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Authors
Bogdanoff, WA
Perez, EI
Lopez, T
et al.

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Structural basis for the escape of human astrovirus from antibody neutralization: broad implications for rational vaccine design

Walter A. Bogdanoff\textsuperscript{a}, Edmundo I. Perez\textsuperscript{a}, Tomás López\textsuperscript{b}, Carlos F. Arias\textsuperscript{b}, and Rebecca M. DuBois\textsuperscript{a#}

\textsuperscript{a}Department of Biomolecular Engineering, University of California Santa Cruz, Santa Cruz, California, USA

\textsuperscript{b}Departamento de Genética del Desarrollo y Fisiología Molecular, Instituto de Biotecnología, Universidad Nacional Autónoma de México, Cuernavaca, Morelos, México

Running Head: Human astrovirus neutralization escape

\textsuperscript{a#}Address correspondence to Rebecca M. DuBois, rmdubois@ucsc.edu.

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Abstract

Human astroviruses are recognized as a leading cause of viral diarrhea worldwide in children, immunocompromised patients, and the elderly. There are currently no vaccines available to prevent astrovirus infection, however antibodies developed by healthy individuals during previous infection correlate with protection from reinfection, suggesting that an effective vaccine could be developed. In this study, we investigated the molecular mechanism by which several strains of human astrovirus serotype 2 (HAstV-2) are resistant to the potent HAstV-2-neutralizing monoclonal antibody PL-2 (mAb PL-2). Sequencing of the HAstV-2 capsid genes reveals mutations in the PL-2 epitope within the capsid’s spike domain. To understand the molecular basis for resistance from mAb PL-2 neutralization, we determined the 1.35 Å-resolution crystal structure of the capsid spike from one of these HAstV-2 strains. Our structure reveals a dramatic conformational change in a loop within the PL-2 epitope due to a serine-to-proline mutation, locking the loop in a conformation that sterically blocks binding and neutralization by mAb PL-2. We show that mutation to serine permits loop flexibility and recovers mAb PL-2 binding. Importantly, we find that HAstV-2 capsid spike containing a serine in this loop is immunogenic and elicits antibodies that neutralize all HAstV-2 strains. Taken together, our results have broad implications for rational selection of vaccine strains that do not contain prolines in antigenic loops, so as to elicit antibodies against diverse loop conformations.
IMPORTANCE

Human astroviruses (HAstVs) infect nearly every person in the world during childhood and cause diarrhea, vomiting, and fever. In this study, we investigated how several strains of HAstV are resistant to a virus-neutralizing monoclonal antibody. We determined the crystal structure of the capsid protein spike domain from one of these HAstV strains and found that a single amino acid mutation induces a structural change in a loop that is responsible for antibody binding. Our findings reveal how viruses can escape antibody neutralization and provide insight for the rational design of vaccines to elicit diverse antibodies that provide broader protection from infection.
INTRODUCTION

Astroviruses are a diverse family of small, non-enveloped, icosahedral positive-sense RNA viruses that infect a wide range of mammalian and avian species (1). In poultry, astroviruses have been associated with a large variety of disease manifestations, growth defects, and mortality (2). In humans, astroviruses are a leading cause of viral diarrhea in children, immunocompromised individuals, and the elderly (1, 3-5), accounting for 2 to 9% of all acute nonbacterial gastroenteritis in children worldwide (6). Human astroviruses (HAstVs) have also been associated with systemic infections and neurological complications such as encephalitis (7-9). There are three distinct phylogenetic clades of HAstV: canonical genotypes (HAstV-1-8) and noncanonical genotypes MLB1-3 and VA1-5 (6, 10, 11). HAstV-1 is the most prevalent serotype worldwide (3, 12), while prevalence of HAstV-2, the subject of the present study, can vary widely (13).

