Loss-of-function mutation in Omicron variants reduces spike protein expression and attenuates SARS-CoV-2 infection

- 3 Michelle N. Vu¹, Dorothea R. Morris^{1, 2}, R. Elias Alvarado^{1,3}, Kumari G. Lokugamage¹, Yiyang
- 4 Zhou⁴, Leah K. Estes¹, Alyssa M. McLeland¹, Craig Schindewolf¹, Jessica A. Plante^{1, 5, 6}, Yani P.
- 5 Ahearn¹, Angelica L. Morgan⁷, William M. Meyers⁷, Jordan T. Murray¹, Scott C. Weaver^{1, 5, 6, 8},
- 6 David H. Walker^{7, 8}, William K. Russell⁴, Andrew L. Routh⁴, Kenneth S. Plante^{1, 5, 6}, Vineet
- 7 Menachery^{1, 5, 6, 8*}
- ⁸ ¹Microbiology and Immunology, University of Texas Medical Branch, Galveston, TX, United
- 9 States.
- ²Pediatrics, University of Texas Medical Branch, Galveston, TX, United States.
- ³Institute for Translational Sciences, University of Texas Medical Branch, Galveston, TX, United
 States.
- ⁴Biochemistry and Molecular Biology, University of Texas Medical Branch, Galveston, TX,
- 14 United States.
- ⁵Institute for Human Infection and Immunity, University of Texas Medical Branch, Galveston, TX,
- 16 United States.
- ¹⁷ ⁶World Reference Center of Emerging Viruses and Arboviruses, University of Texas Medical
- 18 Branch, Galveston, TX, United States.
- ¹⁹ ⁷Pathology, University of Texas Medical Branch, Galveston, TX, United States.
- ⁸Center for Biodefense and Emerging Infectious Disease, University of Texas Medical Branch,
- 21 Galveston, TX, United States.
- 22
- 23 * Corresponding Author: Vineet D. Menachery, vimenach@utmb.edu

24 Abstract

SARS-CoV-2 Omicron variants emerged in 2022 with >30 novel amino acid mutations in the spike 25 protein alone. While most studies focus on the impact of receptor binding domain changes, 26 mutations in the C-terminal of S1 (CTS1), adjacent to the furin cleavage site, have largely been 27 28 ignored. In this study, we examined three Omicron mutations in CTS1: H655Y, N679K, and 29 P681H. Generating a SARS-CoV-2 triple mutant (YKH), we found that the mutant increased spike processing, consistent with prior reports for H655Y and P681H individually. Next, we generated 30 a single N679K mutant, finding reduced viral replication in vitro and less disease in vivo. 31 32 Mechanistically, the N679K mutant had reduced spike protein in purified virions compared to wild-33 type; spike protein decreases were further exacerbated in infected cell lysates. Importantly, 34 exogenous spike expression also revealed that N679K reduced overall spike protein yield independent of infection. Together, the data show that N679K is a loss-of-function mutation 35 reducing overall spike levels during omicron infection, which may have important implications for 36 37 disease severity, immunity, and vaccine efficacy.

38

39 One Sentence Summary

40 Spike substitution N679K attenuates SARS-CoV-2 Omicron variants by decreasing spike protein

and has potential implications for immunity and vaccine efficacy.

42 Introduction

43 Since its introduction, SARS-CoV-2 has continuously evolved giving rise to multiple Variants of Concern (VOCs) with diverse mutations in the spike protein (1). Present as a trimer 44 on virions, spike is composed of S1 and S2 subunits, responsible for receptor binding and 45 46 membrane fusion, respectively (2, 3). The S1 subunit contains the N-terminal domain (NTD), 47 receptor binding domain (RBD), and the C-terminus of the S1 subunit (CTS1), which harbors a furin cleavage site (FCS) in SARS-CoV-2. Following receptor binding, the spike is cleaved at the 48 S1/S2 site by host proteases to expose the fusion machinery for entry. With the diverse mutations 49 50 in the spike protein, most Omicron studies have focused on the RBD and the impact on vaccine-51 or infection-induced immunity. However, mutations surrounding the FCS and S1/S2 cleavage site have been demonstrated to drive SARS-CoV-2 pathogenesis (4-9) and have been largely 52 unstudied in the context of Omicron. 53

54 With this in mind, we set out to evaluate the role of Omicron CTS1 mutations on infection 55 and pathogenesis. Omicron maintains three mutations adjacent to the FCS and S1/S2 cleavage site: H655Y, N679K, and P681H. Both H655Y and P681H have previously been observed in the 56 Gamma and Alpha variants (10, 11); in contrast, N679K is unique to and maintained by all 57 58 Omicron subvariants (12). To evaluate the role of these mutations, we used reverse genetics to generate SARS-CoV-2 mutants with all three CTS1 mutations (YKH) or N679K alone in the 59 original WA1 backbone from early 2020. While YKH modestly increases viral replication and spike 60 processing in vitro, N679K results in a loss-of-function mutation that attenuates viral replication in 61 62 vitro and disease in vivo through reduced spike protein expression. Given the importance of spike protein for immunity, our finding may have major implications for vaccine efficacy and 63 breakthrough infections. 64

65

66 Results

67 H655Y, N679K, and P681H together increase viral replication and spike processing.

While the majority of the > 30 spike mutations Omicron acquired are localized to the RBD. 68 three are harbored in the CTS1 adjacent to the furin cleavage site - H655Y, N679K, and P681H 69 70 (Fig. 1A). Both H655Y and P681H have been observed individually in Gamma and Alpha variants 71 and are associated with increased spike processing. In contrast, N679K is a mutation unique to Omicron and is maintained in all subsequent Omicron subvariants despite involving a single 72 73 nucleotide change (T/C to A/G) in the wobble codon position (12). Importantly, N679K is adjacent 74 to an important O-linked glycosylation site at T678 (13, 14); our group has previously shown this 75 glycosylation is important for SARS-CoV-2 infection and protease usage (8).

