Phosphatidylserine Receptors Enhance SARS-CoV-2 Infection: AXL as a Therapeutic Target for COVID-19

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24 AUTHOR SUMMARY

- 25 Phosphatidylserine (PS) receptors are PS binding proteins that mediate uptake of apoptotic
- 26 bodies. Many enveloped viruses utilize this PS/PS receptor mechanism to adhere to and
- 27 internalize into the endosomal compartment of cells and this is termed apoptotic mimicry. For
- viruses that have a mechanism(s) of endosomal escape, apoptotic mimicry is a productive route
- 29 of virus entry. We evaluated if PS receptors serve as cell surface receptors for SARS-CoV-2
- 30 and found that the PS receptors, AXL, TIM-1 and TIM-4, facilitated virus infection when low
- 31 concentrations of the SARS-CoV-2 cognate receptor, ACE2, was present. Consistent with the
- 32 established mechanism of PS receptor utilization by other viruses, PS liposomes competed with
- 33 SARS-CoV-2 for binding and entry. We demonstrated that this PS receptor enhances SARS-
- 34 CoV-2 binding to and infection of an array of human lung cell lines and is an under-appreciated
- 35 but potentially important host factor facilitating SARS-CoV-2 entry.
- 36 37

38 INTRODUCTION

- 39 Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) emerged in late 2019 and
- 40 guickly spread around the world, resulting in the current public health pandemic. SARS-CoV-2 is
- 41 a beta coronavirus of the sarbecovirus subgenus and is closely related to SARS-CoV, the agent
- 42 responsible for an epidemic in 2003. SARS-CoV-2 is effectively transmitted between humans
- 43 and has infected more than 178 million individuals and caused more than 3.88 million deaths
- 44 worldwide as of June 23, 2021 (WHO). Fortunately, a herculean scientific effort has resulted in
- 45 the development of SARS-CoV-2 vaccines which have been shown to be efficacious, potentially
- 46 stemming the pandemic. Nonetheless, in combination with vaccines, continued development of
- 47 efficacious antivirals is needed, as outbreaks continue in under-vaccinated regions and severe
- 48 disease is not eradicated following vaccination. Towards this goal, a more comprehensive
- 49 understanding of SARS-CoV-2 interactions with host cells will be required.
- 50
- 51 SARS-CoV-2 entry into cells is mediated by the viral spike glycoprotein (S) binding to
- 52 Angiotensin Converting Enzyme 2 (ACE2) (1-3). The S1 subunit of S binds to ACE2 while S2
- 53 mediates membrane fusion. Cleavage at the S1/S2 junction occurs during virus egress from
- 54 producer cells by the host protease furin which facilitates S1 binding to ACE2. A second site
- 55 termed S2' is also cleaved by the host proteases. Cleavage by TMPRSS2 at the cell surface
- 56 promotes fusion of the viral and host plasma membranes (2, 4). Alternatively, SARS-CoV-2
- 57 virions can be internalized via clathrin-mediated endocytosis after ACE2 binding, wherein host

cathepsins (especially cathepsin L) proteolytically cleave S2' (3, 5-7). Adherence factors that
enhance virion binding and increase infectivity have also been identified, namely neuropilin 1
and heparan sulfate (8, 9).

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62 Binding and internalization of a variety of different enveloped viruses occurs through virion 63 associated phosphatidylserine (PS) binding to PS receptors. Members of the TIM family (TIM-1 64 and TIM-4) bind PS directly while another family of PS receptors, the TAM tyrosine kinase 65 receptor family (TYRO3, AXL and MERTK), bind PS indirectly through the adaptor proteins 66 Gas6 and Protein S. These PS receptor mediate binding and internalization of a wide range of 67 viruses, including filoviruses, alphaviruses, and flaviviruses (10-13). TIM-1, TIM-4, and AXL appear to be the most efficacious at mediating viral entry given their prevalent use among 68 69 enveloped viruses (14-16). Once virions are within the endosome, events that result in virion 70 fusion with cellular membranes are virus specific, with filoviruses requiring viral glycoprotein 71 processing followed by interactions with Niemann Pick C1 protein (NPC1) to initiate fusion, 72 whereas flaviviruses rely on endosomal acidification driving glycoprotein conformational 73 changes which mediate fusion (17, 18).

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75 Given that PS receptors mediate entry of other enveloped viruses through interactions with viral 76 membrane PS, we assessed the role of PS receptors on SARS-CoV-2 infection and the 77 mechanism of interaction. We found that plasma membrane-expressed PS receptors by 78 themselves do not result in productive SARS-CoV-2 infection; however, these receptors 79 enhance infection when low levels of ACE2 are expressed. Our findings indicate that these 80 receptors synergize with ACE2 to mediate SARS-CoV-2 entry through PS-dependent 81 interactions. These data are in contrast to the conclusions drawn by another report stating that 82 AXL interacts directly with the SARS-CoV-2 spike protein (19). Appreciation of this route of entry 83 provides an additional pathway that could be therapeutically targeted to inhibit virus entry and 84 subsequent infection.

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87 RESULTS

88 **PS receptors enhance ACE2-dependent SARS-CoV-2 infection**

89 The ability of TIM and TAM family PS receptors to support SARS-CoV-2 infection were initially

90 examined in transfected HEK 293T cells. Wild-type HEK 293T cells do not express significant

91 amounts of ACE2 or PS receptors and are poorly permissive to SARS-CoV-2 infection (3, 20).

92 Expression plasmids encoding ACE2 and/or the PS receptors, AXL or TIM-1, were transfected, 93 resulting in expression of these receptors on the surface of the transfected cells (S1A-B). Dual 94 transfection did not alter expression of ACE2 or PS receptors relative to single transfection. The 95 PS receptors AXL and TIM-1 by themselves did not facilitate infection of SARS-CoV-2 or VSV 96 pseudovirions bearing SARS-CoV-2 spike (VSV/Spike) (Fig. 1A, S1C). However, when low 97 levels of ACE2 (50 to 250 ng of plasmid) were co-expressed with either AXL or TIM-1, the 98 combinations enhanced infection over that observed with ACE2 alone. TIM-1-enhanced ACE2-99 dependent recombinant VSV/Spike (rVSV/Spike) (21) infection, and enhancement occurred 100 over a wider range of ACE2 concentrations than AXL-enhanced ACE2-dependent infection, with 101 AXL consistently enhancing infection when 250 ng of ACE2 plasmid was transfected (Fig. 1B-102 **C**, **S1C**). The more limited ability of AXL to synergize was not due to limiting Gas6 in the media 103 as the addition of Gas6 to media did not enhance the synergy. At higher concentrations of 104 ACE2 plasmid, PS receptor enhancement of infectivity was reduced and, with transfection of 1 105 µg of ACE2 plasmid, no PS receptor enhancement was observed. Thus, only when ACE2 is 106 limiting on the cell surface do PS receptors facilitate infection. Consistent with a role for PS 107 receptors in SARS-CoV-2 entry, we observed enhanced virion attachment to cells when PS 108 receptors were expressed (Fig. 1D). In addition to viral load assessments, supernatants from 109 SARS-CoV-2 infected HEK 293T cultures were evaluated for production of infectious virions at 110 48 hpi by TCID₅₀ assays in Vero E6 cells that express TMPRSS2. Low levels (50 ng) of ACE2 111 transfection increased virion production and this was enhanced by co-expression of TIM-1 (Fig. 112 **1E**). Consistent with the viral load findings, AXL co-expression was not effective at enhancing 113 production of infectious virus when this low level of AXL was transfected. Other PS receptors, 114 TIM-4, TYRO3, and MerTK were examined for their ability of enhance infection. TIM-4 enhanced ACE2-dependent entry of VSV/Spike in a manner similar to TIM-1; however, TYRO3 115 116 and MerTK of the TAM family did not mediate increased entry, despite detectable levels of plasma membrane expression after transfection (Fig. 1F; S1A-B; S1D). The synergy between 117 118 PS receptors and ACE2 was specific for SARS-CoV-2 as infection with VSV-luciferase 119 pseudovirions bearing Lassa virus GP was not affected by expression of these receptors (S1E). 120 These studies indicate that PS receptors synergize with low levels of ACE2 to enhance SARS-121 CoV-2 infection. Further, these data provide evidence that VSV/Spike pseudovirions serve as a 122 useful BSL2 surrogate for SARS-CoV-2 entry events as others have shown (22-24). 123

124 PS receptors bind to virion associated PS, not the SARS-CoV-2 Spike protein

125 We took several different approaches to examine the mechanism by which PS receptors 126 interact with SARS-CoV-2. In the context of other viral pathogens, PS receptors are known to 127 bind to virion membrane associated PS and mediate endosomal internalization of virions, 128 shuttling virus to cognate endosomal receptors. Liposomes composed of PS compete for virion 129 binding to PS receptors (25). To test this with SARS-CoV-2, increasing concentrations of PS or 130 phosphatidylcholine (PC) liposomes were evaluated for their ability to compete with virus for PS 131 binding sites in ACE2 + AXL or ACE2 + TIM-1 transfected HEK 293T cells. PS liposomes effectively blocked VSV/Spike entry, whereas PC liposomes were significantly less effective 132 133 (Fig. 2A-B). We also assessed the activity of a TIM-1 mutant, ND115DN which has a disrupted 134 TIM-1 PS binding pocket, for its ability to facilitate VSV/Spike entry. This TIM-1 mutant did not 135 synergize with ACE2, indicating that the TIM-1 PS binding pocket is critical for this enhanced 136 activity (Fig. 2C) (12, 20).

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138 Others have reported that the N-terminal domain of SARS-CoV-2 spike directly binds to AXL 139 and is important for AXL-mediated entry of SARS-CoV-2 (19). To assess spike/AXL 140 interactions, purified soluble full-length SARS-CoV-2 spike-Fc, spike receptor binding domain-141 Fc (RBD) or spike N-terminal domain-Fc (NTD) was incubated with HEK 293T cells transiently 142 expressing AXL. Flow cytometry was used to detect spike proteins bound to AXL. Parallel ACE2 143 binding to soluble SARS-CoV-2 spike served as a positive control. As the spike NTD-Fc was 144 not expected to bind to ACE2, an ELISA confirmed the ability of a conformationally dependent 145 α -spike NTD monoclonal antibody to bind NTD-Fc, suggesting that NTD-Fc was in its native 146 conformation (S2B). The full-length spike-Fc and the RBD-Fc bound to ACE2, but no 147 interactions were detected between any of the purified spike proteins and AXL (Fig. 2D) despite 148 evidence of robust AXL surface expression on transfected HEK 293T cells (S2A) and the 149 equivalent levels of detection of the purified proteins via ELISAs (S2C). Biolayer interferometry 150 studies confirmed and extended our findings that recombinant AXL does not bind to purified 151 NTD, whereas interaction with the α-spike NTD monoclonal antibody was readily detected (Fig. 152 **2E**). Thus, using two complementary approaches, we were unable to demonstrate direct 153 interactions of AXL with spike. In total, our studies are consistent with PS receptors interacting 154 with SARS-CoV-2 virions through the well-established mechanism of virion-associated PS 155 binding to TIM-1 and AXL.

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157 Redundant routes of virus entry: endosomal vs. plasma membrane mediated infection

ACE2-dependent coronaviruses enter cells through two different routes: 1) An endosomal route of virus uptake that requires low pH-dependent cathepsin L processing of spike, and 2) A plasma membrane route that is dependent upon TMPRSS2 cleavage of spike (3, 26). Others have reported that TMPRSS2-dependent entry is preferentially utilized by the virus when this serine protease is expressed (27). We examined the route of virus entry at play when ACE2, PS receptors and/or TMPRSS2 was expressed.