Several studies demonstrate that the adaptive immune response plays a key role in controlling astrovirus infection and disease. Approximately 75% of people in the United States have developed antibodies against HAstV by the age of ten (14), indicating that an adaptive immune response is mounted in humans. Clinical studies of healthy adult volunteers infected with HAstV revealed that those that had anti-HAstV antibodies experienced little or no disease, whereas those that did not have anti-HAstV antibodies experienced more severe diarrheal disease (15, 16). In another study, immunoglobulin therapy led to the recovery of an immunocompromised patient with a severe and persistent HAstV infection (17). More recently, astrovirus infectivity was tested in rag1 knockout mice (Rag1−/−),
which lack mature B cells or T cells. These studies show that Rag1−/− mice infected with murine astrovirus had higher (2 logs) levels of viral RNA compared to wild-type mice(18). Altogether, these studies support that adaptive immunity is key to the control of astrovirus infection. Furthermore, these studies suggest that a vaccine and therapeutic antibodies could be developed to prevent and/or treat HAstV infections.

Understanding the sites at which antibodies bind HAstV and understanding how the virus might resist antibody neutralization can inform the development of vaccines and antiviral therapeutics. Mature HAstV particles are comprised of a small RNA genome surrounded by a ~35nm T=3 icosahedral capsid protein shell projecting 30 knob-like spikes(19). Our lab and others have recently determined the crystal structure of the HAstV capsid core domain, which forms the icosahedral shell that encapsulates the viral RNA genome(20, 21). Our lab and others have also determined the crystal structure of the HAstV capsid spike domain, which forms the dimeric spike protrusions on the virus surface(20, 22, 23). While both the HAstV capsid core and spike domains are antigenic, the spike domain is ~5-fold more antigenic(20). In addition, the capsid spike fragment binds all four of the previously described HAstV-neutralizing monoclonal antibodies (mAbs), including mAb PL-2 that neutralizes serotype HAstV-2(24, 25).

We recently engineered the single chain variable fragment (scFv) of the PL-2 antibody and used X-ray crystallography to determine the structure of the HAstV capsid spike / scFv PL-2 complex(23, 26). We found that the HAstV-2 capsid spike’s loop 1 plays a major role in antibody PL-2 binding. We also provided evidence that
the HAstV capsid spike is a receptor-binding domain and that antibody PL-2 blocks
spike binding to human cells (23).

In the present study, we investigated the molecular mechanism by which
several strains of HAstV-2 are resistant to neutralization by mAb PL-2. We find that
a single point mutation in the capsid spike loop 1 of these HAstV-2 strains is
responsible for resistance to antibody neutralization. Structural studies reveal how
this mutation prevents antibody binding. Altogether, these studies have broad
implications for rational design of vaccines and therapeutic antibodies.

RESULTS

HAstV-2 strain resistance to antibody PL-2 neutralization. Monoclonal antibody
PL-2 (mAb PL-2) was reported to neutralize infectivity of HAstV-2 strain CDC-Spain
(HAstV-2-CDC-Spain), and our recent structural and functional studies suggested
that mAb PL-2 may neutralize HAstV-2 by blocking a receptor-binding site on the
capsid surface (23, 24). With the intention to directly investigate the mechanism of
mAb PL-2 neutralization, we began our studies by aiming to confirm that mAb PL-2
neutralizes HAstV-2 infection in cell culture. HAstV-2 strain Oxford (HAstV-2-
Oxford) was pre-incubated with increasing concentrations of mAb PL-2 or scFv PL-2
and then added to Caco-2 cells. To our surprise, no neutralization was observed by
either mAb PL-2 or scFv PL-2 (Fig. 1A,B). Similar results were observed with three
other HAstV-2 strains tested (HAstV-2-RIVMa, -RIVMb, and -RIVMc). In comparison,
complete neutralization of HAstV-2-Oxford was obtained with polyclonal antibodies
raised against HAstV-2-Oxford virus, and even partial neutralization was obtained
with polyclonal antibodies raised against HAstV-8-Yuc8 (Fig. 1C). Unfortunately, HAstV-2-CDC-Spain, which was originally used to produce mAb PL-2, was no longer available in the several labs that we queried.