76 Several motifs within the CTS1 spike domain, including the furin cleavage site and the upstream QTQTN motif, are key to spike cleavage and host protease interactions, which drive 77 SARS-CoV-2 infection and pathogenesis. All three Omicron mutations in the CTS1, H655Y, 78 79 N679K, and P681H, are adjacent to or within these motifs and may impact their function (Fig. 1A and 1B). To evaluate this, we generated a mutant SARS-CoV-2 harboring H655Y, N679K, and 80 P681H in the original WA1 backbone (YKH) (Fig. 1C) (15, 16). Plagues produced by the YKH 81 82 mutant were smaller compared to the parental WA1 (WT) (Fig. 1D). However, the YKH mutant did not attenuate stock titers nor replication kinetics in Vero E6 cells as compared to wild-type 83 (WT) SARS-CoV-2 (Fig. 1E and 1F). Notably, while replication was slightly reduced at 24 hpi, 84 85 end point titers for YKH were augmented at 48 hpi in Calu-3 2B4 cells compared to WT (Fig. 1G). 86 The results suggest that the combination of the three mutations alters infection dynamics, which 87 may offer some advantages to the Omicron variant in human respiratory cells (Fig. 1G). As H655Y and P681H have individually been shown to increase spike processing, we next evaluated spike 88 processing on purified virions from YKH and WT infection. Similar to Delta and Omicron, YKH 89 90 spike was more processed than WT (Fig. 1H and 1I). At 24 hpi, the S1/S2 cleavage ratio to full length spike ratio was ~2.4:1 for the YKH spike (55% S1/S2 product, 23% full-length); in contrast, 91

92 WT had roughly equivalent amounts of S1/S2 product and full length. Overall, the combination of

93 H655Y, N679K, and P681H in the YKH mutant resulted in increased viral endpoint yields in human

⁹⁴ respiratory cells and contributed to Omicron's enhanced spike processing.

95 N679K mutation attenuates SARS-CoV-2 infection.

96 The increase in spike processing found in the YKH mutant is consistent with prior work examining H655Y and P681H mutations individually; however, the contribution of N679K had yet 97 to be evaluated. Based on its location adjacent to a key O-linked glycosylation site, we 98 hypothesized that N679K might impact SARS-CoV-2 infection (Fig. 2A). To evaluate potential 99 100 changes, we generated a SARS-CoV-2 mutant with only N679K in the WA1 backbone (N679K) 101 (Fig. 2B). Our initial characterization found that the N679K plague sizes were distinctly smaller at days 2 and 3 post-infection (Fig. 2C), and stock titers were slightly lower than WT (Fig. 2D). 102 103 These differences in plaque size and stock titers are consistent with other observations of most 104 of the Omicron strains (17-20). Notably, unlike the minimal differences seen in YKH replication 105 kinetics, the N679K mutant had attenuated replication in both Vero E6 and Calu-3 2B4 cells at 24 hpi (Fig. 2E and 2F). Although N679K viral titer recovered by 48 hpi, the results suggest that 106 N679K is a loss-of-function mutation in terms of replication in both cell lines. 107

108 We next evaluated N679K in vivo by infecting 3-to-4-week-old golden Syrian hamsters and monitored weight loss and disease over 7 days (Fig. 2G). Hamsters infected with N679K 109 displayed significantly attenuated body weight loss compared to those infected with WT (Fig. 2H). 110 111 Despite the stark attenuation seen in weight loss, N679K viral titers in the lungs were equivalent 112 to WT at 2 dpi and 4 dpi (Fig. 2I). Similarly, N679K viral titers were comparable to WT at 2 dpi in 113 nasal washes; however, the mutant virus resulted in reduced replication at 4 dpi (Fig. 2J). Taken together, our results indicate that N679K has a distinct loss-of-function phenotype in vitro and in 114 vivo; given this result, we hypothesize that N679K loss-of-function is mitigated by the impact of 115 116 the H655Y and P681H mutations.

117 N679K mutation results in decreased spike protein expression.

We next sought to determine the mechanism driving the loss-of-function observed with 118 119 the N679K mutant. Given its location adjacent to the FCS, we first evaluated N679K effects on proteolytic spike processing. Virions were purified from WT, N679K or the Omicron variant BA.1 120 (Omicron) and blotted for spike processing. Nearly identical to YKH, the N679K mutant had 121 122 increased spike processing with a ~2.5:1 ratio of S1/S2 cleavage product to full-length spike compared to 1:1 ratio for WT at 24 hpi (Fig. 3A and 3B). However, we noted distinct differences 123 in total spike protein with N679K and Omicron compared to WT, despite similar levels of 124 125 nucleocapsid protein. Densitometry analysis revealed that the total spike to nucleocapsid (S/N) 126 ratio of N679K and Omicron virions was reduced 21% and 36%, respectively, as compared to WT 127 (Fig. 3C). Overall, our results indicate that the N679K mutant and Omicron variant incorporate less spike protein onto their virions. 128

We then sought to determine if changes in the virion spike were due to changes to total 129 protein expression in the cell or incorporation into the particle. To examine spike protein 130 expression, we measured total spike relative to nucleocapsid in infected Vero E6 cell lysates 24 131 hpi (Fig. 3D and 3E). N679K resulted in a S/N ratio 66% less than WT, displaying an even further 132 decrease in spike protein compared to the reduction in purified virions. Additionally, a similar 133 134 decrease in S/N ratio was observed in Omicron, indicating that the phenotype is maintained in the context of all Omicron mutations (Fig. 3D and 3E). Together, the results indicate that the 135 N679K mutation reduces the Omicron spike protein levels compared to WT. 136

Having established reduced spike protein in the context of N679K, we next wanted to determine if this reduction only occurs in the context of virus infection or is inherent to the protein. Therefore, we introduced the mutation into the Spike HexaPro plasmid to exogenously express spike protein to separate N679K driven changes from other aspects of viral infection (*21*). Vero E6 cells were transfected with the WT or N679K mutant spike HexaPro and harvested at 24 and 48 hours post transfection (hpt). Similar to what was observed in viral infection, N679K spike was reduced 43% at 24 hpt and 46% at 48 hpt (**Fig. 3F and 3G**). Overall, the results suggest that the reduction in Omicron spike is governed by N679K in a manner independent of viral infection.