- 166 ACE2-dependent infection. As anticipated, we found that low levels (10 ng) of TMPRSS2 167 expressing plasmid enhanced VSV/Spike pseudovirion entry in HEK 293T cells co-transfected 168 with 50 ng of ACE2 expressing plasmid (Fig. 3A). However, at higher concentrations of 169 TMPRSS2 plasmid. TMPRSS2 did not enhance infection, perhaps due to aberrant protease 170 activity. Inverting these variables, when 10 ng of TMPRSS2 plasmid was transfected in the 171 presence of low concentrations of ACE2, TMPRSS2 enhanced virus infection (Fig. 3B). At high 172 concentrations of ACE2, virus entry became TMPRSS2-independent in a manner similar to the 173 effects of the PS receptors. Taken with Fig. 1B and C, these studies indicate that the PS 174 receptors and TMPRSS2 can facilitate ACE2-dependent virus infection when ACE2 is limiting. 175 but with increasing ACE2 concentrations the infections become independent of these entry 176 factors. This may be related to effects of soluble ACE2 on entry (28).
- 177

178 To evaluate how PS receptors and/or TMPRSS2 expression would alter the route of ACE2-

dependent infection, HEK 293T cells were transfected PS receptors as before and infected with

180 VSV/Spike in the presence or absence of the cysteine protease inhibitor, E-64, that blocks

181 endosomal cathepsin activity. Non-toxic levels of E-64 were effective at blocking ACE2-

dependent infection (**Fig. 3C, S3**), indicating that virions were entering these cells in a cysteine

183 protease-dependent manner, likely through the endosomal compartment. The enhancement of

virus entry conferred by the combination of PS receptors and ACE2 was also inhibited by E-64,

providing evidence that this is the route of virion uptake that is enhanced by PS receptors.

186 These findings are consistent with earlier reports that PS receptors mediate cargo

187 internalization into the endosomal compartment (12, 13, 29). In cells that expressed ACE2 and

188 TMPRSS2, virus entry was no longer sensitive to E-64 as previously reported (6, 27).

189 VSV/Spike entry in the presence of TMPRSS2, PS receptors, and ACE2, was also insensitive to

190 E-64, suggesting that the TMPRSS2 expression and activity mediates entry at the plasma

191 membrane independent of PS receptor utilization.

192

193 Inhibition of endogenous AXL utilization blocks SARS-CoV-2 entry

194 We next evaluated the ability of endogenously expressed PS receptors to enhance SARS-CoV-195 2 in ACE2 positive cells. Vero E6 cells that express ACE2, AXL, and TIM-1 (S4A) were initially 196 assessed (20). Notably, a large fraction of ACE2 protein is located intracellularly, suggesting a 197 rich reserve of ACE2 is inaccessible to extracellular virions (S4B). Initial studies using PS 198 liposomes confirmed that PS receptors are important for SARS-CoV-2 infection of these cells. 199 Competition studies in Vero E6 cells demonstrated that increasing doses of PS, but not PC. 200 inhibited VSV/Spike infection, similar to our findings in HEK 293T cells (Fig. 4A). PS liposomes 201 also significantly reduced SARS-CoV-2 binding to the surface of Vero E6 cells, implicating PS 202 receptors in attachment and subsequent entry of SARS-CoV-2 (Fig. 4B). These findings 203 reinforce the importance of either AXL, TIM-1, or both for SARS-CoV-2 entry. 204 205 To assess if AXL was important for infection of Vero E6 cells, the selective AXL kinase inhibitor,

bemcentinib, was tested for its ability to block SARS-CoV-2 infection. In a dose dependent

207 manner, bemcentinib profoundly inhibited SARS-CoV-2 virus load and blocked infection of the

208 one-hit VSV/Spike pseudovirion (**Fig. 4C, S4D**). A time-of-addition study indicated that

209 bemcentinib inhibition was most effective at early timepoints during SARS-CoV-2 infection,

consistent with a role of AXL in virus entry (Fig. 4D). Bemcentinib toxicity was tested on human

lung epithelial cells and was nontoxic at the concentrations used (**S4C**). Consistent with an

212 important role for AXL in SARS-CoV-2 infection, RNAseq of infected Vero E6 demonstrated that

213 infection of these cells with a MOI of 0.01 resulted in ~80% of transcripts composed of viral

transcripts 18 hpi (**Fig. 4E**). If these cells were treated with 1 μ M bemcentinib at the time of

infection, the fraction of viral transcripts dropped precipitously, decreasing to ~10% of the totalreads.

217

218 To determine if TIM-1 contributed to SARS-CoV-2 infection of Vero E6 cells, the blocking anti-219 human TIM-1 monoclonal antibody, ARD5, was evaluated for inhibition of recombinant VSV 220 (rVSV) bearing either Ebola GP (EBOV) or spike. While rVSV/EBOV GP was inhibited by ARD5 221 as previously reported (12, 30), it had no effect on rVSV/Spike infection (S4E). Thus, despite 222 robust expression of both PS receptors, AXL is preferentially utilized for SARS-CoV-2 infection 223 in these cells. While preferential PS receptor utilization has been reported for other pathogens 224 (20), our previous studies indicated that TIM-1 rather than AXL was preferentially used, in 225 contrast to our current observations with SARS-CoV-2. Host factors or virion attributes

- 226 determining PS receptor preference are currently unexplored. A broad-spectrum TAM inhibitor,
- BMS777607, modestly reduced virus infection in a dose dependent manner (Fig. 4F),
- discounting the likelihood of off-target effects with bemcentinib.
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230 As the bulk of ACE2 in Vero E6 cells is intracellular (S4B), surface expressed-AXL may be 231 facilitating SARS-CoV-2 uptake into the endosomal compartment where proteolytic processing 232 and ACE2 interactions mediate fusion of the viral envelope and cellular membranes. Previous 233 studies with the betacoronavirus responsible for the 2003-2004 outbreak. SARS-CoV. 234 demonstrated that ACE2 is found abundantly in the endosomal compartment, specifically co-235 localizing with the early endosomal marker EEA1 (31). Further, at 3 hpi, SARS-CoV antigens 236 colocalize with vesicular ACE2 and that ACE2 formed notable vesicular puncta in the infected 237 cells (31). We utilized Stimulated Emission Depletion (STED) microcopy, leveraging the super 238 resolution capabilities of this platform to investigate ACE2 and AXL colocalization in uninfected 239 and infected Vero E6 cells. In uninfected cells, AXL and ACE2 were found on the plasma 240 membrane and intracellularly, but colocalize poorly (Fig. 4G and H). However, as shown in the 241 micrographs (white arrows) and the associated fluorescence intensity plot profiles (S4F yellow 242 lines highlight selected ROI), ACE2 and AXL demonstrate overlapping localization patterns 243 within cytoplasmic punctate structures. Pearson's correlation coefficients of infected and 244 uninfected cells calculated for AXL and ACE2 intensity demonstrate a significant increase in 245 colocalization values between this PS receptor and the cognate SARS-CoV-2 receptor in 246 infected cells, relative to mock counterparts. (Fig. 4H). These data support the possibility that 247 PS receptors enhance SARS-CoV-2 trafficking into these intracellular puncta where ACE2 is 248 abundant.

249

250 AXL promotes SARS-CoV-2 infection in a range of lung cell lines

- In addition to ACE2, many lung cell lines express AXL (19, 32) (S5A). We evaluated these lines
- 252 for their ability to support SARS-CoV-2 infection and whether infection was sensitive to
- bemcentinib inhibition. The panel of lung cells that were selected included A549
- 254 (adenocarcinoma) stably expressing ACE2, HCC1650 (NSCLC), HCC1944 (squamous), H1819
- 255 (adenocarcinoma), H2302 (adenocarcinoma) and Calu3 (adenocarcinoma).
- 256
- 257 These cells were inoculated with SARS-CoV-2 (MOI = 0.5 unless otherwise noted) in the
- 258 presence or absence of a serial dilution of the AXL inhibitor, bemcentinib, or the cysteine
- protease inhibitor, E-64. The cell lines A549^{ACE2}, H1650, HCC1944, H1819, and HCC2302

260 readily supported SARS-CoV-2 infection, and viral loads 24 hpi were decreased in a dose-261 dependent manner, by bemcentinib and E64 (Fig. 5A-E). Infectious SARS-CoV-2 present in 262 HCC2302 cell supernatants at 24 and 48 hpi were also markedly decreased by bemcentinib 263 (Fig. 5F), demonstrating that bemcentinib treatment reduced production of new infectious virus 264 in a dose dependent manner. Further, at 1 µM of bemcentinib, detectable production of any 265 infectious virus was delayed until 48 hours (L.O.D. = $5 \text{ TCID}_{50}/\text{mL}$). In H1650 cells, the ability of 266 bemcentinib to inhibit recently emerged SARS-CoV-2 variants of concern (VOC), Alpha 267 (B.1.1.7) and Beta (B.1.351), was evaluated. While the Alpha VOC replicated poorly in these 268 cells, bemcentinib significantly inhibited virus replication of both variants, providing evidence the 269 efficacy of the AXL inhibitor is not influenced by SARS-CoV-2 adaptative changes (S5B). 270 271 SARS-CoV-2 infection of TMPRSS2⁺ Calu-3 cells (S6A) was not sensitive to bemcentinib or E-272 64, again providing evidence that, in this cell line, the route of virus entry was dominated by the 273 TMPRSS2-dependent path, bypassing the use of PS receptors and the endosomal 274 compartment (Fig. 5G). These findings stand in contrast to SARS-CoV-2 infection of 275 TMPRSS2⁺ H1650 cells that were markedly bemcentinib and E-64 sensitive and were found to 276 be insensitive to camostat inhibition (Fig. 5B, S5C). The paradoxical finding that virus entry into 277 H1650 is sensitive to E64 and bemcentinib despite endogenous TMPRSS2 expression indicates 278 that TMPRSS2-dependent pathways are not always the dominating or default route of SARS-279 CoV-2 entry and suggests that a more complex balance of events controls which pathway is 280 used. Neither the total amount of cell surface expressed ACE2 nor the intracellular versus 281 extracellular ACE2 ratio appears to determine the route of virus uptake (S5D). 282 283 RNA sequencing studies confirmed and extended our findings with bemcentinib in A549^{ACE2} cells. At 24 hpi, 20% of the transcripts in A549^{ACE2} cells mapped to the viral genome. Infection in 284

the presence of 1 μM bemcentinib significantly decreased the number of viral transcripts present
 (Fig. 5H). Further analysis of potential qualitative changes in viral transcripts indicated that

transcript numbers across the genome were reduced, rather than a reduction of specific

288 subgenomic transcripts.

289

To directly evaluate the importance of AXL during infection of human lung cells, CRISPR-Cas9
 technology was used to knock out (KO) AXL expression in H1650 and HCC2302 cells. H1650
 AXL^{neg}, a biologically cloned AXL-null line, was evaluated along with bulk AXL KO populations
 of H1650 and HCC2302, denoted as AXL^{low}. The AXL^{neg} clone, which expressed undetectable

levels of AXL protein (Fig. 6A), supported dramatically lower SARS-CoV-2 virus loads at a

- range of input MOIs (Fig. 6B) and became unresponsive to bemcentinib (Fig. 6C),
- 296 demonstrating an important role for AXL in SARS-CoV-2 infection and indicating that
- 297 bemcentinib specifically targets AXL. Bulk populations of AXL^{low} H1650 and HCC2302 also
- 298 supported reduced levels of SARS-CoV-2 infection and were poorly responsive to bemcentinib
- 299 (S6A-E). Taken together, the data presented here demonstrate that SARS-CoV-2 utilizes AXL
- to enhance virion binding and entry in some lung cell lines, and that this mechanism can be
- effectively disrupted in human lung cells by small molecule inhibitors and genetic ablation ofAXL.
- 303

304 AXL facilitates infection of other betacoronaviruses

305 To assess the role of AXL in the related betacoronavirus, mouse hepatitis virus (MHV strain 306 A59), we investigated the ability of bemcentinib to inhibit infection in C57BL/6J mouse bone 307 marrow derived macrophages (BMDMs) that express AXL (33-35). As MHV uses the mouse 308 receptor CEACAM as its cognate receptor, this model allowed us to examine the role of AXL 309 with a coronavirus that is ACE2-independent (36). Bemcentinib added to BMDM cultures 310 decreased virus load at 24 hours in a dose-dependent and an MOI-dependent manner (S7A-B). 311 At higher MOIs the effect of bemcentinib was diminished. Bemcentinib treatment also inhibited 312 MHV infection in peritoneal macrophages, an MHV permissive population phenotypically and 313 functionally distinct from BMDMs (S7C). These studies provide evidence that AXL facilitates 314 infection of multiple members of this enveloped virus family, independent of the cognate 315 receptor used by the virus.