**Sequence of HAstV-2 strains.** We used RT-PCR to determine the sequence of the capsid spike from each of the four HAstV-2 strains (strains Oxford, RIVMa, RIVMb, and RIVMc). We then aligned these sequences to the reported capsid protein sequence of HAstV-2-CDC-Spain (Fig. 1E). Focusing on the amino acids within the mAb PL-2 epitope, we observed two amino acids that are mutated in all four HAstV-2 strains: Ser463Pro and Glu580Lys. Ser463 falls in the middle of the PL-2 epitope in loop 1 of the capsid spike domain whereas Glu580 resides at the edge of the epitope(23).

**Structure of Spike-2-Oxford.** To understand the molecular basis for resistance to mAb PL-2 neutralization, we expressed and purified recombinant HAstV-2-Oxford capsid spike domain (Spike-2-Oxford) and observed that it elutes as a dimer by size exclusion chromatography, consistent with previous observations for recombinant HAstV-2 capsid spike domain (Spike-2-CDC-Spain)(Fig. 2A)(23). We used an enzyme-linked immunosorbent assay (ELISA) to test if mAb PL-2 binds Spike-2-Oxford. Consistent with the lack of mAb PL-2 neutralization of HAstV-2-Oxford infectivity, we observed no binding of mAb PL-2 to Spike-2-Oxford, whereas dose-dependent binding was observed for Spike-2-CDC-Spain (Fig. 2B). We then crystallized Spike-2-Oxford and solved its structure to 1.35 Å resolution (Fig. 2C,
Table 1). The overall structure of Spike-2-Oxford is very similar to that of Spike-2-
CDC-Spain, with an RMSD of 0.315 Å.

The major structural difference between Spike-2-Oxford and Spike-2-CDC-
Spain occurs in loop 1, which interacts with antibody PL-2 (Fig. 3). We observe that
the Ser463Pro mutation in loop 1 leads to the formation of a short alpha helix,
locking loop 1 into a distinct conformation. Loop 1 adopts this same “down”
conformation in all four molecules of Spike-2-Oxford in the crystallographic
asymmetric unit (Fig. 3A). In contrast, loop 1 in Spike-2-CDC-Spain appears more
flexible and adopts different conformations in each of the four molecules in the
crystallographic asymmetric unit, with two molecules in a “down” conformation,
one molecule in an intermediate conformation, and one molecule in an “up”
conformation (Fig. 3B). Thus, it appears that the flexibility of loop 1 in Spike-2-CDC-
Spain may be required for binding of antibody PL-2. Indeed, we observed that loop 1
adopts a single “up” conformation in all eight molecules of Spike-2-CDC-Spain/scFv
PL-2 complex in the crystallographic asymmetric unit (Fig. 3C). Superposition of the
Spike-2-Oxford structure onto the structure of the Spike-2-CDC-Spain/scFv PL-2
complex reveals how the “down” conformation of loop 1 in Spike-2-Oxford would
sterically clash with antibody heavy chain CDR 3 (Fig. 4). Overall, our structural
studies suggest that the Ser463Pro mutation locks loop 1 into a rigid conformation
that clashes with and completely prevents antibody PL-2 binding.

**Spike-2-Oxford mutant Pro463Ser restores binding to mAb PL-2.** Our structural
studies lead to the hypothesis that HAstV-2-Oxford capsid Pro463 is responsible for
the lack of binding and neutralization by mAb PL-2. To test our hypothesis, we expressed and purified recombinant Spike-2-Oxford Pro463Ser mutant and observed that it is pure and elutes as a dimer by size exclusion chromatography (Fig. 5A and 5B). We then performed an ELISA using purified mAb PL-2 and scFv PL-2 (Fig. 5B-D). We observed that both antibody PL-2 samples now bind to Spike-2-Oxford Pro463Ser mutant at similar levels as binding to Spike-2-CDC-Spain. These results confirm that a single amino acid in the capsid protein of HAstV-2-Oxford is responsible for resistance to binding and neutralization by mAb PL-2.