145 Discussion

Most Omicron studies have focused on determining the impact that the RBD mutations 146 147 have on immune escape, largely overlooking mutations in other spike domains like the CTS1. Harboring the FCS and S1/S2 cleavage site, the CTS1 has been demonstrated as a hotspot for 148 both attenuating and augmenting mutations (4-9). Focusing on Omicron's three CTS1 mutations 149 150 - H655Y, N679K, and P681H, we generated infectious clones with all three (YKH) or N679K alone 151 in the SARS-CoV-2 WA1 background. The combination of YKH produced a modest increase in 152 endpoint titers after infection of human respiratory cells, and augmented spike processing, consistent with prior studies that tested the effects of H655Y and P681H individually (10, 11). 153 154 However, the N679K mutant reduced viral replication in vitro and weight loss in vivo. Mechanistic 155 studies determined that both N679K and Omicron have reduced spike protein incorporated into their virions, less spike protein in infected cell lysates, and inferior production using exogenous 156 spike protein expression systems compared to WT. Our results argue that reduced spike protein 157 158 in the context the N679K mutation attenuates Omicron strains and may have implications for 159 SARS-CoV-2 immunity by reducing spike antigen thus shifting immune recognition.

160 N679K is likely attenuated because of its decrease in spike protein production. Starting with ~30% less spike in its virions, one possibility was a change in spike incorporation. However, 161 an even greater decrease (66%) in spike protein was present in infected Vero E6 cell lysates, 162 163 indicated that overall spike protein levels were affected. To confirm that the reduction in spike was not a product of virus infection or host immune interactions, we exogenously expressed the 164 165 spike protein to demonstrate that the N679K spike protein itself was less stable than the WT 166 control. N679K, a single nucleotide and amino acid change, is unlikely to alter spike RNA 167 expression and may instead act at the protein level. One possible mechanism is that the 168 asparagine-to-lysine change introduces a ubiguitination site that could lead to spike degradation.

Another possible mechanism is that the N679K mutation itself may destabilize the protein structurally. Additionally, the N679K substitution adds another basic amino acid to the stretch including the FCS; the additional positively charged lysine extends the polybasic cleavage motif and may facilitate cleavage by additional host proteases (*22*). Overall, while the exact mechanism is unclear, the N679K mutation results in a less stable spike protein that impacts infection and pathogenesis of SARS-CoV-2.

Surprisingly, N679K is uniformly found in 100% of Omicron sequences in GSAID, despite 175 176 being a single nucleotide change in the wobble position (12). Though attenuated in vitro, N679K 177 does replicate to similar titers as WT in hamster lungs and at day 2 in nasal washes. These results 178 suggest no deficits in transmission and may explain how N679K is maintained despite clear 179 attenuation of SARS-CoV-2 infection. Notably, addition of H655Y and P681H in the YKH mutant rescues replication in Calu-3 cells, suggesting that other Omicron mutations may compensate for 180 181 N679K. However, it is unclear if reverting N679K in the Omicron strains would result in a gain-offunction in terms of in vitro replication or in vivo pathogenesis. While N679K in SARS-CoV-2 WA1 182 produces a clear loss-of-function, the constellation of spike mutations and epistatic interactions 183 may mitigate the deficit in Omicron strains. Importantly, the complete conservation of N679K in 184 185 Omicron also implies some fitness advantage; this may be in transmission, replication in certain cells/tissues, or evasion of host immune responses (23-28). While more studies are required to 186 decipher why it is maintained, N679K is clearly a mutation that impacts overall spike protein 187 188 expression.

In addition to impacting primary infection, the reduction in spike protein may have important implications for SARS-CoV-2 resistance to human immunity. Compared to WT, the N679K mutant produces less spike protein upon infection and can potentially skew the ratio of antibodies targeting spike and nucleocapsid. Prior work with SARS-CoV had shown that an altered spike/nucleocapsid antibody ratio contributed to vaccine failure in aged mice (*29*). Therefore, infection with Omicron could increase N targeting antibodies at the expense of spike

195 antibodies. The result would be less protective neutralizing antibody, which may facilitate more 196 breakthrough infections. Furthermore, SARS-CoV-2 vaccines based on the Omicron spike may produce less spike protein due to N679K mutation. In the context of the mRNA bivalent vaccines, 197 the N679K mutation may alter the 1:1 ratio of WT to Omicron spike protein; N679K may bias 198 199 immune responses towards WT spike protein instead of equally between both spike proteins. In 200 addition, the total amount of spike protein produced may be less than previous vaccines formulations, thus diminishing the overall antibody response. These factors potentially contribute 201 202 to the less than expected increase in immunity against Omicron despite the new bivalent vaccine 203 formulations. Moving forward, reverting K679 back to N679 in vaccine may improve spike protein 204 yields and subsequently improve vaccine response to the Omicron variants.

Together, our results demonstrate that Omicron N679K is a loss-of-function mutation 205 consistently maintained in subvariants. Mechanistically, the N679K mutation attenuates the virus 206 207 in vitro and in vivo by increasing spike degradation. While the N679K mutation is attenuating in isolation, other Omicron mutations like H655Y and P681H may compensate for the N679K loss 208 of function by amplifying spike processing and infection. However, the decreased spike protein 209 expression by N679K may have implications for immunity induced by infection and vaccines. 210 211 Overall, the interplay between the Omicron CTS1 mutations has a significant impact on SARS-212 CoV-2 infection and require additional studies for clarification.

