficient to generate ~100,000 RLUs were incubated with serial dilutions of nanobodies for 60 min at 37°C. Approximately 15,000 HEK293T-ACE2 cells were then added to each well and the plates were incubated at 37°C for 48 hours. Luminescence was then measured using Bright-Glo (Promega) per the manufacturer’s instructions on a GM-2000 luminometer (Promega) with an integration time of 0.3s.
**Flow Cytometry.** Cells were trypsinized and fixed in 4% formaldehyde/PBS and stained with RBD-AS635P under non-permeabilizing conditions or with Ty1-AS635P under permeabilizing conditions. Fluorescence was quantified using a BD FACSCelesta and the FlowJo software package.

**Immunofluorescence.** Vero E6 cells were seeded onto coverslips in a 24-well plate and incubated overnight at 37°C/5% CO₂. Cells were infected with SARS-CoV-2 at a MOI of 1 for 24 h. Cells were fixed with 4% (v/v) formaldehyde, permeabilized in 0.5% Triton X-100 and blocked in 5% horse serum. Cells were incubated with anti-dsRNA antibody (1:2000, J2 Scicons) for 1 hour at room temperature followed by 1 hour staining with the secondary antibody anti-mouse-Alexa Fluor 488 (1:2000, Thermo Fisher Scientific), Hoechst (1:1000, Invitrogen) and Ty1-AS635P (0.05 µg/mL). Coverslips were mounted in mounting media and images were obtained using Zeiss Axiovert microscope and processed using Adobe Photoshop.

**Biophysical Biolayer interferometry (BLI) and isothermal titration calorimetry (ITC).** BLI was performed using single-use high-precision streptavidin biosensors (SAX) on an eight-channel Octet RED instrument according to manufacturer’s protocols (Fortebio). Assays were performed in 2xPBS comprising 0.05% Tween-20 (PBST). Biotinylated nanobodies Ty1 and NP-VHH1 were loaded at concentrations between 30 and 250 nM followed by quenching using biocytin to reach final sensor loads of between 0.15 and 0.7 nm. For the comparative binding test, the eight sensors were divided into two sets, each comprising double sample as well as single reference and single control sensors. Sample and reference sensors were loaded with respective nanobodies. The SAX control was only quenched. Loading of the two sets was performed consecutively to reach similar immobilization levels, while subsequent association and dissociation phases were performed simultaneously. For association, the sample and control sensors were dipped into RBD, while the reference sensor was dipped into PBST. For titration experiments, all sensors were loaded simultaneously. During association one of the sensors was used as reference and only dipped into PBST. Raw data were pre-processed, analyzed and fitted by applying the 1:1 binding model as implemented in the manufacturer’s software. Bayesian analysis to obtain the two-dimensional distribution of $K_D$ and $k_{off}$-rate values were performed using Evilfit. The shown titration data were processed applying reference sensor subtraction and Savitzky-Golay filter operations.

For ITC, proteins were exchanged to 2xHBS-buffer (50 mM HEPES, 300 mM NaCl, pH 7.5) and isolated as single peak populations by Superdex-200 HR10/300 size-exclusion chromatography. ITC measurements were performed using an ITC200 calorimeter (GE Healthcare). The cell temperature was set to 37°C and the syringe stirring speed to 750 rpm. Before each experiment, the RBD and nanobodies
were loaded into the cell and syringe at concentrations of 4 μM and 75 μM, respectively. Data and binding parameters were analyzed using the MicroCal PeakITC software (Malvern). The integrated heat versus molar ratio plots of the Ty1:RBD interactions were obtained by subtracting the Ty1 dilution heat uptake from the binding data. The NP-VHH1:RBD data were only baseline-corrected, since dilution effects were not evident.

Raw and processed BLI/ITC data were imported into Rstudio for visualization and further analysis\textsuperscript{32-34}. Data along with analysis R scripts will be made publicly available via Github and/or DataDryad.

**Cryo-EM sample preparation and imaging.** Spike trimer (0.7 mg/ml) and Ty1 (1.3 mg/ml) were mixed in a 1:8 molar ratio and incubated on ice for 5 minutes. A 3-μl aliquot of the sample solution was applied to glow-discharged CryoMatrix holey grids with amorphous alloy film (Zhenjiang Lehua Technology) in a Vitrobot Mk IV (Thermo Fisher Scientific) at 4 degrees and 100% humidity (blot 10 s, blot force 3).