Spike-2-CDC-Spain elicits HAstV-2 cross-neutralizing antibodies. We hypothesized that Spike-2-CDC-Spain, which contains Ser463 in loop 1, would elicit antibodies against diverse conformations of loop 1 that can neutralize the infectivity of all four HAstV-2 strains. To test this, we immunized rabbits with purified, recombinant Spike-2-CDC-Spain and found that the anti-Spike-2-CDC-Spain polyclonal antibodies effectively neutralized all four strains of HAstV-2 in a concentration-dependent manner (Fig. 1D). Although the spike most likely contains more than one neutralizing epitope, this study does show that the Spike-2-CDC-Spain, which contains Ser463, is able to elicit antibodies that neutralize HAstV-2 strains containing Pro463. Moreover, this is the first demonstration that recombinant HAstV capsid spike can elicit HAstV-neutralizing antibodies and may be an effective immunogen in a HAstV subunit vaccine.

DISCUSSION
Here, we investigated the molecular mechanism by which four strains of HAstV-2 (HAstV-2-Oxford, -RIVMa, -RIVMb, and -RIVMc) are resistant to the HAstV-2-CDC-Spain-neutralizing monoclonal antibody PL-2. Serotype HAstV-2 has been reported to have especially high genetic heterogeneity between strains (13). We describe the crystal structure of Spike-2-Oxford and find that Pro463 induces a rigid helix in loop 1, locking it in a “down” conformation that sterically clashes and prevents binding by mAb PL-2. We further show that the point mutant Pro463Ser recovers binding by mAb PL-2. Finally, we show that recombinant Spike-2-CDC-Spain is immunogenic and elicits antibodies that neutralize all four HAstV-2 strains.

Viruses constantly evolve under the pressure of a host’s immune system, and one of the most efficient ways to evade immunity is by acquiring mutations that sterically clash with antibody binding. While serines and prolines are often interchangeable amino acids due to their similarity in size, we find that mutation of serine to proline in a protein loop can induce a major conformational change. Proline does not follow the typical Ramachandran plot, due to its cyclic side chain that limits motion in its $\psi$ and $\phi$ angles, reducing flexibility to the polypeptide backbone. Because loop structures are often associated with antigenic epitopes, it is possible that viruses mutate amino acids in antigenic loops to prolines to evade binding by some antibodies. Indeed, serine codons, as well as threonine, alanine, leucine, histidine, glutamine, and arginine codons, are most susceptible of becoming a proline codon, since in these cases it requires only a single nucleotide mutation.
One might initially conclude from these studies that vaccines strain antigens with serines in antigenic loops should not be chosen because they are at risk for antibody escape, however we would actually argue the opposite. Specifically, we propose that antigens with serines in antigenic loops should be chosen for vaccine strains because they would elicit more diverse polyclonal antibodies against the diverse conformations in the loop, including conformations observed upon mutation to proline. Indeed, our structures reveal that, with a serine, the loop will sometimes adopt the conformation that is restrained by a proline mutation. In support of this concept, we show that recombinant Spike-2-CDC-Spain, which has a serine in loop 1, elicits antibodies that neutralize all four HAstV-2 strains with prolines in loop 1.

Notably, we show that recombinant HAstV capsid spike is immunogenic and elicits neutralizing antibodies. These findings demonstrate the high potential to develop a subunit vaccine to prevent HAstV diarrheal disease. In addition, the affordable, simple, and scalable strategy of producing large amounts of recombinant HAstV capsid spike in *E. coli* hints at the feasibility of global vaccination. Of course, future studies will be required to determine if a vaccine comprised of capsid spikes from multiple serotypes is required to elicit a broadly protective polyclonal antibody response.