213 References

- 1. E. Callaway, Heavily mutated Omicron variant puts scientists on alert. *Nature* **600**, 21 (2021).
- S. Belouzard, J. K. Millet, B. N. Licitra, G. R. Whittaker, Mechanisms of Coronavirus Cell Entry Mediated by the Viral Spike Protein. *Viruses* 4, 1011-1033 (2012).
- M. N. Vu, V. D. Menachery, Binding and entering: COVID finds a new home. *PLOS Pathogens* 17, e1009857 (2021).
- B. Coutard *et al.*, The spike glycoprotein of the new coronavirus 2019-nCoV contains a furin-like
 cleavage site absent in CoV of the same clade. *Antiviral Research* 176, 104742 (2020).
- 5. B. A. Johnson *et al.*, Loss of furin cleavage site attenuates SARS-CoV-2 pathogenesis. *Nature*, (2021).
- Y. Liu *et al.*, Delta spike P681R mutation enhances SARS-CoV-2 fitness over Alpha variant. *bioRxiv*,
 2021.2008.2012.456173 (2021).
- 7. T. P. Peacock *et al.*, The furin cleavage site in the SARS-CoV-2 spike protein is required for
 transmission in ferrets. *Nat Microbiol* 6, 899-909 (2021).
- 8. M. N. Vu *et al.*, QTQTN motif upstream of the furin-cleavage site plays a key role in SARS-CoV-2
 infection and pathogenesis. *Proceedings of the National Academy of Sciences* **119**, e2205690119
 (2022).
- S.-Y. Lau *et al.*, Attenuated SARS-CoV-2 variants with deletions at the S1/S2 junction. *Emerging Microbes & Infections* 9, 837-842 (2020).
- 23210.A. Escalera *et al.*, Mutations in SARS-CoV-2 variants of concern link to increased spike cleavage233and virus transmission. *Cell Host & Microbe* **30**, 373-387.e377 (2022).
- 23411.Y. Liu *et al.*, Delta spike P681R mutation enhances SARS-CoV-2 fitness over Alpha variant. *Cell*235*Reports* **39**, (2022).
- 12. J. Hadfield *et al.*, Nextstrain: real-time tracking of pathogen evolution. *Bioinformatics* 34, 4121 4123 (2018).
- C. Gao *et al.*, SARS-CoV-2 Spike Protein Interacts with Multiple Innate Immune Receptors. *bioRxiv*,
 2020.2007.2029.227462 (2020).
- M. Sanda, L. Morrison, R. Goldman, N- and O-Glycosylation of the SARS-CoV-2 Spike Protein. *Anal Chem* 93, 2003-2009 (2021).
- 242 15. X. Xie *et al.*, An Infectious cDNA Clone of SARS-CoV-2. *Cell Host Microbe* **27**, 841-848 e843 (2020).
- 24316.X. Xie *et al.*, Engineering SARS-CoV-2 using a reverse genetic system. Nature Protocols 16, 1761-2441784 (2021).
- 24517.T. P. Peacock *et al.*, The altered entry pathway and antigenic distance of the SARS-CoV-2 Omicron246variant map to separate domains of spike protein. *bioRxiv*, 2021.2012.2031.474653 (2022).
- 24718.B. Meng *et al.*, Altered TMPRSS2 usage by SARS-CoV-2 Omicron impacts infectivity and248fusogenicity. *Nature* 603, 706-714 (2022).
- 249
 19.
 K. S. Phadke, N. B. A. Higdon, B. H. Bellaire, In vitro comparison of SARS-CoV-2

 250
 variants. *bioRxiv*, 2023.2003.2011.532212 (2023).
- 20. H. Zhao *et al.*, SARS-CoV-2 Omicron variant shows less efficient replication and fusion activity
 when compared with Delta variant in TMPRSS2-expressed cells. *Emerging Microbes & Infections* 11, 277-283 (2022).
- 254 21. C.-L. Hsieh *et al.*, Structure-based design of prefusion-stabilized SARS-CoV-2 spikes. *Science* 369, 1501-1505 (2020).
- 25. Z. Mustafa, H. Kalbacher, T. Burster, Occurrence of a novel cleavage site for cathepsin G adjacent
 to the polybasic sequence within the proteolytically sensitive activation loop of the SARS-CoV-2
 Omicron variant: The amino acid substitution N679K and P681H of the spike protein. *PLOS ONE* 17, e0264723 (2022).

- 260 23. S. Yuan *et al.*, Pathogenicity, transmissibility, and fitness of SARS-CoV-2 Omicron in Syrian
 261 hamsters. *Science* **377**, 428-433 (2022).
- 262 24. C. P. McCormack *et al.*, Modelling the viral dynamics of the SARS-CoV-2 Delta and Omicron 263 variants in different cell types. *bioRxiv*, 2023.2003.2015.529513 (2023).
- 264 25. K. P. Y. Hui *et al.*, SARS-CoV-2 Omicron variant replication in human bronchus and lung ex vivo.
 265 *Nature* 603, 715-720 (2022).
- 26. A. M. Carabelli *et al.*, SARS-CoV-2 variant biology: immune escape, transmission and fitness.
 267 Nature Reviews Microbiology **21**, 162-177 (2023).
- 268 27. V. Papanikolaou *et al.*, From delta to Omicron: S1-RBD/S2 mutation/deletion equilibrium in SARS 269 CoV-2 defined variants. *Gene* 814, 146134 (2022).
- 270 28. L. B. Shrestha, C. Foster, W. Rawlinson, N. Tedla, R. A. Bull, Evolution of the SARS-CoV-2 omicron
 271 variants BA.1 to BA.5: Implications for immune escape and transmission. *Reviews in Medical*272 *Virology* 32, e2381 (2022).
- 273 29. D. Deming *et al.*, Vaccine Efficacy in Senescent Mice Challenged with Recombinant SARS-CoV
 274 Bearing Epidemic and Zoonotic Spike Variants. *PLOS Medicine* **3**, e525 (2006).
- 27530.J. Harcourt *et al.*, Severe Acute Respiratory Syndrome Coronavirus 2 from Patient with276Coronavirus Disease, United States. *Emerging infectious diseases* **26**, 1266-1273 (2020).
- 27731.B. A. Johnson *et al.*, Nucleocapsid mutations in SARS-CoV-2 augment replication and278pathogenesis. *PLOS Pathogens* **18**, e1010627 (2022).