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318 **DISCUSSION**

Here, we demonstrate that PS receptors, AXL, TIM-1 and TIM-4, synergize with ACE2 to mediate SARS-CoV-2 infection of HEK 293T cells when ACE2, the cognate receptor for the virus, was expressed at low levels. PS receptors enhanced virion binding to cells in a PSdependent manner. At higher levels of ACE2 expression, a role for the PS receptors was no longer observed. Similar findings were observed for TMPRSS2-facilitated, ACE2-dependent infection, indicating that when ACE2 is expressed on the plasma membrane at high concentrations these host proteins that assist SARS-CoV-2 entry are no longer required. 327 A recent study reported that AXL mediates SARS-CoV-2 infection (19). That report suggested 328 that AXL-mediated virus entry is independent of ACE2 and that AXL binds to the N-terminal 329 domain of SARS-CoV-2 spike. Conclusions from our studies indicate that AXL and other PS 330 receptors mediate enhancement of SARS-CoV-2 infection through interactions with virion-331 associated PS in an ACE2-dependent manner. We report several lines of evidence that are 332 consistent with our contention that the PS receptors interact with virion-associated PS. First, PS 333 liposomes abrogate binding and entry in a dose dependent manner. Second, interfering with PS/PS receptor complexes by mutating the TIM-1 PS binding pocket abrogates SARS-CoV-2 334 335 infection. These data are consistent with and support the well-established mechanism of PS 336 receptor enhancement of enveloped virus infection (11-13, 37, 38). Third, we directly tested 337 whether AXL binds to purified Spike or NTD by flow cytometry and biolayer interferometry 338 assays and were unable to detect any interaction. Finally, the ability of diverse PS receptors to 339 enhance SARS-CoV-2 infection in HEK 293T cells lend support to the contention that these 340 receptors interact with PS rather than viral spike on the surface of SARS-CoV-2 to mediate 341 productive infection. Thus, we conclude that AXL does not interact with SARS-CoV-2 spike, nor 342 does it mediate virus entry unilaterally. It should be noted that this is the first example of an 343 enveloped virus that utilizes PS receptors in conjunction with low/moderate expression of a high 344 affinity surface receptor.

345

346 A previous study concluded that the related coronavirus, SARS-CoV, was not productively 347 internalized by PS receptors (11). However, with insights from our studies, an alternative 348 explanation is that PS receptor enhancement of coronavirus entry is ACE2-dependent and 349 sufficient quantities of ACE2 on the plasma membrane abrogate a role for PS receptors. Thus, 350 PS receptors only facilitate SARS-CoV-2 entry under conditions where ACE2 is expressed at 351 suboptimal levels, conditions that were likely not evaluated in the cited study but are found on 352 ACE2 expressing cells such as Vero E6 cells and many patient derived lung cell lines. As ACE2 353 expression is low within the lung, such suboptimal conditions may be highly relevant during 354 SARS-CoV-2 infection (39, 40).

355

PS receptors have previously been shown to interact with PS on the surface of other enveloped
viruses such as filoviruses, alphaviruses and flaviviruses and mediate internalization into
endosomes (11, 41). However, PS receptor-dependent entry is a mechanism that is functionally
out-competed by high-affinity viral glycoprotein-host receptor interactions, such as that of Lassa
virus with α-dystroglycan (16, 20). In the case of Lassa virus entry PS receptors seem to serve

as a backup entry mechanism, as these receptors only mediate virus internalization when the
 high affinity surface receptor, α-dystroglycan, is not expressed.

363

364 Our studies provide evidence that PS receptors enhance SARS-CoV-2 binding to cells and 365 mediate internalization into endosomes where cysteine proteases potentiate spike protein 366 triggering and subsequent fusion events. Consistent with the utilization of this uptake pathway, 367 the cysteine protease inhibitor E-64 effectively blocked ACE2 or ACE2/PS receptor entry in HEK 368 293T cells. This is also supported by our super resolution microscopy observations that AXL 369 and ACE2 colocalize during SARS-CoV-2 infection. However, in HEK 293T, the route of virus 370 entry changes upon expression of TMPRSS2; virion entry is no longer sensitive to E-64. As 371 others have reported, these findings are consistent with TMPRSS2-dependent entry dominating 372 as the route of entry when TMPRSS2 is expressed (24, 42, 43). We also investigated virus infection of a variety of lung lines that endogenously express TMPRSS2, AXL, and ACE2. While 373 374 findings with Calu-3 cells were similar to that we observed in TMPRSS2-transfected HEK 293T 375 cells, the other TMPRSS2⁺ lung lines, such as H1650 cells, remained sensitive to E-64 and the 376 AXL signaling inhibitor, bemcentinib. Differences in the ability of TMPRSS2 expression to 377 control the route of entry may be due to a fine balance of surface expression of the various 378 receptors and should be explored in more detail.

379

380 Our data indicate that AXL serves as the most important PS receptor for SARS-CoV-2 infection 381 of the TIM and TAM families and our studies with MHV implicated AXL in facilitating infection of 382 additional coronaviruses. While AXL is abundant on lung epithelial cells, it is also present in 383 many organs in the body, with the exclusion of neural tissues (32, 44). Thus, it is likely a role for 384 AXL in SARS-CoV-2 infection is not only relevant to lung cell populations, but ACE2-expressing 385 tissues suspected to be affected by COVID-19 such as the heart and kidneys (45, 46). We 386 surmise that AXL-inhibiting therapeutics could function in tandem with other antivirals, protecting 387 a number of organs from infection. Our data suggest that the efficacy of bemcentinib will persist 388 as the virus evolves, inhibiting the VOCs Alpha and Beta effectively. By targeting host proteins 389 such as AXL we dramatically reduce the potential selection for pathogen mutants that reduce or 390 abolish antiviral activity. Given that currently utilized small molecule therapeutics such as 391 remdesivir targeting viral proteins have shown limited efficacy and the benefits of antibody-rich 392 convalescent plasma is minimal, AXL inhibition by small molecule inhibitors such as 393 bemcentinib offers a novel route of attack to reduce SARS-CoV-2 entry and disease (47, 48). 394

395 The preferential utilization of AXL rather than TIM-1 by SARS-CoV-2 in Vero E6 cells was 396 unexpected. In our previously studies, other enveloped viruses that utilize PS receptors, such as 397 filoviruses, use TIM-1 preferentially when both proteins are expressed (12, 20). Further, a recent 398 study identified that the TIM-1 IgV domain that contains the PS binding pocket serves as an 399 effective inhibitor of enveloped virus infection regardless of the PS receptor utilized for virus 400 uptake (49), consistent with the good affinity the TIM-1 PS binding pocket has for PS (50). 401 Nonetheless, when TIM-1 is not present in cells and AXL is the sole PS receptor expressed, 402 AXL is used by filoviruses and flaviviruses (11, 13, 51-53). The subpar utilization of AXL 403 reported for other viruses may be due to the requirement for the adaptor protein, Gas6, to also 404 be present, expression patterns, or steric factors. As multiple proteins from a variety of different 405 PS receptors families can mediate uptake of apoptotic bodies, it is no surprise that PS receptor 406 interactions with viruses are likewise intricate. Further studies are needed to understand the 407 preferential use of AXL by SARS-CoV-2. 408

- Bemcentinib, an orally bioavailable small molecule inhibitor of AXL, is currently in Phase II trials
- 410 for non-small cell lung cancer (NSCLC) and a variety solid and hematological cancers
- 411 (ClinicalTrials.gov IDs: NCT03184571, NCT03184558). However, multiple screens have
- 412 identified bemcentinib as inhibitory to SARS-CoV-2 infection, bolstering this mechanism of entry
- 413 (54, 55). Two phase 2 clinicals trial evaluating efficacy of bemcentinib in hospitalized COVID-19
- 414 patients are ongoing, with the first recently reporting short-term efficacy results
- 415 (https://clinicaltrials.gov/ct2/show/NCT04890509). In this exploratory, open-label study
- bemcentinib was added to standard-of care (SoC) therapy to hospitalized patients (56). Though
- the primary endpoints (time to improvement by 2 points on WHO ordinal scale or time to
- discharge) showed a marginal benefit of bemcentinib treatment, there was evidence of
- 419 potentially meaningful clinical benefit in a key secondary endpoint which was avoidance of
- 420 deterioration. These interim data are promising and support further clinical investigations of this
- 421 AXL inhibitor for treating SARS-CoV-2.
- 422

The robust body of PS receptor research completed in the last decade and historical patterns of zoonotic events (Ebola virus, Zika virus, coronaviruses) suggest that future emergent viral pathogens are likely to utilize PS receptors to enhance entry and infection. The observations reported here that PS receptors are utilized by a novel pandemic coronavirus support this conclusion. This confluence of information provides insights into a new class of potential therapeutics to stem future outbreaks, namely drugs aimed at inhibiting PS receptor activity.

- 429 Further studies are required to determine the role of TIM and TAM use *in vivo*; however, our
- 430 studies demonstrate a role of PS receptors in SARS-CoV-2 infection in relevant cell populations
- and further extend the importance of PS receptors in enveloped virus entry to coronaviruses.
- 432

433 STAR METHODS

434

435 **RESOURCE AVAILABILITY**

- 436 Lead contact:
- 437 Further information and request for resources and reagents should be directed to and will be
- 438 fulfilled by the lead contact Wendy J. Maury (<u>wendy-maury@uiowa.edu</u>).
- 439
- 440 Materials availability:
- 441 This study did not generate new reagents.
- 442

443 EXPERIMENTAL MODEL AND SUBJECT DETAILS

444

Ethics statement: This study was conducted in strict accordance with the Animal Welfare Act and the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health (University of Iowa (UI) Institutional Assurance Number: #A3021-01). All animal procedures were approved by the UI Institutional Animal Care and Use Committee (IACUC) which oversees the administration of the IACUC protocols and the study was performed in accordance with the IACUC guidelines (Protocol #8011280, Filovirus glycoprotein/cellular protein interactions).

452

453 **Mice**

- 454 The mice (6-8 weeks old, female) used in these studies were obtained from the Jackson
- 455 Laboratory (C57BL6/J). The protocol (#8011280) was approved by the Institutional Animal Care
- 456 and Use Committee at the University of Iowa.
- 457

458 **Primary Cells and Immortal Cell Lines**

- 459 Bone marrow derived macrophages (BMDM) were isolated and cultured in RPMI-1640
- supplemented with 10% Fetal Bovine Serum (FBS), 0.5 µg/mL of penicillin and streptomycin
- 461 (pen/strep) and 50 ng/mL murine M-CSF. Vero E6 cells (ATCC CRL-1586), Vero TMPRSS2,
- 462 Vero E6 and HEK 293T (ATCC CRL-11268) were cultured in Dulbecco's modified Eagle's

463 medium (DMEM, GIBCO, Grand Island, NY) supplemented with 5-10% FBS and 1% 464 penicillin/streptomycin (GIBCO). Blasticidin (5 µg/mL) was added to media supporting Vero E6 TMPRSS2 cell growth. A549^{ACE2} cells were generated by transduction of A549 (ATCC CCL-465 466 185) with a codon-optimized ACE2 encoding lentivirus and selection with 10 µg/mL blasticidin. 467 Clonal populations were isolated and ACE2 expression verified by western blot. H1650, 468 HCC2302, HCC1944 and H1819 human lung lines were maintained in RPMI with 5-10% FBS 469 and pen/strep. Cell lines were periodically tested for mycoplasma contamination (E-Myco kit, 470 Bulldog Bio, Portsmouth, NH) and cured of contamination before use (Plasmocin, Invitrogen, 471 San Diego CA). Cell lines were authenticated periodically by ATCC (A549 and HEK 293T) or 472 the lab responsible for their generation (H1650, HCC2302, HCC1944, H1819, and Vero 473 TMPRSS2).