**METHODS**

**Cells, viruses, and reagents.** C2Bbe1 cells (ATCC), derived from the human colon adenocarcinoma Caco-2 cell line, were propagated in a 10% CO₂ atmosphere at 37°C in Dulbecco’s modified Eagle’s medium-High Glucose (DMEM-HG) (Sigma),
supplemented with non-essential amino acids (Gibco) and 15% fetal bovine serum (FBS) (Cansera). HAstV-2 strain Oxford (HAstV-2-Oxford) was obtained from J.B. Kurtz (Dept. of Virology, John Radcliffe Hospital, Oxford, UK). HAstV-2 strains RIVMa, RIVMb, and RIVMc were obtained from S. Guix (Dept. Microbiologia, Facultat de Biologia, Universitat de Barcelona). All viral strains were activated and grown as described (27), except that 200 ug/ml of trypsin was used to active the virus infectivity. Anti-Spike-2-CDC-Spain polyclonal sera was generated by immunization of mice with recombinant HAstV-2-CDC-Spain capsid spike (see below). For this, female BALB/c mice (8 weeks old) were immunized intraperitoneally with 50 ug of recombinant HAstV capsid spike at 2-week intervals (four times total). The first immunization was done in Freund’s complete adjuvant, the second and third with incomplete Freund’s adjuvant, and the fourth with no adjuvant. Four days after the fourth immunization the mice were bled to death. Anti-Core-1 polyclonal sera used to detect HAstV infectivity in C2Bbe1 cells was generated by immunization of New Zealand rabbits with recombinant HAstV-1 capsid core (amino acids 80-429) (20). The generation of polyclonal antibodies to Spike-2-CDC-Spain and to Core-1 was approved by the Bioethics Committee of the Institute of Biotechnology UNAM (#296).

Neutralization assays. The indicated concentration of antibody or scFv was pre-incubated with an m.o.i. of 0.002 of the indicated HAstV-2 strain for 1 h at room temperature. The virus-antibody mix was then added to confluent monolayers C2Bbe1 cells grown in 96 multi-well plates, and incubated for 1 h at 37°C. After this
time the cells were washed three times with PBS and the infection was left to proceed for 18 h at 37°C. Infected cells were detected by an immunoperoxidase focus-forming assay, as described (28), with the following modifications. At 18 h post infection the cells were fixed for 20 min at room temperature with 2% formaldehyde, washed with PBS, and permeabilized using 0.2% Triton X-100 (in PBS) for 15 min, at room temperature. For detection of infected cells, a polyclonal serum directed to HAstV-1 capsid core was used (see above). Experiments were performed in biological triplicates.

**PCR and sequencing.** For HAstV-2 sequencing, RNA was isolated from viral lysates using PureLink® viral RNA/DNA Mini Kit (Invitrogen), and cDNA was synthesized with SuperScript™ III reverse transcriptase (Thermo Fisher Scientific) using as primer the sequence 5’-GCGGTCTCCAGAAAGTTTG-3’ (HastV2LW) corresponding to the nucleotide position 2369 to 2387 of the HAstV-2 capsid gene (accession number 8497068). For PCR amplification, Vent® DNA Polymerase (New England BioLabs) and the oligonucleotides HAstV2LW and HAstV2Up 5’-CAGTTCACTCAAATGAACCA-3’, corresponding to nucleotides 1215 to 1234 of the HAstV-2 capsid gene (accession number L06802.1), were used. The PCR product was purified using the DNA clean and Concentrator-5 kit (Zymo Research) and sequenced in the sequencing facility of the Instituto de Biotecnología, UNAM. The HAstV-2-Oxford sequence was deposited in GenBank (accession number KY964327).
Expression and purification of Spike-2. Synthetic genes codon-optimized for *Escherichia coli* encoding HAstV-2-CDC-Spain capsid spike amino acids 431 to 674 (GenBank accession AAA62427.1), and HAstV-2-Oxford capsid spike (Accession KY964327) were purchased. To make spike expression plasmids, genes were cloned into pET52b (Addgene) in-frame with a C-terminal thrombin cleavage site and a 10-histidine purification tag. To make Spike-2-Oxford Pro463Ser mutant expression plasmid, the Phusion Site-Direct Mutagenesis Kit (Thermo Scientific) was used with phosphorylated mutagenesis primers. All plasmids were verified by DNA sequencing. Plasmids were transformed into *E. coli* strain BL21(DE3), and protein production was induced with 1mM isopropyl-β-D-thiogalactopyranoside at 18°C for 16 h. *E. coli* cells were lysed by ultrasonication in 20 mM Tris-HCl, pH 8.0, 500 mM NaCl, and 20 mM imidazole (Buffer A) containing 2 µM MgCl$_2$, 1250 U benzonase (Millipore), and 1X protease inhibitor cocktail Set V EDTA-Free (Millipore). Proteins were batch purified from soluble lysates by TALON metal affinity chromatography and eluted with Buffer A containing 500 mM imidazole. Proteins were dialyzed into 20 mM Tris-HCl, pH 8.0 and 25 mM NaCl and purified by anion exchange chromatography on a HiTrap Q FF column with a gradient elution to 20 mM Tris-HCl, pH 8.0 and 1 M NaCl. Proteins were buffer exchanged into PBS and further purified by size-exclusion chromatography on a Superdex 200 column in PBS.