279

280

281 Acknowledgments

- 282 Research was supported by grants from NIAID of the NIH to (R01-AI153602 and R21-AI145400
- to VDM; R24-AI120942 (WRCEVA) to SCW). ALR was supported by a UTMB Institute for Human
- 284 Infection and Immunity grant and the Sealy and Smith Foundation. Research was also supported
- by STARs Award provided by the University of Texas System to VDM and Data Acquisition award
- provided by the Institute for Human Infections and Immunity at UTMB to MNV. Trainee funding
- provided by NIAID of the NIH to MNV (T32-Al060549).
- 288 Mass spectrometry experiments and analysis was provided by the Mass Spectrometry Facility at
- the University of Texas Medical Branch (https://www.utmb.edu/MSF). Figures were created with
- 290 BioRender.com

291 Competing Interest Statement

- 292 VDM has filed a patent on the reverse genetic system and reporter SARS-CoV-2. MNV and VDM
- have filed a provisional patent on a stabilized SARS-CoV-2 spike protein. Other authors declareno competing interests.
- 295 Author contributions
- 296 Conceptualization: MNV, VDM
- 297 Formal analysis: MNV, VDM
- 298 Funding acquisition: MNV, ALR, SCW, VDM
- 299 Investigation: MNV, DRM, REA, KL, CS, JAP, LKE, AMM, YPA, WMM, ALM, JM, DHW, KP, ALR
- 300 Methodology: MNV, KSP, VDM
- 301 Project Administration: MNV, VDM
- 302 Supervision: MNV, SCW, DHW, KSP, VDM

- 303 Visualization: MNV, DHW, VDM
- 304 Writing original draft: MNV, VDM
- 305 Writing review and editing: MNV, REA, CS, DHW, VDM, SCW

306 Figure Legends

307 Figure 1. The combination of Omicron mutations H655Y, N679K, and P681H increases viral

308 replication and spike processing.

- 309 (A) Comparison of CTS1 region near the S1/S2 cleavage site between SARS-CoV-2 variants.
- 310 (B) Structure of loop containing the S1/S2 cleavage site on SARS-CoV-2 spike protein. The
- residues that are mutated in Omicron are shown: H655 (magenta), N679 (green), and P681
- 312 (blue). The furin cleavage site RRAR (cyan) and QTQT motif (red) are also shown.
- 313 **(C)** Schematic of WT and YKH SARS-CoV-2 mutant genomes.
- 314 (D) WT and YKH SARS-CoV-2 plaques on Vero E6 cells at 2 dpi.

315 (E) Viral titer from WT and YKH virus stocks representing the highest yield generated from
 316 TMPRSS2-expressing Vero E6 cells.

317 (F-G) Replication kinetics of WT and YKH in Vero E6 (F) and Calu-3 2B4 (G) cells. Cells were

infected at an MOI of 0.01 infectious units/cell (n=3). Data are mean ± s.d. Statistical analysis
measured by two-tailed Student's t-test.

320 **(H)** Purified WT, YKH, Delta isolate (B.1.617.2), and Omicron (BA.1) virions from Vero E6 321 supernatant were probed with α -Spike and α -Nucleocapsid (N) antibodies in Western blots. Full-322 length spike (FL), S1/S2 cleavage product, and S2' cleavage product are indicated.

(I) Densitometry of FL and S1/S2 cleavage product was performed, and quantification of FL and
 S1/S2 cleavage product percentage of total spike is shown. Quantification was normalized to N
 for viral protein loading control. WT (black), YKH (blue), Delta isolate (purple), Omicron (orange).
 Results are representative of two experiments.

327 Figure 2. N679K attenuates SARS-CoV-2 replication and disease.

- 328 (A) Structural modeling of O-linked glycosylation of threonine 678 (yellow) of QTQTN motif (red)
- and the residues mutated in Omicron H655 (magenta), N679 (green), and P681 (blue) with
- N679 adjacent to the glycosylation. The furin cleavage site RRAR is also shown (cyan).
- 331 **(B)** Schematic of WT and N679K SARS-CoV-2 genomes.
- (C) WT and N679K SARS-CoV-2 plaques on Vero E6 cells at 2 (left) and 3 dpi (right). Average
 plaque sizes noted below.

(D) Viral titer from WT and N679K virus stocks with the highest yield generated form TMPRSS2 expressing Vero E6 cells.

- (E-F) Replication kinetics of WT and N679K in Vero E6 (E) and Calu-3 2B4 (F) cells. Cells were
 infected at an MOI of 0.01 infectious units/cell (n=3). Data are mean ± s.d. Statistical analysis
 performed using two-tailed Student's t-test.
- (G) Schematic of experimental design for golden Syrian hamster infection with WT (black) or
 N679K (green) SARS-CoV-2. Three- to four-week-old male hamsters were infected with 10⁵ pfu
 and monitored for weight loss over 7 days. At 2, 4, and 7 dpi, nasal washes and lungs were
 collected for viral titer, and lung was collected for histopathology.

343 **(H)** Weight loss of hamsters infected with WT (black) or N679K (green) SARS-CoV-2 over 7 days.

- Data are mean ± s.e.m. Statistical analysis measured by two-tailed Student's t-test.
- (I-J) Viral titers of lungs (I) and nasal washes (J) collected at 2 and 4 dpi from hamsters infected
 with WT (black) or N679K (green) SARS-CoV-2. Data are mean ± s.d. Statistical analysis
 measured by two-tailed Student's t-test.

348 Figure 3. N679K results in decreased spike expression on virions and in infected cells.

(A) Purified WT, N679K, and Omicron (BA.1) virions from Vero E6 supernatants were probed with

α-Spike and α-Nucleocapsid (N) antibodies in Western blots. Full-length spike (FL), S1/S2
 cleavage product, and S2' cleavage product are indicated.

- (B) Densitometry of spike processing from purified virions applied to Western blots in (A) was
 performed, and quantification of FL and S1/S2 cleavage product percentage of total spike is
 shown. Quantification was normalized to N as viral protein loading control. WT (black), N679K
 (green), Omicron (orange). Results are representative of two experiments.
- 356 (C) Densitometry of spike expression from purified virion Western blots in (A) was performed, and

357 guantification of total spike protein to nucleocapsid ratio is shown. Spike/N ratio is relative to WT.

358 WT (black), N679K (green), Omicron (orange). Results are representative of two experiments.

359 **(D)** Vero E6 cells were infected with WT, N679K, or Omicron at an MOI of 0.01 infectious 360 units/cell. Cell lysate was collected at 24 hpi and probed with α -Spike and α -Nucleocapsid (N) 361 antibodies in Western blots. Full-length spike (FL), S1/S2 cleavage product, and S2' cleavage 362 product are indicated.

(E) Densitometry of spike expression from infected cell lysate Western blots in (D) was performed,
 and quantification of total spike protein to nucleocapsid ratio is shown. Spike/N ratio is relative to
 WT. WT (black), N679K (green), Omicron (orange). Results are representative of three biological
 replicates.

