#### 1 Full Title:

# Synthetic Heparan Sulfate Mimetic Pixatimod (PG545) Potently Inhibits SARS-CoV-2 By Disrupting The Spike-ACE2 interaction

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# 6 **Short title:** Heparan Sulfate Mimetic Pixatimod Potently Inhibits SARS-CoV-2

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- 47 Summary
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Heparan sulfate (HS) is a cell surface polysaccharide recently identified as a co-receptor with the 49 ACE2 protein for recognition of the S1 spike protein on SARS-CoV-2 virus, providing a tractable 50 new target for therapeutic intervention. Clinically-used heparins demonstrate inhibitory activity, 51 but world supplies are limited, necessitating alternative solutions. Synthetic HS mimetic pixatimod 52 is a drug candidate for cancer with immunomodulatory and heparanase-inhibiting properties. Here 53 we show that pixatimod binds to and destabilizes the SARS-CoV-2 spike protein receptor binding 54 domain (S1-RBD), and directly inhibits its binding to human ACE2, consistent with molecular 55 modelling identification of multiple molecular contacts and overlapping pixatimod and ACE2 56 binding sites. Assays with multiple clinical isolates of live SARS-CoV-2 virus show that pixatimod 57 potently inhibits infection of monkey Vero E6 and human bronchial epithelial cells at 58 concentrations within its safe therapeutic dose range. Furthermore, in a K18-hACE2 mouse model 59 pixatimod demonstrates that pixatimod markedly attenuates SARS-CoV-2 viral titer and COVID-60 19-like symptoms. This demonstration of potent anti-SARS-CoV-2 activity establishes proof-of-61 concept for targeting the HS-Spike protein-ACE2 axis with synthetic HS mimetics. Together with 62 other known activities of pixatimod our data provides a strong rationale for its clinical investigation 63 as a potential multimodal therapeutic to address the COVID-19 pandemic. 64

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66 *199 words* 

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# 69 Introduction

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The coronavirus disease 2019 (COVID-19) pandemic caused by the severe acute respiratory 71 syndrome coronavirus 2 (SARS-CoV-2) has according to the World Health Organisation recently 72 surpassed 110 million cases and 2.4 million deaths world-wide. Although vaccines against COVID-73 19 are currently being developed and deployed, given the severe pathophysiology induced by 74 SARS-CoV-2 (1), there is a clear need for therapeutic alternatives to alleviate and stop the COVID-75 19 epidemic that complement vaccination campaigns. Heparan sulfate (HS) is a highly sulfated 76 glycosaminoglycan found on the surface of most mammalian cells which is used by many viruses 77 as an entry receptor or co-receptor (2), including coronaviruses (3). Various compounds that mimic 78 cellular HS such as clinically-used heparins have been investigated and have been shown to block 79 infectivity and cell-to-cell spread in a multitude of different viruses, including SARS-associated 80 81 coronavirus strain HSR1 (4). The glycosylated spike (S) protein of SARS-CoV-2 mediates host cell infection via binding to a receptor protein, angiotensin-converting enzyme 2 (ACE2) (5). Analysis 82 of the sequence and experimentally determined structures of the S protein reveals that the receptor 83 binding domain (RBD) of the S1 subunit contains a HS binding site. Recent studies have clearly 84 85 demonstrated binding of heparin and HS to S1 RBD (6-9), including induction of significant conformational change in the S1 RBD structure (6), and also revealed that HS is a co-receptor with 86 ACE2 for SARS-CoV-2 (10). Collectively these data strongly suggest that blocking these 87

interactions with heparins and HS mimetics has potential as an effective strategy for COVID-19
therapy. Although heparins have major potential for repurposing for such applications, limitations
in the global supply of natural product heparins will greatly restrict its availability (11), highlighting
an urgent need to find synthetic alternatives.