**ELISA.** Purified spike proteins at a concentration of 5 µg/mL in PBS (150 µL total) were incubated overnight at room temperature in 96-well ELISA microtiter plates. Plates were then washed three times with PBS containing 0.05% Tween 20 (PBST).
Wells were blocked by adding 150 µL of 5% BSA in PBS and incubating at room temperature for 1 hr followed by three PBST washes. Antibody samples mAb PL-2 or scFv PL-2 were diluted to 5 µg/mL with 1% BSA in PBS, and serially diluted 1:3 with 1% BSA in PBS. Wells were incubated with 150 µL antibody for 1 hr at room temperature and the plates were washed three times with PBST. For ELISAs where the primary antibody was mAb PL-2, plates were incubated for 1 hr at room temperature with 150 µL HRP-conjugated, secondary goat anti-mouse IgG antibody diluted 1:5,000 in 1% BSA in PBS. For ELISAs where the primary antibody was scFv PL-2, plates were incubated for 1 hr at room temperature with 150 µL HRP-conjugated Strep-Tactin protein, diluted 1:5,000 in 1% BSA in PBS. Plates were washed three times with PBST and developed by adding peroxidase substrate o-phenylenediamine dihydrochloride (OPD) in 0.05 M phosphate-citrate buffer, pH 5.0 and 1.5% hydrogen peroxide for 10 min at room temperature. The reactions were stopped by incubation with 2N sulfuric acid for 10 min at room temperature, and the absorbance was measured at 490 nm.

Structure determination of Spike-2-Oxford. Purified Spike-2-Oxford in PBS was concentrated to 28.3 mg/mL. Crystals were grown by hanging drop vapor diffusion at 22°C with a well solution of 20% PEG 3350, 0.2 M magnesium acetate, and 0.1 M HEPES buffer, pH 7.5. Crystals were transferred into a cryoprotectant solution of 25% PEG 3350, 25% glycerol, 0.2M magnesium acetate, 0.1M HEPES, pH 7.5, and flash frozen in liquid nitrogen. Diffraction data were collected at cryogenic temperature at the Advanced Photon Source on beamline 23-ID-B using a
wavelength of 1.033 Å. Diffraction data from a single crystal were processed with
iMosflm(29) and Scala(30)(Table 1). The Spike-2-Oxford structure was solved by
molecular replacement, using the HAstV-2-CDC-Spain capsid spike (PDB ID
5KOU)(22) and the program PHASER(31). The Spike-2-Oxford structure was refined
and manually rebuilt using PHENIX(32) and Coot(33), respectively. The final Spike-
2-Oxford structure had two dimers in the asymmetric unit of the crystal.

**Accession code.** Coordinates and structure factors have been deposited in the
Protein Data Bank under accession code 5W1N.