367 **(F)** Vero E6 cells were transfected with Spike HexaPro WT and N679K and cell lysate was 368 collected at 8, 24, and 48 hpt. Lysates were probed with α -Spike and α -GAPDH antibodies in 369 Western blots.

- 370 (G) Densitometry of spike expression from transfected cell lysates by Western blot in (G) was
- 371 performed, and quantification of relative total spike protein is shown. Spike protein levels were
- 372 normalized to GAPDH and are relative to WT. WT (black), N679K (green). Results are
- 373 representative of three biological replicates.

374 Methods

375 Cell Culture

Vero E6 cells were grown in high glucose DMEM (Gibco #11965092) with 10% fetal bovine serum

and 1x antibiotic-antimycotic. TMPRSS2-expressing Vero E6 cells were grown in low glucose

378 DMEM (Gibco #11885084) with sodium pyruvate, 10% FBS, and 1 mg/mL Geneticin[™] (Invitrogen

379 #10131027). Calu-3 2B4 cells were grown in high glucose DMEM (Gibco #11965092) with 10%

- defined fetal bovine serum, 1 mM sodium pyruvate, and 1x antibiotic-antimycotic.
- 381 Viruses

382 The SARS-CoV-2 infectious clones were based on the USA-WA1/2020 sequence provided by 383 the World Reference Center of Emerging Viruses and Arboviruses and the USA Centers for Disease Control and Prevention (30). Mutant viruses (YKH and N679K) were generated with 384 restriction enzyme-based cloning using gBlocks encoding the mutations (Integrated DNA 385 Technologies) and our reverse genetics system as previously described (15, 16). Virus stock was 386 387 generated in TMPRSS2-expressing Vero E6 cells to prevent mutations from occurring at the FCS. Viral RNA was extracted from virus stock and cDNA was generated to verify mutations by 388 389 Sanger sequencing.

Delta isolate (B.1.617.2) was obtained from the World Reference Center of Emerging Viruses and
Arboviruses. Infectious clone of Omicron (BA.1) was obtained from Dr. Pei Yong Shi and Dr.
Xuping Xie.

393 In vitro Infection

Vira infections in Vero E6 and Calu-3 2B4 were carried out as previously described (8). Briefly, growth media was removed, and cells were infected with WT or mutant SARS-CoV-2 at an MOI of 0.01 for 45 min at 37°C with 5% CO₂. After absorption, cells were washed three times with PBS and fresh complete growth media was added. Three or more biological replicates were collected at each time point and each experiment was performed at least twice. Samples were titrated with plague assay or focus forming assays.

400 Plaque Assay

Vero E6 cells were seeded in 6-well plates and grown to 80-100% confluency in complete growth media. Ten-fold serial dilutions in PBS were performed on virus samples. Growth media was removed from cells and 200 μ l of inoculum was added to monolayers. Cells were incubated for 404 45 min at 37°C with 5% CO₂. After absorption, 0.8% agarose overlay was added, and cells were 405 incubated at 37°C with 5% CO₂ for 2 days. Plaques were visualized with neutral red stain. Average 406 plaque size was determined using ImageJ.

407 Focus Forming Assay

408 Focus forming assays (FFAs) were performed as previously described (31). Briefly, Vero E6 cells 409 were seeded in 96-well plates to be 100% confluent. Samples were 10-fold serially diluted in serum-free media and 20 µl was to infect cells. Cells were incubated for 45 min at 37°C with 5% 410 CO2 before 100 µl of 0.85% methylcellulose overlay was added. Cells were incubated for 24 h 45 411 412 min at 37°C with 5% CO₂. After incubation, overlay was removed, and cells were washed three 413 times with PBS before fixed and virus inactivated by 10% formalin for 30 min at room temperature. Cells were then permeabilized and blocked with 0.1% saponin/0.1% BSA in PBS before incubated 414 with α -SARS-CoV-2 Nucleocapsid primary antibody (Cell Signaling Technology) at 1:1000 in 415 416 permeabilization/blocking buffer overnight at 4°C. Cells are then washed three times with PBS before incubated with Alexa FluorTM 555-conjugated α -mouse secondary antibody (Invitrogen 417 #A28180) at 1:2000 in permeabilization/blocking buffer for 1 h at room temperature. Cells were 418 419 washed three times with PBS. Fluorescent foci images were captured using a Cytation 7 cell 420 imaging multi-mode reader (BioTek), and foci were counted manually.

421 Hamster Infection

Three- to four-week-old male golden Syrian hamsters (HsdHan:AURA strain) were purchased from Envigo. All studies were conducted under a protocol approved by the UTMB Institutional Animal Care and Use Committee and complied with USDA guidelines in a laboratory accredited by the Association for Assessment and Accreditation of Laboratory Animal Care. Procedures

involving infectious SARS-CoV-2 were performed in the Galveston National Laboratory ABSL3 facility. Hamsters were intranasally infected with 10^5 pfu of WT or N679K SARS-CoV-2 in 100 µl. Infected hamsters were weighed and monitored for illness over 7 days. Hamsters were anesthetized with isoflurane and nasal washes were collected with 400 µl of PBS on endpoint days (2, 4, and 7 dpi). Hamsters were euthanized by CO₂ for organ collection. Nasal wash and lung were collected to measure viral titer and RNA. Left lungs were collected for histopathology.

432 Virion Purification

Vero E6 cells were grown in T175 flasks to be 100% confluent at time of infection. Cells were infected with 50 µl of virus stock in PBS for 45 min at 37°C with 5% CO₂, and growth media with 5% FBS was added after absorption. Supernatant was harvested at 24 hpi and clarified by low-speed centrifugation. Virions were purified from supernatant by ultracentrifugation through a 20% sucrose cushion at 26,000 rpm for 3 hrs using a Beckman SW28 rotor. Pellets were resuspended with 2x Laemmli buffer to obtain protein samples for Western blot.