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Pixatimod (PG545) is a clinical-stage HS mimetic with potent anti-cancer (12,13), and anti-93 inflammatory properties (14). However, significant antiviral and virucidal activity for pixatimod 94 has also been reported against a number of viruses that use HS as an entry receptor with  $EC_{50}$ 's 95 ranging from 0.06 to 14 µg/mL. This includes HSV-2 (15), HIV (16), RSV (17), Ross River, 96 Barmah Forest, Asian CHIK and chikungunya viruses (18), and Dengue virus (19). In vivo efficacy 97 has been confirmed in a prophylactic mouse HSV-2 genital infection model (15), a prophylactic 98 Ross River virus mouse model (18) and a therapeutic Dengue virus mouse model (19). Pixatimod 99 has been evaluated in a Phase Ia clinical trial in patients with advanced solid tumours where it 100 demonstrated a tolerable safety profile and some evidence of disease control (13). It has been safely 101 administered to over 80 cancer patients in Phase I studies as a monotherapy or in combination with 102 nivolumab (NCT02042781 and ACTRN12617001573347), prompting us to examine its anti-viral 103 activity against SARS-CoV-2. 104

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Here we provide evidence of a direct and destabilizing interaction of pixatimod with the S1 spike 106 protein RBD, supported by molecular modelling data. Additionally, pixatimod was able to inhibit 107 the interaction of S1-RBD with ACE2 and also Vero cells which are known to express the ACE2 108 109 receptor, indicating a direct mechanism of action. We established that pixatimod is a potent inhibitor of attachment and invasion of Vero cells and human bronchial epithelial cells by multiple 110 clinical isolates of live SARS-CoV-2 virus, and reduces its cytopathic effect, at concentrations 111 within the known therapeutic range of this drug. Finally, we observed marked attenuation of SARS-112 CoV-2 viral RNA load and COVID-19-like symptoms in the K18-hACE2 mouse model of 113 infection. Our data demonstrate that synthetic HS mimetics can target the HS-Spike protein-ACE2 114 axis to inhibit SARS-CoV-2 infection. They provide strong support for clinical investigation of the 115 potential of pixatimod as a novel therapeutic intervention for prophylaxis and treatment of COVID-116 19, and have implications for wider applications against other HS-binding viruses and emerging 117 global viral threats. 118

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#### 132 **Results**

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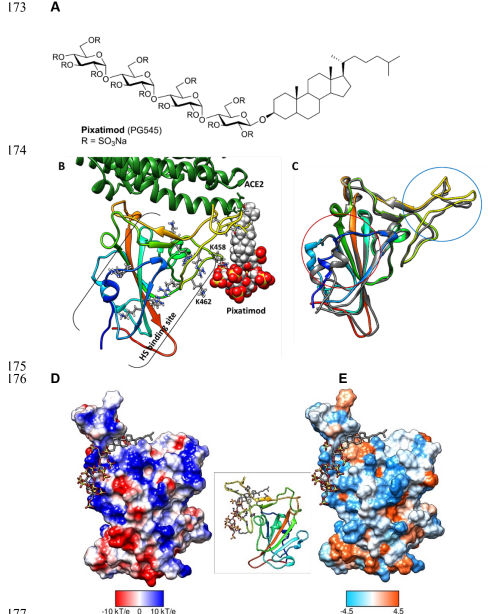
# 134 Modelling of pixatimod-spike protein interactions

We initially used molecular dynamics (MD) simulations to map the potential binding sites of 135 pixatimod (Fig 1A) on the S1 RBD surface (Fig 1B) of monomeric spike. A total of 24 unique 136 residues of RBD are known to interact with ACE2 based on the X-ray structures (Fig 1B). 137 Interestingly, a few of these residues (Tyr489, Phe456, Leu455, Ala475) are also predicted to be 138 involved in binding to pixatimod. Amino acids making significant interactions with pixatimod were 139 140 identified on the basis of their individual contributions to the total interaction energy, considering only the residues that contribute less than -1.0 kcal/mol. The decomposition approach was helpful 141 for locating residues of the RBD domain such as Lys458, Ser459, Lys462 and Asn481 that 142 transiently interact to form hydrogen bonds or ionic interactions with the sulfated tetrasaccharide 143 moiety of pixatimod (Fig 1D). The free energy of binding is -10 kcal/mol, wherein van der Waals 144 energies make the major favourable contribution to the total free energy. The cholestanol residue 145 also formed stabilizing interactions with Tyr489, Phe456, Tyr473, Ala475, Gln474 and Leu455 146 (Fig 1E). Furthermore, the standard deviation of backbone RMSD around residues Leu455-Pro491 147 and the N-terminal of RBD (Thr333-Thr345) among the four repeated MD trajectories were 148 approximately 2Å, indicating significant conformational change in the region. RMSF calculations 149 of main-chain atoms showed significant atomic fluctuations (>1.5 Å) for Lvs458, Asn460, Lvs462, 150 Arg466, Ser477 and Asn481 upon binding to the ligand pixatimod. These results indicate that a 151 conformational change may be induced by binding of pixatimod to S1 RBD (Fig 1C). 152