Cryo-EM data collection was performed with EPU 2.7 (Thermo Fisher Scientific) using a Krios G3i transmission-electron microscope (Thermo Fisher Scientific) operated at 300 keV in the Karolinska Institutet 3D-EM facility. Images were acquired in nanoprobe EFTEM mode with a slit width of 10 eV using a GIF 967 energy filter (Ametek) and a K3 detector (Ametek) during 2.4 seconds with a dose rate of 4.1 e-/px/s resulting in a total dose of 38 e-/Å\(^2\) fractionated into 40 movie frames. Motion correction, CTF-estimation, fourier binning (to 1.02 Å/px), picking and extraction in 428 pixel boxes were performed on the fly using Warp\textsuperscript{35}.

A total of 13,589 micrographs were selected based on an estimated resolution cutoff of 4 Å and defocus below 2 microns and 573,036 particles were picked by Warp. Extracted particles were imported into cryoSPARC v2.15.0\textsuperscript{36} for 2D classification, 3D classification and non-uniform 3D refinement. The particles were processed with C1 symmetry throughout. After 2D classification (300 classes) 354,678 particles were retained and used to build three ab-initio 3D reconstructions. These were further processed for heterogeneous refinement that resulted in one reconstruction showing high-resolution structural features in the core of the spike. One round of homogenous refinement followed by non-uniform refinement resulted in a final reconstruction to an overall resolution of 2.9 Å (0.143 FSC) using 210,832 particles. To minimise the effects of loss of delocalised information due to tight boxing\textsuperscript{37}, the particles were re-extracted with 600 pixel box size. In addition, localized reconstruction\textsuperscript{38} were performed using particles where all parts of the spike except the N-terminal domains, the RBDs and the nanobodies had been subtracted\textsuperscript{39}. The combined effects of these three approaches significantly increased the level of density detail in the upper part of the spike.
Model building and structure refinement. A structure of the 2019-nCoV spike protein trimer\textsuperscript{19} (PDB: 6VS6) was used as a starting model for model building. The model was extended and manually adjusted in COOT\textsuperscript{40}. The Nanobody structure was homology modelled using SWISS-MODEL\textsuperscript{41} taking PDB:5JMR\textsuperscript{42} as a template. The missing regions of the RBD domains were built based on the RBD-ACE2 crystal structure (PDB: 6LZG)\textsuperscript{43}. Structure refinement and manual model building were performed using COOT and PHENIX\textsuperscript{44} in interspersed cycles with secondary structure and geometry restrained. All structure figures and all EM density-map figures were generated with UCSF ChimeraX\textsuperscript{45}.

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Author Contributions

L.H., L.V.P., D.J.S., T.S., A.M.M., M.C. and B.M. performed experiments and analyzed data. H.D. and B.M.H. collected and processed cryo-EM data, built the model and performed the structural interpretation of the data with the corresponding paper sections. A.A. and G.B.K.H. gave critical advice. L.H., B.M. and G.M.M. conceived the study and wrote the initial draft. All authors contributed to the final draft.

Competing Interests Statement

The authors declare no conflicts of interest.
Figure 1 - Nanobody discovery.

a, Overview of the nanobody generation process. b, Variant frequencies quantified by NGS across successive enrichment steps. Identified using RBD bait, Ty1 exhibits the greatest total fold change of all nanobodies, increasing in proportion over 10,000-fold between initial and final libraries. c, Sequence of Ty1. Complementarity-determining regions (CDRs) are indicated. Color scheme according to ClustalX.
Figure 2 - Nanobody Ty1 neutralizes SARS-CoV-2 and specifically recognizes SARS-CoV-2 spike glycoprotein.

a, VSV G or SARS-CoV-2 spike pseudotyped lentivirus (PSV) was incubated with a dilution series of Ty1 or control nanobody (influenza NP-VHH1). Infectivity relative to cells infected with pseudotyped virus in the absence of nanobody is shown. Neutralization by Ty1 was repeated in duplicate across 6 assays, and the error bars represent the standard deviation. b, Cells were transfected with a plasmid harboring the SARS-CoV-2 spike for 24 h. Cells were fixed, permeabilized and stained with Ty1-AS635P (black and red) or left unstained (grey). Cells were analyzed by flow cytometry. Cell counts are presented as % of max (representative histogram). c, Vero E6 cells were infected with SARS-CoV-2 at a MOI of 1 for 24 hours. Cells were fixed, permeabilized and stained for DNA (blue), dsRNA (green) and with Ty1-AS635P (red). Pictures were taken by fluorescence microscopy and representative examples are shown. d, ACE2 expressing HEK293T cells were trypsinized, fixed and stained with RBD-AS635P alone, or preincubated with IAV NP-VHH1 or Ty1. Cells were analyzed by flow cytometry.
Figure 3 - RBD bound to Ty1 with 5-10 nM affinities in biophysical BLI and ITC measurements