474

475 AXL knockout HCC2302 and H1650 were generated by transduction of parental cells with a 476 Cas9 encoding lentivirus (kind gift of Aloysius Klingelhutz, University of Iowa) and selection in 477 10 µg/mL blasticidin for 10 days. Then cells were transduced with an Invitrogen LentiArray 478 CRISPR gRNA lentivirus targeting AXL, and subsequently selected in puromycin at 2µg/mL for 5 days. Cells were then lifted, stained for AXL, and sorted for AXL^{low} cells at the University of 479 480 Iowa Flow Cytometry Core on a FACSAria Fusion (Becton, Dickinson and Company, Franklin Lakes, New Jersey). Bulk populations of AXL^{low} cells were used for experiments as noted, and 481 Clone #4 was generated by sorting single AXL^{low} cells into a 96 well plate. AXL expression was 482 verified by flow cytometry on a FACSVerse (Becton, Dickinson and Company). 483 484

484

485 Viruses

486 Studies used the 2019n-CoV/USA-WA-1/2020 strain of SARS-CoV-2 (Accession number:

487 MT985325.1) which was propagated on Vero E6 cells. Briefly, Vero E6 cells were inoculated

488 with an MOI of 0.001 in DMEM supplemented with 2% FCS and pen/strep. Media was removed

and refreshed 24 hpi. When cells exhibited severe cytopathic effect, generally 72 hpi, cells were

490 freeze-thawed once, transferred to a conical tube, centrifuged at 1000g for 10 minutes, and

491 supernatants were filtered through a 0.45 μm filter. Virus was sequenced via Sanger

sequencing periodically for furin cleavage site mutations (none were detected) and only low

493 passage stocks were used.

494

- 495 MHV (A59) stocks were generously provided by Dr. Stanley Perlman. Viral stocks were
- 496 generated on Vero E6 and the TCID₅₀ was determined on HeLa-mCECAM1 cells by
- 497 identification of cytopathic effect at 5 days.
- 498
- 499 Stocks of recombinant vesicular stomatitis virus that expressed SARS-CoV-2 Spike containing
- 500 the D614G mutation and nano-luciferase (rVSV/Spike) (kind gift of Dr. Melinda Brindley, Univ.
- 501 GA) were generated in either Vero E6 or Vero E6 TMPRSS2 cells. Cells were infected with a
- 502 low MOI (~0.005) of virus and input was removed after ~12 h. Upon evidence of cytopathology,
- supernatants were collected over a three-day period, filtered through a 0.45 µm filter and frozen
- at -80°C until purified. Supernatants were thawed and centrifuged overnight at 7000 x g to
- 505 concentrate the virus. The virus pellet was resuspended in endotoxin-free PBS and layered over
- a 20% sucrose/PBS cushion. Virus was pelleted through the cushion by centrifugation at
- 507 28,000 rpm in a SW60Ti rotor (Beckman). The virus pellet was resuspended in PBS and the
- 508 TCID50 was determined on Vero E6 cells.
- 509
- 510 All viral titers were determined by a modified Spearman-Karber method as previously described 511 and reported as infectious units (IU)/mL (57).
- 512

513 METHOD DETAILS

514 Inhibitors

- 515 Bemcentinib (BGB324, R428) was provided by BerGenBio ASA (Bergen, Norway) and
- 516 dissolved in DMSO for *in-vitro* studies. BMS777607 (Millipore Sigma) was dissolved in DMSO.
- 517 E-64 (Millipore Sigma) was dissolved in DMSO.
- 518

519 **RNA isolation and qRT PCR**

- 520 Total RNA for PCR was extracted from cells or tissue using TRIzol (Invitrogen, Cat# 15596018)
- 521 according to the manufacturer's protocol. Total isolated RNA (1µg) was reverse transcribed to
- 522 cDNA using the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems Cat#
- 523 4368814). The resulting cDNA was used for amplification of selected genes by real-time
- 524 quantitative PCR using Power SYBR Green Master Mix (Applied Biosystems, Cat# 4368708).
- 525 Data were collected on QuantStudio 3 and Ct values determined with the QuantStudio Data
- 526 Analysis software (Applied Biosystems). Averages from duplicate wells for each gene were
- 527 used to calculate relative abundance of transcripts relative to housekeeping genes (HPRT,
- 528 GAPDH, mouse cyclophilin) and presented as $2^{-\Delta\Delta CT}$.

529

530 VSV/Spike pseudovirus production

531 The production of SARS-CoV-2-Spike vesicular stomatitis virus (VSV/Spike) pseudovirions has 532 been described previously (Hoffmann et al., 2020). Briefly, HEK 293T cells were seeded in 10 533 cm tissue culture plates (CellTreat; Ref# 229692). After 48 hours cells were transiently 534 transfected with a SARS-CoV-2-Spike pCG1 plasmid (a kind gift from Dr. Stefan Pohlman as 535 described in Hoffmann et al., 2020) using a standard polyethyleneimine (PEI) protocol. For this 536 transfection, one tube was prepared with 16 up of plasmid diluted in 1.5 mL of OPTI-Mem 537 (Gibco; Ref# 31985-070). The second tube was prepared with PEI (1mg/mL) diluted in 1.5 mL 538 of OPTI-Mem at a concentration of 3 µl/1 µg of DNA transfected. The tubes were then 539 combined, vortexed for 10-15 seconds and left to incubate at room temperature for 15 minutes. 540 The mixture was then added dropwise to the HEK 293T cells and returned to incubator 541 overnight. Twenty-four hours after transfection the cells were infected with a stock of replication 542 insufficient VSV virions expressing firefly luciferase that were pseudotyped with Lassa virus 543 glycoprotein on the viral membrane surface. The infection was incubated for ~6 hours at 37°C, 544 was removed from the cells, and fresh media was added. Viral collection took place at 24- and 545 48-hours post-infection. Media supernatants were removed from the flasks, briefly spun down to 546 remove cellular debris (180 x g for 1 minute) and filtered through a 0.45 µm syringe-tip disk 547 PVDF filter (CellTreat; Ref# 229745). Supernatants were then concentrated by a 16-hour 548 centrifugation step at 5380 x g at 4°C. Pseudovirions were purified through a 20% sucrose 549 cushion via ultracentrifugation at 28,000 rpm for two hours at 10°C in a Beckman Coulter 550 SW60Ti rotor. Pseudovirus was then resuspended in 1x PBS and stored at -80°C. Pseudovirus 551 was titered using end point dilution on Vero E6 cells. All infections of cells using SARS-CoV-2-S 552 pseudotyped virions were conducted at a volume of virus that gave a relative light unit (RLU) of 553 roughly 100,000 - 200,000 RLU.

554

555 HEK 293T transfections and plasmids

All transfections were performed in HEK 293T cells, with a total plasmid concentration of 2 μ g. Cells were seeded into 6-well tissue culture plate (Dot Scientific; Ref# 667106) at a density of 5 x 10⁵ cells/well. Forty-eight hours after cell seeding, cells were transfected with CMV-driven expression vectors of ACE2, TIM-1, TIM-4, AXL, TYRO3, MerTK and TMPRSS2 (see plasmid details below) with a standard PEI transfection protocol. For this transfection one tube was prepared with plasmid DNA and 150 mM NaCI at a concentration of 25 μ I/1 μ g of DNA transfected. A second tube was prepared with 150 mM NaCI at a concentration of 25 μ I/1 μ g of

563 DNA transfection along with PEI (1 mg/mL) at a concentration of 3 µl/ 1 µg of DNA transfected.

Tubes were combined, vortexed vigorously for 10-15 seconds and incubated at room

temperature for 15 minutes. Mixtures were added dropwise to HEK 293T cells. For all

566 experiments using ACE2 50 ng of ACE2 plasmid were transfected, unless otherwise noted. For

567 PS receptors, 1000ng of plasmid was transfected unless otherwise noted. All transfections were

568 brought up to 2 µg total transfected DNA with a PCD3.1 empty expression vector.

569

570 ACE2 and PS receptor expression detection via flow cytometry

To detect cell surface expression of ACE and PS receptor on transfected HEK293Ts, WT
VeroE6s, and AXL knock down/out H1605 and HCC2302 clones, cells were lifted using 1x

573 Versene (GIBCO; Ref# 15040066) at 37°C for ~15 minutes and placed in 5mL polystyrene

574 round-bottom tubes (Falcon Ref# 352052). Cells were washed once with FACS buffer (1x

575 Sterile PBS, 2% FBS, 0.01% sodium azide). Cells were incubated for 30 minutes on ice with

576 primary antibodies diluted in FACS buffer against ACE or PS receptors. Primary antibodies

577 were diluted to 0.75 µg/mL in FACS buffer prior to incubation. Specific primary antibodies used

as follows: goat anti-ACE2 (R&D AF933), goat anti-AXL (R&D154), goat anti-TIM-1 (R&D 1750),

579 goat anti- Tyro3 (R&D AF859), goat anti-TIM-4 (R&D 2929), goat anti-MerTK (R&D AF891),

rabbit anti-TMPRSS2 (Abcam ab92323). Cells were washed once with FACS buffer. Cells were

then incubated with secondary antibodies at a 1:1000 dilution in FACS buffer on ice for 30

582 minutes. Secondaries used were donkey anti-goat IgG (H+L) Alexa Fluor 647 (Jackson Immuno

583 Research; Ref# 705-605-003) and donkey anti-rabbit IgG (H+L) Alexa Fluor 647 (Invitrogen;

A32733). Flowcytometry was performed on a Becton Dickinson FACS Calibur and analyzed by

585 Flow Jo.

586

587 To examine both the surface and intracellular expression of hACE2 on H1650, Calu3, HCC1944, HCC2302, A549^{ACE2} and Vero E6 cells we performed the following protocol. Briefly, 588 589 cells were staining with Fixable Viability Dye eFluor 780 (eBiosciences), goat anti-human ACE2 590 (R&D AF933) followed by secondary were donkey anti-goat IgG (H+L) Alexa Fluor 647 591 (Jackson Immuno Research; Ref# 705-605-003) To measure the intracellular expression of 592 hACE2, cells were surface stained with Fixable Viability Dye eFluor 780, fixed (PFA 4%), 593 permeabilized (1X PBS + 0.5%Tween20) and stained intracellularly using goat anti-human 594 ACE2 (R&D AF933) followed by secondary donkey anti-goat labeled with AF647. Unstained 595 cells, cells plus viability dye and cells plus secondary antibody/viability dye were included as a

control in every staining. Samples were measured on a FACSverse cytometer (BD Biosciences)and data were analyzed with Flowjo software (BD Biosciences).

598

599 VSV/SARS-CoV-2-spike pseudovirion studies with inhibitors

600 Following the transfection of HEK 293T cells, cells were incubated for 24 hours. At that time 601 cells were lifted with 0.25% Trypsin (GIBCO; Ref# 25200-056) and plated at a density of 2 X 10⁴ 602 cells/well on opaque, flat-bottomed, 96-well plates (Falcon; Ref# 353296). Each transfection 603 was plated into at least 3 wells to create experimental replicates. Cells were incubated for an 604 additional 24 hours. At that time, cells were infected with VSV-luciferase/SARS-CoV-2 Spike. 605 Cells were incubated for an additional 24 hours. For experiments done with inhibitors, cells were 606 treated with the concentrations of inhibitors noted in the figure panel immediately prior to being 607 infected with pseudotyped virions. After 24 h, virus-containing media was removed and replaced 608 with 35 µl of 1x Passive lysis buffer (Promega; Ref# E194A). Plates underwent three freeze-609 thaw cycles consisting of freezing on dry-ice for 15 minutes followed by thawing at 37°C for 15 610 minutes. We followed the protocol for measuring firefly luciferase as reported previously 611 (Johnson et al., 2017). For this method 100 µl of luciferin buffer (100 µl of luciferin buffer (15 612 mM MgSO₄, 15mM KPO₄ [pH7.8], 1 mM ATP, and 1mM dithiothreitol) and 50 µl of 1mM d-613 luciferin potassium salt (Syd Laboratories; Ref# MB000102-R70170)) were added to each well 614 and luminescence was read via Synergy H1 Hybrid reader (BioTek Instruments). Relative 615 luminescence units were read out. Results analyzed by normalizing values to mock transfection

- 616 with no protease inhibitors.
- 617

618 HEK 293T SARS-CoV-2 infection studies

619 Following PEI transfection with plasmids as described in the previous section, cells were lifted

- 620 with 0.25% trypsin and plated into 48-well plates at a density of 6 x 10⁴ (Dot Scientific; Ref#
- 621 667148). In our BSL3 facility, transfected HEK 293T cells were infected at a MOI = 0.5 with
- 622 SARS-CoV-2. Cells were incubated at 37°C or 24 hours and then treated with TRIzol RNA
- 623 isolation reagent and removed from the BSL3 facility. RNA extraction and cDNA generation
- 624 proceeded as described. qRT-PCR was conducted on the cDNA using SARS-CoV-2-Spike and
- 625 GAPDH primers. Data analyzed using the $\Delta\Delta$ Ct method as described above.
- 626

627 Virion binding assays

- 628 For Vero E6 binding studies, cells were grown to confluence in 48 well plates. Media was
- 629 replaced with DMEM supplemented with 2% FBS and the indicated compounds. Cells were

630 incubated at 10°C (preventing internalization and entry) until equilibrated and SARS-CoV-2 was 631 added at MOI 5. Plates were returned to 10°C for 1 hour. Media was then removed and cells 632 were washed three times with cold DPBS (GIBCO, Cat# 14190144), removing any unbound 633 virus. Then 0.05% Trypsin-EDTA (GIBCO, Cat# 25300054) was added to control wells for 5 634 minutes at 37°C, and washed. After the final wash, all media was removed and replaced with 635 TRIzol (Invitrogen, Cat# NCC1701D). RNA was isolated and analyzed as described. Binding 636 studies in HEK 293T cells were performed using rVSV-SARS-CoV-2 Spike virions (rVSV/Spike) 637 and binding was performed at room temperature in IMMULON 2HB flat bottom plates (Thermo 638 Scientific, Waltham, MA).