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An alternate heparin binding site is reported around residues Arg403, Arg406, Arg408, Gln409, Lys417, Gln493, Gln498 (8). One of the replicates indicated a second binding mode wherein the tetrasaccharide of pixatimod was found to interact around this region (Supplementary Materials, **Fig S1**), however, the free energy of binding was > +13 kcal/mol indicating much less favourable binding to this site. Overall, our modelling data strongly support the notion of direct binding of pixatimod via multiple amino acid contacts in S1 RBD, potentially resulting in induction of a conformational change and interference with binding to ACE2.

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179 180 Figure 1: Molecular dynamics modelling defines direct interactions of pixatimod with S1 RBD: A, Structure of 181 pixatimod. B, Model showing interactions of pixatimod with the RBD domain of spike protein. The sulfated tetrasaccharide partially occupies the HS/heparin binding site. The lipophilic tail of pixatimod wraps around the 182 183 hydrophobic loop, thereby creating a steric clash with the helix of ACE2 protein (shown in inset-green ribbon). C, 184 Superimposition of the X-ray structure (PDB: 6LZG) and one of the snapshots from MD simulations (ligand not shown) 185 suggest conformational change around the loop region (blue circle) and the N-terminal helix as highlighted (red circle). 186 D, coulombic surface and E, hydrophobic surface binding mode of pixatimod to S1 RBD. Both surfaces are oriented in the same direction as shown in the ribbon diagram of the protein in the middle. The sulfated tetrasaccharide interacts 187 188 with the basic regions on S1 RBD whereas cholestanol residue prefers hydrophobic region for interactions. Coulombic surface coloring defaults:  $\varepsilon = 4r$ , thresholds  $\pm 10$  kcal/mol were used. Blue indicates surface with basic region whereas 189 190 red indicates negatively charged surface. The hydrophobic surface was colored using the Kyte-Doolittle scale wherein 191 blue, white and orange red colour indicates most hydrophilicity, neutral and hydrophobic region, respectively. UCSF 192 Chimera was used for creating surfaces and rendering the images. Hydrogens are not shown for clarity.

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# 197 **Pixatimod interacts with spike protein**

Spectroscopic studies with circular dichroism (CD) were undertaken to investigate direct binding 198 of pixatimod to recombinant spike protein receptor binding domain (S1 RBD) (Fig 2A,B), the 199 region which interacts with the ACE2 receptor on human cells. CD spectroscopy in the far UV 200 201 region ( $\lambda = 190 - 260$  nm) detects conformational changes in protein secondary structure that occur in solution and can infer binding by an added ligand. Such secondary structural changes can be 202 quantified using spectral deconvolution. SARS-CoV-2 EcS1-RBD (produced in E. coli, see 203 Methods) underwent conformational change in the presence of either pixatimod or heparin as a 204 comparator sulfated molecule known to bind the RBD (6-10), consisting of decreased  $\alpha$ -helical 205 content for pixatimod and increased  $\alpha$ -helical content for heparin (Fig 2C). A decrease in global  $\beta$ -206 sheet content is observed for both pixatimod and heparin, along with increases in turn structure (Fig 207 **2C**). 208

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210 We explored the effects of pixatimod on protein stability using differential scanning fluorimetry

211 (DSF) in which the thermal denaturation of a protein is monitored in the presence of a hydrophobic

fluorescent dye (20). Binding of pixatimod induced a notably large reduction in melting temperature

213 ( $\Delta T_m$ ) of 9.1°C (**Fig 2D**; p=0.0001), indicating major destabilisation of the mammalian expressed

214 S1-RBD (mS1 RBD) protein. In contrast, heparin at an equivalent dose only partially destabilised

the RBD protein, evidenced by a small side peak shifted by  $\sim$ 5-6°C indicating populations of RBD in a bound and unbound state (**Fig 2D**).