a, RBD bound to surface-immobilized Ty1 (red curves), but not to NP-VHH1 (blue curves). Almost equal nanobody immobilization levels of about 0.7 nm were obtained by first loading Ty1 and then NP-VHH1. In BLI, binding of molecules over time is recorded as sensorgrams recording the shift in wavelengths (unit: nm) due to an increase in the optical thickness of the surface layer. b, (1st panel) RBD titration sensorgrams obtained at high salt concentrations revealed concentration-dependent responses. Sensorgrams are color-coded according to the log2 RBD concentration scale. Standard and Bayesian 1:1 binding models are shown as grey and black solid lines. (2nd panel) Pseudo-equilibrium response values plotted against the logarithmic RBD concentration revealed sigmoidal binding curves that were fit to the single-site interaction model yielding $K_D$-values in the low nM range. $K_D$-values and standard deviations are shown as solid and dotted lines, respectively. (3rd panel) Sensor immobilization levels are shown as jittered box plots. (4th panel) Two-dimensional distribution of dissociation rate ($k_{off}$) and affinity ($K_D$) values obtained from the Bayesian and standard 1:1 model fits, visualized as densities according to depicted normalized height scale and single black crosses, respectively. See related Figure S2A. Plots for RBD/Ty1 titrations at low salt condition with same legends and scales. c, (1st panel) ITC demonstrated high affinity binding of Ty1 to RBD with fitted $K_D$- and binding enthalpy ($DH$) mean values of 9 nM (with estimated bounds of 1 and 70 nM) and -10 ± 0.5 kcal/mole, respectively, for two binding experiments (red). Injection of NP-VHH1 into RBD yielded heat changes at background level (blue). (2nd panel) Baseline-corrected heat changes plotted for two Ty1/RBD, a single Ty1/HBS buffer
and NP-VHH1/RBD titration experiments. Negative and positive heat changes are colored according to the red-to-blue gradient. See related Figure S2B: Same figure on larger scale to highlight the Ty1 into buffer dilution spikes.
Figure 4 - Ty1 binds to the RBD in ‘up’ and ‘down’ conformation and prevents ACE2 engagement

a, Cryo-EM reconstruction to an overall resolution of 2.9 Å (0.143 FSC) of the spike trimer with three bound molecules of Ty1. b, Atomic model (cartoon representation) of trimeric spike in complex with three molecules of Ty1. Three chains of spike are shown in three different colors. The RBD of chain A (light green) is present in ‘up’ conformation while the other two RBDs are captured in ‘down’ conformation. The ACE2 interaction surface of RBD1 and the Ty1 interaction surface is highlighted. Magnified view of RBD2 (in ‘down’ conformation) and Ty1 interaction is shown. CDR1,2 and 3 of Ty1 are shown in blue, green and red respectively. c, Ty1 shows a two-pronged inhibition of ACE2 receptor binding through binding the RBD in the ‘up’ conformation and by binding to the neighbouring RBD in the ‘down’ conformation. Binding of Ty1 to RBDs (both ‘up’ and ‘down’) would make ACE2 interaction surface inaccessible for ACE2.
Supplemental Figure 1 - Increased neutralization of a Ty1-Fc construct

SARS-CoV-2 spike pseudotyped lentivirus (PSV) was incubated with a dilution series of Ty1 or Ty1-Fc. Infectivity relative to cells infected with pseudotyped virus in the absence of nanobody is shown. Neutralization was repeated in duplicate across 3 assays, and the error bars represent the standard deviation.
Supplemental Figure 2

a, related to Fig 3b, plots for RBD/Ty1 titrations at low salt condition with same legends and scales.

b, related to Fig 3c, Same data plotted on larger scale to highlight the Ty1 into buffer dilution spikes.

c, Samples after ITC were analyzed by SDS-PAGE.