639

640 **RNA sequencing and analysis**

641 Following indicated treatments and infections that were performed in triplicate or quadruplicate, 642 Vero E6 and A549^{ACE2} cells in 6 well formats were homogenized using QIAShredder tubes and 643 total RNA isolated using the RNEasy kit with DNase treatment (Qiagen). High quality RNA 644 samples that were verified by a Bioanalyzer (Agilent) was guantified and used as input to 645 generate mRNA-seg libraries for the Illumina platform. Paired-end reads were performed at the 646 Paired-end sequencing reads were subject to alignment to suitable reference genomes: human 647 GRCh38 (GCA 000001405.15 - A549 cells), green monkey (Chlsab1/GCA 000409795.2 - Vero 648 E6 cells) and SARS-CoV-2 (MN985325 - both A549 and Vero E6 cells). Alignments to human 649 and monkey genomes were performed using hisat2 v2.0.5, while to viral genome using bowtie2 650 v2.2.9. Aligned reads were counted using featureCounts from subread package v1.5.2. Counted 651 reads were normalized in R, using DESeg2 v1.30.0 and subjected to statistical analysis. The 652 statistical analysis included computation of median based fold changes, Student t-test p values 653 and false discovery rate (multiple testing correction).

654

655 **Purified spike protein flow cytometry binding studies**

The NTD-Fc and RBD-Fc constructs were kindly provided by Tom Gallagher. They contain the

- 657 Fc region of human IgG1 fused to the N-terminal domain of SARS-CoV-2 Spike (residues 1-
- 658 309) or the RBD-containing C-terminal domain of the S1 subunit (residues 310-529). We also
- generated an Fc-Spike construct that contains the Fc region of human IgG1 fused to a
- 660 cleavage-negative form of the Spike ectodomain (subunits S1 and S2, corresponding to
- positions 1-1274). To eliminate the polybasic furin cleavage site of Spike, we replaced the Arg-
- 662 Arg-Ala-Arg motif at positions 682-685 with Ser-Ser-Ala-Ser. All proteins were produced by
- transient transfection of 293T cells using polyethylenimine. Proteins were harvested in 293S

ProCDM and purified using Protein A beads. Eluted products were dialyzed against phosphate
buffered saline (pH 7.4). All proteins were analyzed by SDS-PAGE and gels were silver stained
to verify the purity of the eluted product.

667

668 We measured the binding efficiency of anti-NTD antibody AM121 (Acro Biosystems) to the 669 Spike-based constructs using an enzyme-linked immunosorbent assay (ELISA), as previously 670 described (58, 59). For this purpose, the NTD-Fc, RBD-Fc or Spike-Fc suspended in PBS were 671 attached to 96-well protein-binding plates by incubation at isomolar concentrations (2, 1,37 and 672 5 µg/mL of the probes, respectively). The next day, wells were washed once with buffer 673 containing 140 mM NaCl, 1.8 mM CaCl2, 1 mM MqCl2, 25 mM Tris pH 7.5, 20 mg/mL BSA and 674 1.1% low-fat milk. The anti-NTD antibody suspended in the same buffer was then added to the 675 wells at 0.5 µg/mL. Binding of the anti-NTD antibody was detected using a goat anti-human 676 kappa light chain conjugated to horseradish peroxidase (HPR) (BioRad). To normalize for the 677 amount of the bound probes, we also quantified the amount of probe bound to the wells by 678 incubation with an HRP-conjugated goat anti-human antibody preparation. Binding of the HRP-679 conjugated antibodies was measured by luminescence using SuperSignal West Pico enhanced 680 chemiluminescence reagents and a Synergy H1 microplate reader, as previously described 681 (60).

682

683 To determine binding of the above probes to AXL, we used flow cytometry. Briefly, HEK 293T 684 cells were seeded in 6-well plates (8.5E5 cells per well) and transfected the next day with 1.5 µg 685 of empty vector or plasmids that express AXL or the full-length form of human angiotensin 686 converting enzyme 2 (ACE2) using JetPrime transfection reagent (PolyPlus). Three days after 687 transfection, cells were detached using PBS supplemented with 7.5 mM EDTA and washed 688 once with washing buffer (PBS supplemented with 5% newborn calf serum). Cells were then 689 incubated with the NTD-Fc, RBD-Fc or Spike-Fc probes (at 5 µg/mL) or anti-AXL antibody (at 690 $0.75 \,\mu\text{g/mL}$) in the same buffer for one hour on ice and were washed four times with washing buffer. To detect binding of the Fc probes, we used a goat anti-human polyclonal antibody 691 692 preparation conjugated to Alexa 647. To detect binding of the anti-AXL antibody, we used a 693 goat anti-donkey polyclonal antibody preparation conjugated to Alexa 594. Secondary 694 antibodies were added at a 1:500 dilution and incubated with the cells on ice for one hour. Cells 695 were then washed and analyzed by flow cytometry. Staining was evaluated on a Becton 696 Dickinson FACS Calibur and analyzed by Flow Jo.

697

698 Biolayer Interferometry

- 699 Biolayer interferometry was performed on an OctetRed96 (Pall Forte-Bio, USA) using NiNTA-700 (Forte-Bio 185101) or Streptavidin- (Forte-Bio 18-5019) coated Dip and Read biosensors for 701 immobilisation of His-tagged or biotin-tagged proteins respectively. All samples were diluted in 702 Kinetic Buffer (0.1% BSA, 0.02% Tween 20, 0.05% Sodium azide in PBS). Biosensors were 703 equilibrated in Kinetic Buffer, and protein (His-tagged SARS-Cov-2 S1 protein NTD, Acro 704 Biosystems, S1D-C52H6; irrelevant control His-tagged β 4 integrin fibronectin type III domain, 705 denerous gift from Petri Kursala laboratory: biotinylated tilvestamab anti-AXL antibody. 706 BerGenBio, Norway) at a concentration of 0.7µM was loaded for 10 minutes. A baseline was 707 taken for 2 minutes in Kinetic Buffer before testing association of the target binding protein for 708 10 minutes (AXL extracellular domain fused to human IgG Fc region, BerGenBio ASA, 2µM; 709 positive control Anti-SARS-Cov2 spike NTD Neutralizing antibody, Acro Biosystems, AM121, 0.1µM; IqG1 isotype control antibody, BioXcell, BE0297, 0.1µM). Results were analyzed in 710 711 Prism 9.1.1 for MacOS (GraphPad Software, USA) by aligning to the baseline values 712 immediately prior to association and subtracting signal from isotype control.
- 713

714 STED sample preparation and image acquisition

12mm #1.5 coverslips (Fisher Scientific; Ref# 12-545-81P) were coated with 0.1% bovine
Achilles' tendon collagen diluted in 1x sterile PBS. Collaged solution was incubated at 37°C for
2 hours before being plated to 12mm coverslips. Collagen was allowed to incubate on
coverslips 12 hours at 37°C. Slips were then rinsed with 1x PBS and dried in 37°C incubator.
Slips were stored in sterile 1x PBS until use.

720

Vero E6 cells were plated onto collaged coated slips at 30,000 cells/slip. 24 hours after plating, cells were either left uninfected, or infected with SARS-CoV02 (WA-1) at an MOI = 0.01 and incubated for 24 hours. At that time cells were washed once with sterile 1x PBS and then fixed with 4% PFA solution (Electron Microscopy Sciences; Ref# 15710) for 10 minutes at room temperature. Following PFA fixation, cells were washed three times with 1x PBS and stored at 4°C until use.

727

For immunofluorescent staining, coverslips were incubated for 2 hours at RT with a blocking

- buffer consisting of 1% Triton X-100, 0.5% sodium deoxycholate, 1% egg albumin and 0.01%
- 730 sodium azide all suspended in 1x PBS. After 2-hour blockade, coverslips were incubated
- 731 overnight at 4°C with primary antibodies against hACE2 (goat anti-hACE2; R&D AF933), and

732 hAXL (rabbit anti-hAXL; Cell Signaling C89E7). Cells were washed three times with 1x PBS for 733 5 minutes each. Samples with goat anti-hACE2 primary antibodies were first incubated with 734 donkey anti-goat IgG (H+L) Alexa Fluor 568 (Invitrogen; A-11057) for 1 hour at RT. Cells were 735 washed three times with 1x PBS for 5 minutes. Coverslips incubated with donkey anti-goat 736 Alexa Fluor 568 were then incubated with 5% NGS (Sigma; G9023) for one hour at RT. Cells 737 were washed three times 1x PBS then incubated for one-hour room temperature with goat anti-738 rabbit IgG (H+L) Alexa Fluor Plus 488 (Invitrogen; A32731TR). After washing three times for 5 739 minutes with 1x PBS cells were fixed to glass microscopy slides (Fisher Scientific; 12-550-15) 740 with 12ul of Prolong Glass mounting medium (Invitrogen; P36982). Coverslips cured for 48 741 hours before imaging.

742 743

744 with an HC PL APO C32 100X/1.4 oil objective lens, and LAS X software (Leica Microsystems; 745 version 3.5.5.19976) in the Central Microscopy Research Facility at the University of Iowa. 746 Excitation was performed using a white light laser set to 20% intensity. Depletion was performed 747 with a 660nm laser set at 7% intensity for Alexa Fluor 488, and 775 nm with 16.5% laser for 748 Alexa Fluor 568 fluorophore. Depletion lasers were aligned using the auto STED alignment tool 749 in the LAS X software. Laser strength and gain were adjusted to prevent pixel saturation. 750 Images collected with two-line averages and 2 frame accumulations. Post image analyses 751 include deconvolution using Huygens Professional Software and the deconvolution Wizard auto 752 functionality (Scientific Volume Imaging). Fluorescence intensity plot profiles were created using

Image acquisition was performed on the Leica SP8 3X STED confocal microscope equipped

- 753 ColorProfiler plug-in for ImageJ (Dimiter Prodanov). Pearson's correlation coefficients were
- performed by using freehand ROI selection tool in ImageJ to outline individual cells and
- performing colocalization calculations using the Colocalization Test plugin (Tony Collins et al.).
- 756 Depicted are the R values for 20 cells across five separate fields imaged from one coverslip.
- 757

758 QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical analysis was completed in GraphPad Prism v9.0.2 (GraphPad Software, San Diego, CA). Quantification of flow cytometry data was completed in FlowJo v10.7.1 (Becton, Dickinson & Company, Ashland, OR). Statistical significance was defined as p < 0.05, and denoted by a single asterisk (*). Details regarding statistical tests used and exact values of n can be found in the corresponding figure legends. All data presented is representative of n = 3 independent experiments unless otherwise noted.