439 Western Blot

Protein levels were determined by SDS-PAGE followed by western blot analysis as previously 440 described (8). In brief, sucrose-purified SARS-CoV-2 virions were inactivated by resuspending in 441 442 2x Laemmli buffer and boiling. SDS-PAGE gels were run with equal volumes of samples on Mini-PROTEAN TGX gels (Bio-Rad #4561094) followed by transfer onto PVDF membrane. 443 Membranes were incubated with α-SARS-CoV S primary antibody (Novus Biologicals #NB100-444 56578) at 1:1000 dilution in 5% BSA in TBST to measure spike protein processing and 445 446 expression. For loading control, α-SARS Nucleocapsid primary antibody (Novus Biologicals 447 #NB100-56576) at 1:1000 in 5% BSA in TBST was used for viral loading control and α -GAPDH primary antibody (Invitrogen #AM4300) at 1:1000 in 5% BSA in TBST for cellular loading control. 448 Primary antibody incubation was followed by HRP-conjugated α -rabbit secondary antibody (Cell 449 450 Signaling Technology #7074) or HRP-conjugated α-mouse secondary antibody (Cell Signaling 451 Technology #7076) at 1:3000 in 5% milk in TBST. Chemiluminescence signal was developed

- 452 using Clarity Western ECL substrate (Bio-Rad #1705060) or Clarity Max Western ECL substrate
- 453 (Bio-Rad #1705062) and imaged with a ChemiDoc MP System (Bio-Rad). Densitometry analysis
- 454 was performed using ImageLab 6.0.1 (Bio-Rad).
- 455 Spike HexaPro Cloning and Transfection

456 SARS-CoV-2 S HexaPro was a gift from Jason McLellan (Addgene plasmid #154754) (21). The

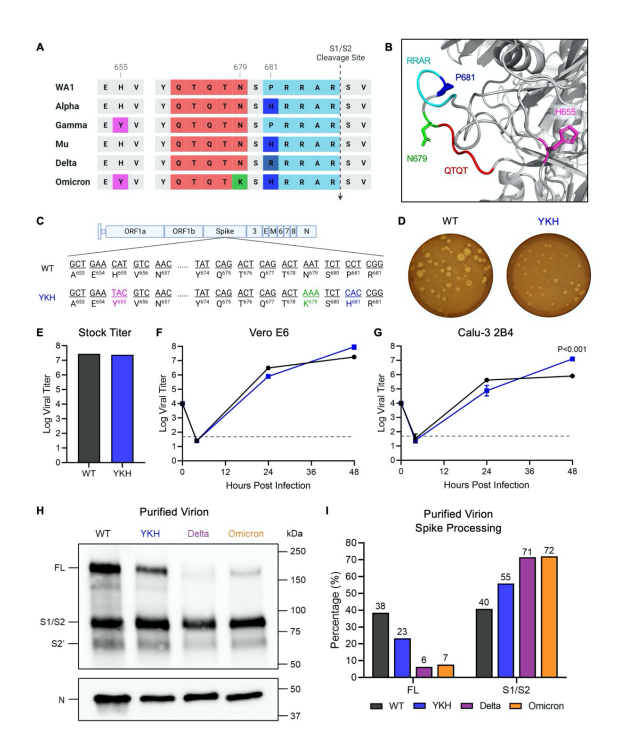
457 N679K mutation was cloned into spike HexaPro using a gBlock encoding the mutation (Integrated

- 458 DNA Technologies) and restriction enzyme-based cloning. Sequences were verified by Sanger 459 sequencing.
- Vero E6 cells were grown in 24-well plates to be 100% confluent at time of transfection. Cells were transfected with spike HexaPro WT or N679K plasmid and Lipofectamine 2000 following manufacturer's instructions (Invitrogen). Briefly, 100 ng of spike HexaPro plasmid and 1.5 µl of Lipofectamine 2000 were separately diluted in 50 µl Opti-MEM (Gibco #31985070) before mixing together. After 20 min of room temperature incubation, 100 µl of the transfection mixture was added to cells, and cells were incubated at 37°C with 5% CO₂. Cell lysate was harvested with 2x Laemmli buffer at 24 and 48 hours post transfection to be analyzed by Western blot.

467 Structural Modeling

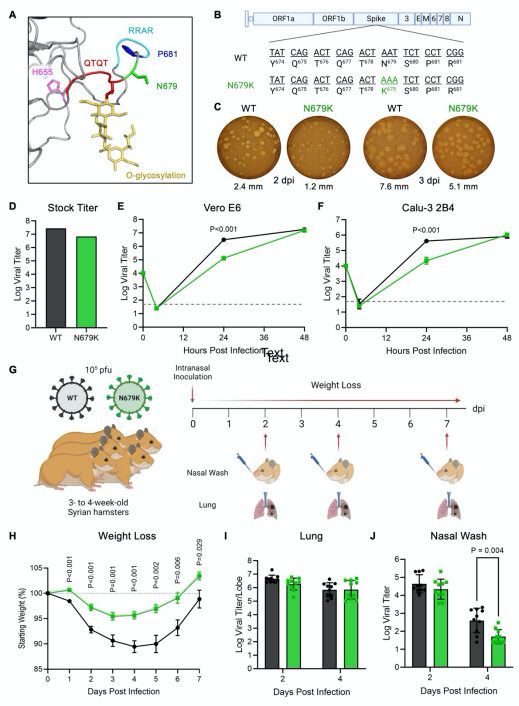
468 Structural models previously generated were used as a base to visualize residues mutated in 469 Omicron (8). Briefly, structural models were generated using SWISS-Model to generate homology 470 models for WT and glycosylated SARS-CoV-2 spike protein on the basis of the SARS-CoV-1 471 trimer structure (Protein Data Bank code 6ACD). Homology models were visualized and 472 manipulated in PyMOL (version 2.5.4) to visualize Omicron mutations.