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FIG 1 Antibody neutralization of HAstV-2 strains and alignment of HAstV-2 capsid spike sequences. (A) Infectivity of indicated HAstV-2 strains pre-incubated with mAb PL-2. (B) Infectivity of indicated HAstV-2 strains pre-incubated with scFv PL-2. (C) Infectivity of HAstV-2-Oxford pre-incubated with anti-HAstV-2-Oxford or anti-HAstV-2-Yuc8 polyclonal sera. (D) Infectivity of indicated HAstV-2 strains pre-
incubated with anti-Spike-2-CDC-Spain polyclonal sera. All infectivity experiments were performed in biological triplicates and the error bars represent the standard error of the mean. (E) Sequence alignment of Spike-2-CDC-Spain, Spike-2-Oxford, and Spike-2-RIVMa-c. Conserved, strongly similar, weakly similar, and non-conserved amino acids are colored red, dark pink, light pink, and white, respectively. Alignments and mapping of conservation onto the structure was performed with the online ENDscript server. Black boxes highlight amino acids in the antibody PL-2 epitope. A blue star indicates the location of Ser463 or Pro463.

**FIG 2 Purification, antibody binding, and structure of Spike-2-Oxford.** (A) Superdex 200 16/600 size-exclusion chromatography trace of Spike-2-CDC-Spain. (B) ELISA results showing specific binding of mAb PL-2 to Spike-2-CDC-Spain and no binding to Spike-2-Oxford. The yellow “Control” sample refers to ELISA wells coated with PBS buffer (no Spike protein). (C) Crystal structure of Spike-2-Oxford, with half of the dimer in gray and the other half in light blue. The gray half has labeled β-sheets, α-helices and loops.
FIG 3 Structural rigidity of loop 1 in Spike-2-Oxford. (A) Structural alignment of all four molecules of Spike-2-Oxford in the crystallographic asymmetric unit. (B) Structural alignment of all four molecules of Spike-2-CDC-Spain in the crystallographic asymmetric unit. (C) Structural alignment of all eight molecules of Spike-2-CDC-Spain in the crystallographic asymmetric unit from the Spike-2-CDC-Spain / scFv PL-2 complex structure.
**FIG 4** Spike-2-Oxford loop 1 sterically clashes with scFv PL-2 binding. The structure of Spike-2-Oxford (cyan) was superimposed onto the structure of Spike-2-CDC-Spain (red) bound to the scFv PL-2 (with the heavy chain colored blue and the light chain colored purple). **Top right:** zoom in showing Spike-2-CDC-Spain Ser463 and “up” conformation of loop 1. **Bottom right:** zoom in showing Spike-2-Oxford Pro463 inducing a helix and “down” conformation in loop 1, which clashes with antibody heavy chain (blue) CDR H3.

**FIG 5** Purification and antibody binding of Spike-2-Oxford Pro463Ser mutant. (A) Superdex 200 16/600 size-exclusion chromatography trace of Spike-2-Oxford Pro463Ser mutant. (B) Coomassie-stained non-reducing SDS-PAGE of mAb PL-2 and scFv PL-2 (left). Coomassie-stained reducing SDS-PAGE of Spike-2-CDC-Spain, Spike-2-Oxford, and Spike-2-Oxford P463S mutant. (C) ELISA results showing specific binding of mAb PL-2 to Spike-2-Oxford Pro463Ser mutant. (D) ELISA results showing specific binding of scFv PL-2 Spike-2-Oxford Pro463Ser mutant.
Table 1 Data collection and refinement statistics\(^a\)

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<tr>
<td>Favored (%)</td>
<td>97.7</td>
</tr>
<tr>
<td>Allowed (%)</td>
<td>2.3</td>
</tr>
<tr>
<td>Outliers (%)</td>
<td>0</td>
</tr>
</tbody>
</table>

\(^a\)Data from one crystal was used for structure determination. Values in parentheses are for the highest-resolution shell.