765

766 AUTHOR CONTRIBUTIONS

- 767 See Supplemental Figure 8 for author contributions matrix.
- 768

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- 779
- 780 Declaration of Interests: G.G., D.M., and E.C. are employees of BerGenBio ASA, a company
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- 784
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- 789 microscopy core.
- 790
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- 795 Veteran's Administration Medical Center.
- 796

797 FIGURE LEGENDS

- 798 Figure 1: PS receptors synergize with ACE2, enhancing SARS-CoV-2 infection of HEK
- 799 **293T cells**. A) Cells transfected with expression PS receptor plasmids, AXL or TIM-1, with or

- 800 without 50 ng of ACE2 and infected 48 hours later with SARS-CoV-2 (MOI = 0.5). Viral loads
- 801 were determined 24 hours following infection. **B-C)** PS receptors, TIM-1 (**B**) and AXL (**C**),
- 802 enhance rVSV-luciferase/Spike infection at low concentrations of ACE2 are transfected. D)
- 803 Virus binding of cells transfected with PS receptor plasmids with or without 50 ng of ACE2.
- 804 rVSV/Spike was bound to transfected cells at 48 hpi and bound virus was measured via RT-
- qPCR. E) Supernatants from SARS-CoV-2 infected (MOI = 0.5) transfected HEK 293T cells
- 806 were titered 48 hpi on Vero E6-TMPRSS2 and TCID₅₀ calculated by Spearman-Karber
- 807 equation. These studies were performed with transfection of 50 ng of ACE2 plasmid. F) HEK
- 808 293T cells transfected with expression PS receptor plasmids, TYRO3 or TIM-4, with or without
- 50 ng of ACE2 and infected 48 hours later with SARS-CoV-2 (MOI = 0.5). Viral loads were
- 810 determined 24 hours following infection.
- Data shown are pooled from at least 3 independent experiments (A, B, C, D, E, F). Data
- represented as means ± SEM. Student's t-test (A,E) and multiple t-test (B,C), One-Way ANOVA
- 813 with multiple comparisons (**D&F**); asterisks represent p < 0.05.
- Figure 2: PS receptors interact with SARS-CoV-2 by binding to PS. A-B) PS liposomes
- 815 interfere with rVSV-luciferase/Spike infection. HEK 293T cells transfected with TIM-1 plasmid
- and 50 ng of ACE2 plasmid (**A**) or AXL plasmid and 50 ng of ACE2 plasmid (**B**) were infected
- 817 with rVSV-luciferase/Spike in the presence of increasing concentrations of PS or PC liposomes
- and assessed for luciferase activity at 24 hours following infection. **C)** HEK 293T cells were
- transfected with WT or PS binding pocket mutant TIM-1 plasmids with or without 50 ng of ACE2
- 820 expressing plasmid and infected 48 hours later with rVSV-luciferase/Spike pseudovirions.
- 821 Luminescence fold change were compared to mock transfected lysates that were set to a value
- of 1. D) AXL is unable to directly interact with purified, soluble SARS-CoV-2 spike/Fc. HEK 293T
- cells transfected with AXL or ACE2 were incubated with soluble spike protein (S1/S2)-Fc, S1
- 824 RBD-Fc or S1 NTD-Fc and subsequently incubated with an Alexa 647 secondary. Spike protein
- binding was detected by flow cytometry. **E)** AXL does not bind to the NTD of SARS-CoV-2
- spike. Biolayer interferometry association curves show that immobilized AXL-Fc fails to interact
- 827 with purified NTD of spike.
- 828 Data are pooled from at least 3 independent experiments (A, B) or are representative of at least
- 829 3 experiments (**C**, **D**, **E**). Data represented as means ± SEM. Multiple t-test (**A**, **B**), One-way
- ANOVA with multiple comparisons (**C**); asterisks represent p < 0.05.
- 831 Figure 3: The route of SARS-CoV-2 entry is altered by TMPRSS2 expression. A) HEK 293T
- cells were transfected with ACE2 and TMPRSS2 as noted and infected at 48 h with VSV-
- 833 luciferase/Spike. At 24 hpi, luminescence activity was determined. Findings are shown relative

- to empty vector (Mock) transfected cells. Panel depicts one representative experiment. Students
- t-tests. **B)** TMPRSS2 expression enhances rVSV-luciferase/Spike entry at low levels of ACE2
- 836 expression. HEK 293T cells were transfected as indicated and pseudovirion entry assessed by
- 837 measuring luminescence activity at 24 hpi. C) Transfected HEK 293T cells were transfected and
- 838 infected with VSV-luciferase/Spike at 48 h in the presence or absence of E-64 (300 μM).
- 839 Luciferase activity was determined 24 hpi.
- Data are pooled from at least 3 independent experiments (**B**, **C**) or are representative of at least
- 3 experiments (A). Data represented as means ± SEM. Student's T-tests (A) Multiple t-tests (B),
- Two-way ANOVA with row-wise multiple comparisons (**C**); asterisks represent p < 0.05.
- 843 Figure 4: AXL has a prominent role in SARS-CoV-2 entry in Vero E6 cells. A) PS liposomes
- interfere with SARS-CoV-2 pseudovirion entry. Vero E6 cells were treated with PS or PC
- 845 liposomes and incubated with VSV-GFP/Spike pseudovirions for 24 hours. Entry was detected
- by GFP fluorescence. **B)** PS liposomes disrupt SARS-CoV-2 binding. Vero E6 cells were
- incubated with SARS-CoV-2 (MOI = 5) at 10°C for 1 hour, washed extensively, and viral load
- assessed by RT-qPCR. **C)** AXL signaling inhibitor bemcentinib inhibits SARS-CoV-2 infection in
- Vero E6 cells. Cells were treated with bemcentinib and infected with SARS-CoV-2 (MOI = 0.01).
- Viral loads were measured 24 hpi by RT-qPCR. **D)** Bemcentinib inhibition of SARS-CoV-2
- 851 infection is most efficacious at early time points during infection. Vero E6 cells were challenged
- with SARS-CoV-2 (MOI = 0.01) and treated with either the vehicle control or 1 μ M bemcentinib
- 853 at the indicated time. Viral loads were measured 24 hpi by RT-qPCR. E) Vero E6 cells were
- treated with 1 μM bemcentinib, infected with SARS-CoV-2 (MOI = 0.01) and mRNA harvested
- 18 hpi. mRNA was deep sequenced on an Illumina platform, and viral loads were calculated by
- alignment to the SARS-CoV-2 genome. F) Broad spectrum TAM inhibitor BMS-777607 inhibits
- 857 SARS-CoV-2 infection in Vero E6 cells. Cells were treated with inhibitor at indicated
- 858 concentrations for 1 hour, challenged (MOI = 0.01), and viral loads measured 24 hpi by RT-
- qPCR **G**) STED micrographs showing staining for ACE2 (red) and AXL (green) and merged in
- 860 Vero E6 cells. Insets are enlarged images from regions highlighted by yellow rectangles. White
- arrows indicate shared vesicular structures between the two channels. Yellow arrowheads
- indicate objects that are only seen in one channel. Plot profiles are shown in **S4F**, representing
- signal intensity along the yellow lines in the merged panels. **H)** Pearson's correlation coefficients
- of ACE2 and AXL were calculated for n=20 mock and infected cells (ROI determined by cell
 borders).

- Data are pooled from at least 3 independent experiments (**B**, **D**, **F**) or are representative of at
- least 3 experiments (**A**, **C**, **G**, **H**). Data are represented as means ± SEM. Multiple t-tests (**A**)
- student's t-test (**B**, **C**, **F**, **H**); asterisks represent p < 0.05.
- 869 Figure 5: AXL inhibition reduces SARS-CoV-2 infection in human lung cells. A-F) SARS-
- 870 CoV-2 infection is reduced by AXL inhibition in multiple human lung cell lines. In order:
- A549^{ACE2}, H1650, HCC1944, H1819, HCC2302, Calu3 were treated with the indicated inhibitors
- for 1 hour and challenged with SARS-CoV-2 (MOI = 0.5) for 24 hours. Viral load was assessed
- by RT-qPCR. **G)** HCC2302 cells were treated with bemcentinib at the indicated concentrations
- for 1 hour and infected with SARS-CoV-2 (MOI = 0.5). Input virus was removed 6 hpi and
- supernatant was collected at 24 and 48 hpi and titered by TCID₅₀ assays on Vero E6-TMPRSS2
- 876 cells. $TCID_{50}$ /mL was calculated by the Spearmann-Karber method. **H)** A549^{ACE2} were treated
- 877 with bemcentinib as indicated, infected with SARS-CoV-2 (MOI = 0.5) and mRNA harvested 24
- hpi. mRNA was sequenced, and viral loads calculated by alignment to the SARS-CoV-2
- 879 genome.
- 880 Data are pooled from at least 3 independent experiments (F) or are representative of at least 3
- experiments (A, B, C, D, E, G). Data represented as means ± SEM. Student's t-test; asterisks
 represent p < 0.05.
- 883 Figure 6: AXL knockout reduces viral loads and ablates inhibition by bemcentinib. A)
- 884 H1650 AXL knockout cells were generated by lentiviral transduction of Cas9 and gRNA
- targeting AXL, followed by selection, enrichment, and biological cloning. Shown are flow
- 886 cytometry histograms depicting AXL surface staining (black) and secondary only background
- 887 (grey), demonstrating loss of AXL expression. **B)** H1650 AXL^{neg} and H1650 Cas9 (parental)
- 888 lines were challenged with SARS-CoV-2 at indicated MOIs. At 24hpi, viral loads were assessed
- by RT-qPCR. **C)** H1650 parental and AXL^{neg} lines were treated with indicated concentration of
- bemcentinib for 1 hour and subsequently challenged with SARS-CoV-2 (MOI = 0.5) and viral
- 891 loads determine by RT-qPCR 24 hpi.
- Data are pooled from at least 3 independent experiments (**B**, **C**) or are representative of at least
- 3 experiments (A). Data represented as means ± SEM. Multiple t-tests; asterisks represent p <
 0.05.
- 895

896 SUPPLEMENTAL FIGURE LEGENDS

- 897 Supplemental Figure 1: PS receptors synergize with ACE2, enhancing SARS-CoV-2
- 898 infection of HEK 293T cells. A) Representative surface staining of receptors transfected into

- cells. B) Surface expression (MFI) of proteins in mock transfected (empty vector) and
- 900 transfected HEK 293T at 48 hours after transfection. Background fluorescence is shown for
- 901 secondary antibodies used in experiment (α-goat or rabbit secondaries). **C)** HEK 293T cells,
- 902 transfected PS receptors as noted with or without 250 ng of ACE2 were transduced with rVSV-
- 903 luciferase/Spike. Transduction was assessed 24 hours later via luminescence. D) Expression of
- 904 MerTK did not affect rVSV-luciferase/Spike transduction in the presence of 250 ng of
- 905 transfected ACE2 plasmid. E) Expression of ACE2, TIM-1 or AXL did not enhance infection of
- 906 VSV-luciferase/Lassa virus GP pseudovirions. HEK 293T cells were transfected with PS
- 907 receptor plasmids and 50 ng of ACE2 and infected 48 hours later. Panels C-E are shown as fold
- 908 change of luciferase activity in cell lysates relative to mock transfected lysates that were set to a909 value of 1.
- 910 Data shown are pooled from at least three independent experiments (**C**, **D**, and **E**). Data
- 911 represented as means ± SEM. One-Way ANOVA with multiple comparisons (C, D), Student's t-
- 912 test (**E**); asterisks represent p < 0.05.
- 913 Supplemental Figure 2: PS receptors interact with SARS-CoV-2 by binding to PS. A) AXL
- 914 surface expression in transfected HEK 293T cells. **B**) Soluble purified S1/S2-Fc and NTD-Fc are
- 915 detected by an NTD monoclonal antibody by ELISA. **C**) All spike-Fc proteins bind and are
- 916 detected at equivalent levels of ELISA plates.
- 917 Supplemental Figure 3: The route of SARS-CoV-2 entry is altered by TMPRSS2
- 918 **expression.** ATPLite cytotoxicity assay in H1650 cells, 24 hours following treatment with E64.
- 919 Data are represented as means +/- SEM.
- 920 Supplemental Figure 4: AXL has a prominent role in SARS-CoV-2 entry in Vero E6 cells.
- 921 A) ACE2, AXL and TIM-1 surface expression MFI in Vero E6 cells, as assessed by flow
- 922 cytometry. Background fluorescence is shown for secondary antibodies used in experiment. B)
- 923 Cell surface versus intracellular ACE2 expression in VeroE6 cells. Indicated cells were lifted,
- 924 permeabilized as noted, and stained with anti-ACE2 unconjugated primary antibodies and Alexa
- 925 647 secondaries. **C)** Bemcentinib toxicity 24 hours after treatment was measured by ATPlite
- assay in H1650 cell line. **D)** VSV-GFP/Spike entry was measured by flow cytometry 24 hours
- after challenge to Vero E6 cells treated with bemcentinib. **E)** Vero E6 were treated with ARD5
- 928 (TIM-1 blocking antibody) 1 hour before infection with rVSV /SARS-CoV-2 Spike or rVSV
- 929 /EBOV-GP (MOI = 0.01). Viral load was measured 24 hpi by RT-qPCR. F) Plot profiles of ACE2
- and AXL intensity are shown in from STED micrographs in Fig. **4G**, representing signal intensity
- along the yellow lines in the merged panels.