473



474

475 Figure 1. The combination of Omicron mutations H655Y, N679K, and P681H increases viral replication and spike processing. 476 477 (A) Comparison of CTS1 region near the S1/S2 cleavage site between SARS-CoV-2 variants. (B) Structure of loop containing the S1/S2 cleavage site on SARS-CoV-2 spike protein. The residues that are mutated in Omicron are shown - H655 (magenta), N679 478 (green), and P681 (blue). The furin cleavage site RRAR (cyan) and QTQT motif (red) are also shown. (C) Schematic of WT and YKH 479 SARS-CoV-2 mutant genomes. (D) WT and YKH SARS-CoV-2 plagues on Vero E6 cells at 2 dpi. (E) Viral titer from WT and YKH 480 virus stock with the highest yield generated from TMPRSS2-expressing Vero E6 cells. (F-G) Growth kinetics of WT and YKH in Vero 481 482 E6 (F) and Calu-3 2B4 (G) cells. Cells were infected at an MOI of 0.01 (n=3). Data are mean ± s.d. Statistical analysis measured by two-tailed Student's t-test. (H) Purified WT, YKH, Delta isolate (B.1.617.2), and Omicron (BA.1) virions from Vero E6 supernatant were probed with α-Spike and α-Nucleocapsid (N) antibodies in Western blots. Full-length spike (FL), S1/S2 cleavage product, and S2' 483 484 485 486 cleavage product are indicated. (I) Densitometry of FL and S1/S2 cleavage product was performed, and quantification of FL and S1/S2 cleavage product percentage of total spike is shown. Quantification was normalized to N for viral protein loading control. WT (black), YKH (blue), Delta isolate (purple), Omicron (orange). Results are representative of two experiments.



487 488 Figure 2. N679K attenuates SARS-CoV-2 replication and disease when isolated.

489 (A) Structural modeling of O-linked glycosylation of threonine 678 (yellow) of QTQTN motif (red) and the residues mutated in Omicron 490 - H655 (magenta), N679 (green), and P681 (blue) – with N679 adjacent to the glycosylation. The furin cleavage site RRAR is also 491 shown (cyan). (B) Schematic of WT and N679K SARS-CoV-2 genomes. (C) WT and N679K SARS-CoV-2 plaques on Vero E6 cells 492 493 at 2 dpì (left) and 3 dpi (right). Average plaque size noted below. (D) Viral titer from WT and N679K virus stock with the highest yield generated form TMPRSS2-expressing Vero E6 cells. (E-F) Growth kinetics of WT and N679K in Vero E6 (E) and Calu-3 2B4 (F) cells. 494 495 Cells were infected at an MOI of 0.01 (n=3). Data are mean ± s.d. Statistical analysis measured by two-tailed Student's t-test. (G) Schematic of experiment design for golden Syrian hamster infection with WT (black) or N679K (green) SARS-CoV-2. Three- to four-496 week-old make hamsters were infected with 105 pfu and monitored for weight loss over 7 days. At 2, 4, and 7 dpi, nasal wash and 497 lung was collected for viral titer, and lung was collected for histopathology. (H) Weight loss of hamsters infected with WT (black) or 498 N679K (green) SARS-CoV-2 over 7 days. Data are mean ± s.e.m. Statistical analysis measured by two-tailed Student's t-test. (I-J) 499 Viral titer of lung (I) and nasal wash (J) collected at 2 and 4 dpi from hamsters infected with WT (black) or N679K (green) SARS-CoV-500 2. Data are mean ± s.d. Statistical analysis measured by two-tailed Student's t-test.

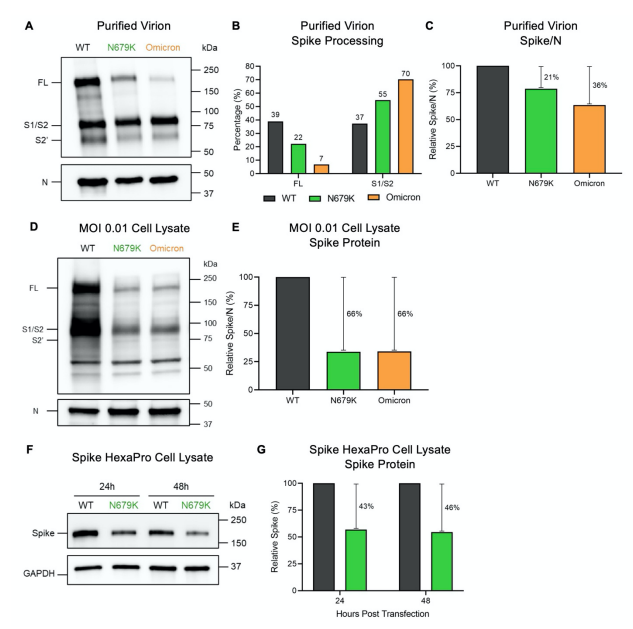


Figure 3. N679K results in decreased spike expression on virions and in cell lysate. (A) Purified WT, N679K, and Omicron (BA.1) virions from Vero E6 supernatant were probed with α-Spike and α-Nucleocapsid (N) antibodies in Western blots. Full-length spike (FL), S1/S2 cleavage product, and S2' cleavage product are indicated. (B) Densitometry of spike processing from purified virion Western blot in (A) was performed, and quantification of FL and S1/S2 cleavage product percentage of total spike is shown. Quantification was normalized to N as viral protein loading control. WT (black), N679K (green), Omicron (orange). Results are representative of two experiments. (C) Densitometry of spike expression from purified virion Western blot in (A) was performed, and quantification of total spike protein to nucleocapsid ratio is shown. Spike/N ratio is relative to WT. WT (black), N679K (green), Omicron (orange). Results are representative of two experiments. (D) Vero E6 cells were infected with WT, N679K, or Omicron at an MOI of 0.01. Cell lysate was collected at 24 hpi and probed with α-Spike and α-Nucleocapsid (N) antibodies in Western blots. Full-length spike (FL), S1/S2 cleavage product, and S2' cleavage product are indicated. (E) Densitometry of spike expression from infected cell lysate Western blot in (D) was performed, and quantification of total spike protein to nucleocapsid ratio is shown. Spike/N ratio is relative to WT. WT (black), N679K (green), Omicron (orange). Results are representative of three biological replicates. (F) Vero E6 cells were transfected with Spike HexaPro WT and N679K and cell lysate was collected at 8, 24, and 48 hpt. Lysates were probed with α-Spike and α-GAPDH antibodies in Western blots.(G) Densitometry of spike expression from transfected cell lysate Western blot in (G) was performed, and quantification of relative total spike protein is shown. Spike protein levels were normalized to GAPDH 517 and are relative to WT. WT (black), N679K (green). Results are representative of three biological replicates.