- Data in A, C, D and E are shown as means ± SEM. Multiple t-tests were performed in C and
- 933 Student's t-test was performed in D and E; asterisks represent p < 0.05.

934 Supplemental Figure 5: AXL inhibition reduces SARS-CoV-2 infection in human lung

- 935 cells. A) Multiple SARS-CoV-2 permissive cell lines were stained for extracellular ACE2, AXL,
- 936 TIM-1, and TMPRSS2 protein, and expression was quantified by flow cytometry. Shown are
- 937 flow cytometry histograms depicting target surface staining (black line) and secondary only
- background (grey shade) **B)** H1650 cells were treated with 1 µM of bemcentinib and infected
- 939 with one of three different variants of SARS-CoV-2: WA-1; B.1.1.7 or B.1.351 (MOI=0.5 for all
- variants). RNA was isolated at 24 hpi and assessed for virus load. C) H1650 cells were infected
- 941 with SARS-CoV-2 (MOI = 0.5) after treatment with the indicated concentration of camostat for 1
- hour. Viral loads 24hpi were measured by qRT-PCR. D) Extracellular and intracellular staining
- 943 of ACE2 are shown in multiple cell lines. Presented as frequency positive cells.
- Data represented as means ± SEM. Student's t-test; asterisks represent p < 0.05.
- 945 Supplemental Figure 6: AXL knockout reduces viral loads and ablates inhibition by
- 946 **bemcentinib. A)** H1650 and HCC2302 AXL knockout cells were generated by lentiviral
- 947 transduction of Cas9 and gRNA targeting AXL, followed by selection. These are designated
- ⁹⁴⁸ "Bulk AXL^{low}" Shown are flow cytometry histograms depicting AXL surface staining (black) and
- secondary only background (grey), demonstrating complete loss of AXL expression in H1650
- 950 AXL^{neg}. **B)** H1650 AXL^{low} and H1650 Cas9 (parental) lines were challenged with SARS-CoV-2 at
- 951 indicated MOIs for 24 hpi and viral loads assessed by RT-qPCR. C) H1650 parental and AXL
- 952 lines were treated with indicated concentration of bemcentinib for 1 hour and subsequently
- 953 challenged with SARS-CoV-2 (MOI = 0.5) and viral loads determine by RT-qPCR 24 hpi. **D-E**)
- 954 As in B-C with HCC2302 cells.
- Data are pooled from at least 3 independent experiments (**B**, **C**, **D**, **E**) or are representative of at
- least 3 experiments (A). Data represented as means ± SEM. Multiple t-tests were performed;
- 957 asterisks represent p < 0.05.
- 958 **Supplemental Figure 7: A)** Bone marrow derived macrophages from C57bl6/J mice were
- 959 treated as indicated for 1 hour, challenged with MHV (stain A59) at the indicated MOI. Viral
- 960 loads were assessed 24 hpi by RT-qPCR. B) BMDMs were treated with indicated
- 961 concentrations of bemcentinib for 1 hour, infected with MHV (MOI = 0.001) for 24 hours and
- viral load assessed by RT-qPCR. C) As in B, MHV infection of peritoneal macrophages (MOI
- 963 =0.001) treated with indicated concentrations of bemcentinib.

- 964 Data shown are representative of 3 independent experiments. Data represented as means ±
- 965 SEM. Student's t-test; asterisks represent p < 0.05.
- 966

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Figure 1: PS receptors synergize with ACE2, enhancing SARS-CoV-2 infection.

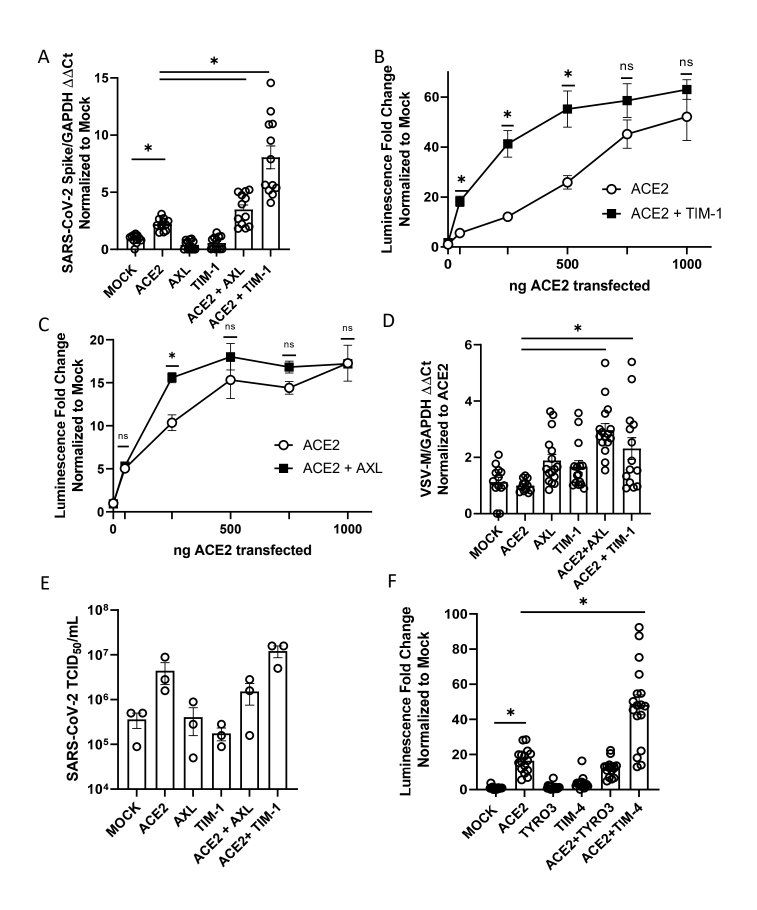


Figure 1: PS receptors synergize with ACE2, enhancing SARS-CoV-2 infection of HEK 293T cells. A) Cells transfected with expression PS receptor plasmids, AXL or TIM-1, with or without 50 ng of ACE2 and infected 48 hours later with SARS-CoV-2 (MOI = 0.5). Viral loads were determined 24 hours following infection. B-C) PS receptors, TIM-1 (B) and AXL (C), enhance rVSV-luciferase/Spike infection at low concentrations of ACE2 are transfected. D) Virus binding of cells transfected with PS receptor plasmids with or without 50 ng of ACE2. rVSV/Spike was bound to transfected cells at 48 hpi and bound virus was measured via RT-qPCR. E) Supernatants from SARS-CoV-2 infected (MOI = 0.5) transfected HEK 293T cells were titered 48 hpi on Vero E6-TMPRSS2 and TCID₅₀ calculated by Spearman-Karber equation. These studies were performed with transfection of 50 ng of ACE2 plasmid. F) HEK 293T cells transfected with expression PS receptor plasmids, TYRO3 or TIM-4, with or without 50 ng of ACE2 and infected 48 hours later with SARS-CoV-2 (MOI = 0.5). Viral loads were determined 24 hours following infection.

Data shown are pooled from at least 3 independent experiments (A, B, C, D, E, F). Data represented as means ± SEM. Student's t-test (A,E) and multiple t-test (B,C), One-Way ANOVA with multiple comparisons (D&F); asterisks represent p < 0.05.



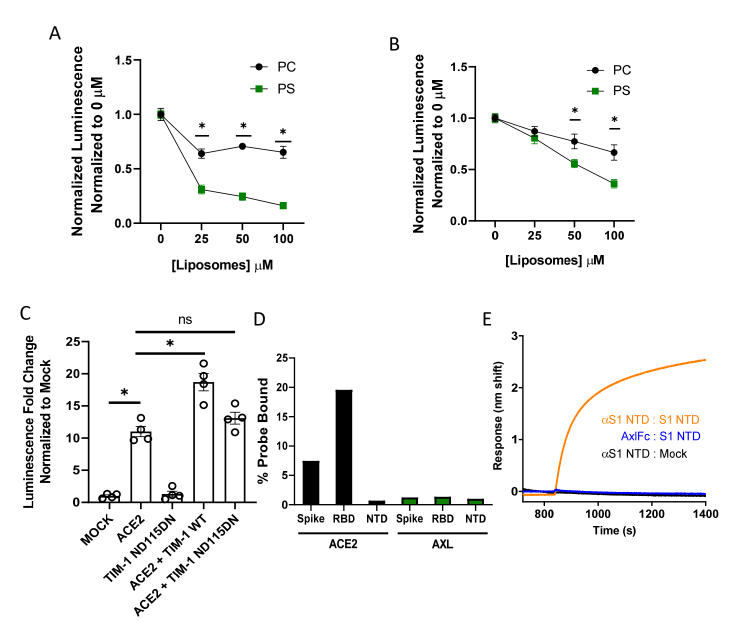


Figure 2: PS receptors interact with SARS-CoV-2 by binding to PS. A-B) PS liposomes interfere with rVSV-luciferase/Spike infection. HEK 293T cells transfected with TIM-1 plasmid and 50 ng of ACE2 plasmid (**A**) or AXL plasmid and 50 ng of ACE2 plasmid (**B**) were infected with rVSV-luciferase/Spike in the presence of increasing concentrations of PS or PC liposomes and assessed for luciferase activity at 24 hours following infection. **C)** HEK 293T cells were transfected with WT or PS binding pocket mutant TIM-1 plasmids with or without 50 ng of ACE2 expressing plasmid and infected 48 hours later with rVSV-luciferase/Spike pseudovirions. Luminescence fold change were compared to mock transfected lysates that were set to a value of 1. **D)** AXL is unable to directly interact with purified, soluble SARS-CoV-2 spike/Fc. HEK 293T cells transfected with AXL or ACE2 were incubated with soluble spike protein (S1/S2)-Fc, S1 RBD-Fc or S1 NTD-Fc and subsequently incubated with an Alexa 647 secondary. Spike protein binding was detected by flow cytometry. **E)** AXL does not bind to the NTD of SARS-CoV-2 spike. Biolayer interferometry association curves show that immobilized AXL-Fc fails to interact with purified NTD of spike.

Data are pooled from at least 3 independent experiments (**A**, **B**) or are representative of at least 3 experiments (**C**, **D**, **E**). Data represented as means \pm SEM. Multiple t-test (**A**, **B**), One-way ANOVA with multiple comparisons (**C**); asterisks represent p < 0.05.

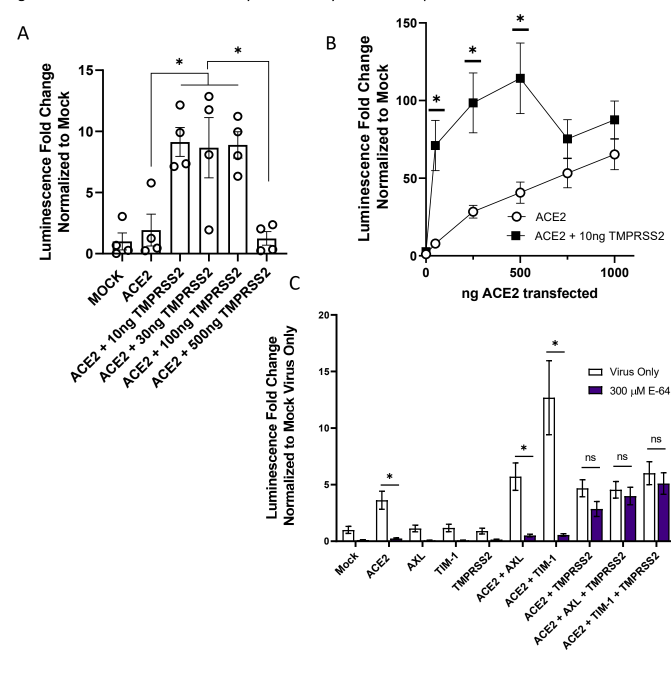


Fig. 3. The route of SARS-CoV-2 entry is altered by TMPRSS2 expression

Figure 3: The route of SARS-CoV-2 entry is altered by TMPRSS2 expression. A) HEK 293T cells were transfected with ACE2 and TMPRSS2 as noted and infected at 48 h with VSV-luciferase/Spike. At 24 hpi, luminescence activity was determined. Findings are shown relative to empty vector (Mock) transfected cells. Panel depicts one representative experiment. Students t-tests. **B)** TMPRSS2 expression enhances rVSV-luciferase/Spike entry at low levels of ACE2 expression. HEK 293T cells were transfected as indicated and pseudovirion entry assessed by measuring luminescence activity at 24 hpi. **C)** Transfected HEK 293T cells were transfected and infected with VSV-luciferase/Spike at 48 h in the presence or absence of E-64 (300 μM). Luciferase activity was determined 24 hpi.

Data are pooled from at least 3 independent experiments (**B**, **C**) or are representative of at least 3 experiments (**A**). Data represented as means \pm SEM. Student's T-tests (**A**) Multiple t-tests (**B**), Two-way ANOVA with row-wise multiple comparisons (**C**); asterisks represent p < 0.05.



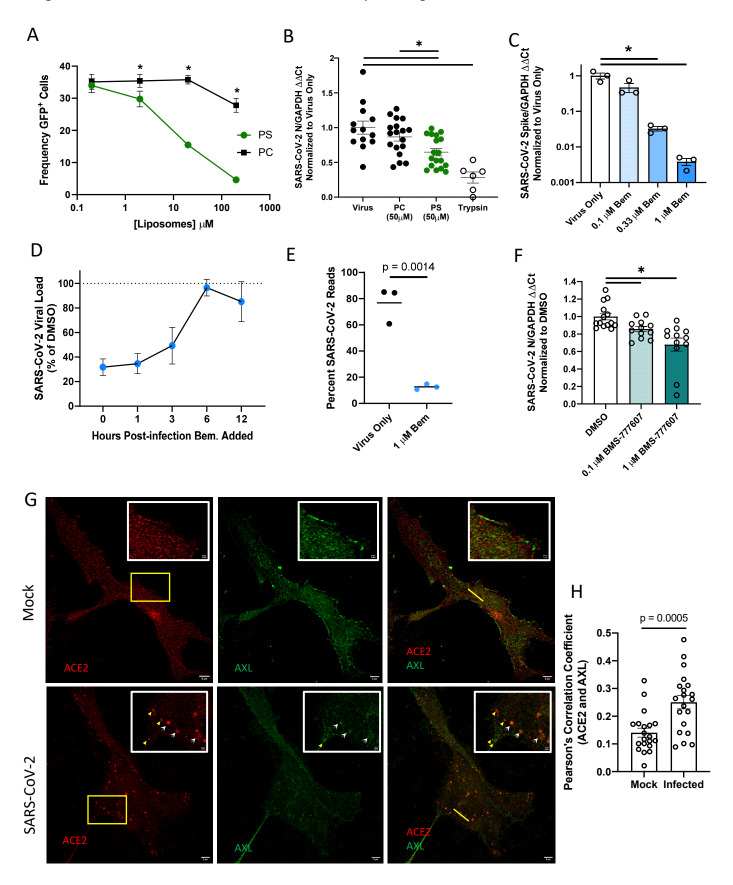
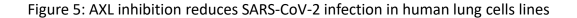


Figure 4: AXL has a prominent role in SARS-CoV-2 entry in Vero E6 cells. A) PS liposomes interfere with SARS-CoV-2 pseudovirion entry. Vero E6 cells were treated with PS or PC liposomes and incubated with VSV-GFP/Spike pseudovirions for 24 hours. Entry was detected by GFP fluorescence. B) PS liposomes disrupt SARS-CoV-2 binding. Vero E6 cells were incubated with SARS-CoV-2 (MOI = 5) at 10°C for 1 hour, washed extensively, and viral load assessed by RT-qPCR. C) AXL signaling inhibitor bemcentinib inhibits SARS-CoV-2 infection in Vero E6 cells. Cells were treated with bemcentinib and infected with SARS-CoV-2 (MOI = 0.01). Viral loads were measured 24 hpi by RT-gPCR. D) Bemcentinib inhibition of SARS-CoV-2 infection is most efficacious at early time points during infection. Vero E6 cells were challenged with SARS-CoV-2 (MOI = 0.01) and treated with either the vehicle control or $1 \mu M$ bemcentinib at the indicated time. Viral loads were measured 24 hpi by RT-gPCR. E) Vero E6 cells were treated with 1 μ M bemcentinib, infected with SARS-CoV-2 (MOI = 0.01) and mRNA harvested 18 hpi. mRNA was deep sequenced on an Illumina platform, and viral loads were calculated by alignment to the SARS-CoV-2 genome. F) Broad spectrum TAM inhibitor BMS-777607 inhibits SARS-CoV-2 infection in Vero E6 cells. Cells were treated with inhibitor at indicated concentrations for 1 hour, challenged (MOI = 0.01), and viral loads measured 24 hpi by RT-qPCR G) STED micrographs showing staining for ACE2 (red) and AXL (green) and merged in Vero E6 cells. Insets are enlarged images from regions highlighted by vellow rectangles. White arrows indicate shared vesicular structures between the two channels. Yellow arrowheads indicate objects that are only seen in one channel. Plot profiles are shown in **S4F**, representing signal intensity along the yellow lines in the merged panels. H) Pearson's correlation coefficients of ACE2 and AXL were calculated for n=20 mock and infected cells (ROI determined by cell borders).

Data are pooled from at least 3 independent experiments (**B**, **D**, **F**) or are representative of at least 3 experiments (**A**, **C**, **G**, **H**). Data are represented as means \pm SEM. Multiple t-tests (**A**) student's t-test (**B**, **C**, **F**, **H**); asterisks represent p < 0.05.



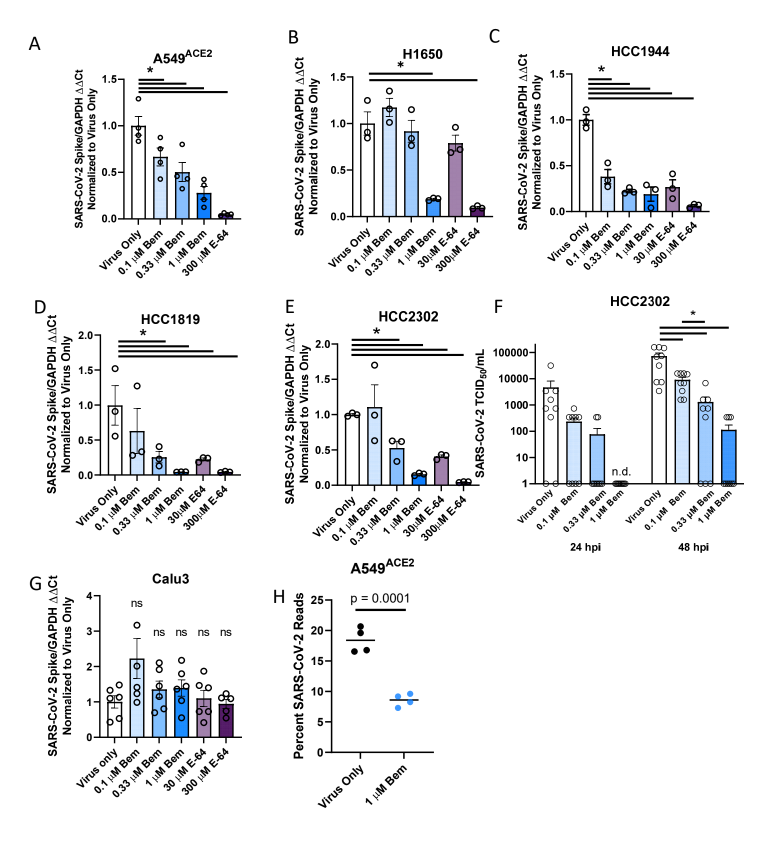


Figure 5: AXL inhibition reduces SARS-CoV-2 infection in human lung cells. A-F) SARS-CoV-2 infection is reduced by AXL inhibition in multiple human lung cell lines. In order: A549^{ACE2}, H1650, HCC1944, H1819, HCC2302, Calu3 were treated with the indicated inhibitors for 1 hour and challenged with SARS-CoV-2 (MOI = 0.5) for 24 hours. Viral load was assessed by RT-qPCR. **G)** HCC2302 cells were treated with bemcentinib at the indicated concentrations for 1 hour and infected with SARS-CoV-2 (MOI = 0.5). Input virus was removed 6 hpi and supernatant was collected at 24 and 48 hpi and titered by TCID₅₀ assays on Vero E6-TMPRSS2 cells. TCID₅₀/mL was calculated by the Spearmann-Karber method. **H)** A549^{ACE2} were treated with bemcentinib as indicated, infected with SARS-CoV-2 (MOI = 0.5) and mRNA harvested 24 hpi. mRNA was sequenced, and viral loads calculated by alignment to the SARS-CoV-2 genome.

Data are pooled from at least 3 independent experiments (**F**) or are representative of at least 3 experiments (**A**, **B**, **C**, **D**, **E**, **G**). Data represented as means \pm SEM. Student's t-test; asterisks represent p < 0.05.

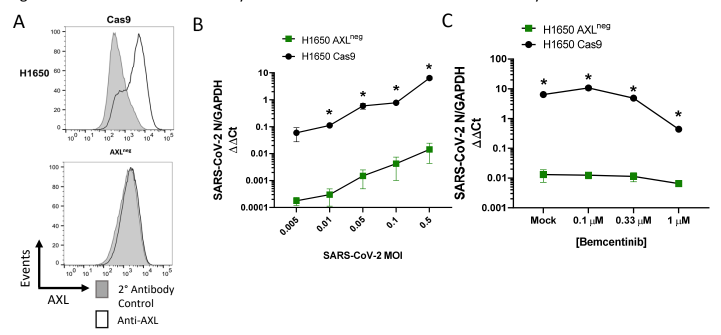


Figure 6: AXL knockout dramatically reduces viral loads and ablates inhibition by bemcentinib.

Figure 6: AXL knockout reduces viral loads and ablates inhibition by bemcentinib. A)

H1650 AXL knockout cells were generated by lentiviral transduction of Cas9 and gRNA targeting AXL, followed by selection, enrichment, and biological cloning. Shown are flow cytometry histograms depicting AXL surface staining (black) and secondary only background (grey), demonstrating loss of AXL expression. **B)** H1650 AXL^{neg} and H1650 Cas9 (parental) lines were challenged with SARS-CoV-2 at indicated MOIs. At 24hpi, viral loads were assessed by RT-qPCR. **C)** H1650 parental and AXL^{neg} lines were treated with indicated concentration of bemcentinib for 1 hour and subsequently challenged with SARS-CoV-2 (MOI = 0.5) and viral loads determine by RT-qPCR 24 hpi.

Data are pooled from at least 3 independent experiments (**B**, **C**) or are representative of at least 3 experiments (**A**). Data represented as means \pm SEM. Multiple t-tests; asterisks represent p < 0.05.