The observed changes demonstrate that the SARS-CoV-2 S1 RBD interacts with pixatimod in aqueous conditions of physiological relevance. Notably, the conformational changes and destabilization observed were distinct for pixatimod compared to heparin, suggesting distinct interactions (**Fig 2**). Consistent with the modelling results, these data confirm direct interactions of pixatimod with S1 RBD, resulting in induction of a conformational change, consistent with the notion that HS mimetics such as pixatimod have the potential to interfere with S1-RBD interactions with ACE2.

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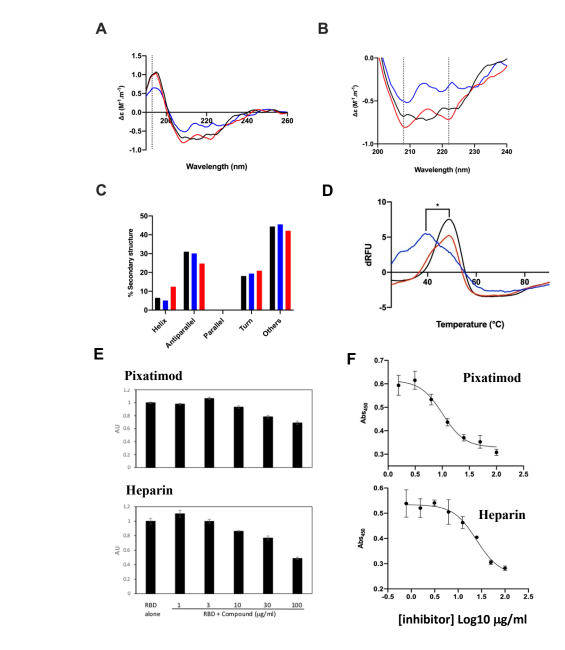


Figure 2: Pixatimod interacts with SARS-CoV-2 S1-RBD and inhibits binding to cells and ACE2 receptor. A, Circular dichroism spectra (190 - 260 nm) of SARS-CoV-2 EcS1-RBD alone (black), or with pixatimod (blue) or heparin (red). The vertical dotted line indicates 193 nm. B, The same spectra expanded between 200 and 240 nm. Vertical dotted lines indicate 222 nm and 208 nm. C, Secondary structure content analysed using BeStSel for SARS-CoV-2 EcS1-RBD.  $\alpha$ -helical secondary structure is characterized by a positive band at ~193 nm and two negative bands at ~208 and ~222 nm (analysis using BeStSel was performed on smoothed data between 190 and 260 nm). D, Differential scanning fluorimetry of binding of pixatimod (blue; 10µg) or heparin (red; 10µg) to mS1-RBD (1µg; black line, protein-only control). \*T<sub>m</sub> values for RBD alone (48.4°C, SD = 0.3) and in the presence of pixatimod (39.3°C, SD = 1) were statistically different, t(4) = 15.25, p = 0.0001. E: dose response effects of pixatimod (E) and unfractionated porcine mucosal heparin (F) on binding of EcS1-RBD to Vero cells. Data were normalised to control with no addition of EcS1-RBD. AU, arbitrary units of fluorescence. n=3 +/- CV. F, Competitive ELISA assay using biotinylated human ACE2 protein immobilized on streptavidin coated plates, to measure inhibition of binding of mS1-RBD in the presence of various concentrations of inhibitor compounds. Pixatimod (IC<sub>50</sub>, 10.1 µg/ml) and porcine mucosal heparin (IC<sub>50</sub>, 24.6  $\mu$ g/ml). n=3, +/-SD; representative example shown. 

# 260 **Pixatimod inhibits S1-RBD cell-binding**

We next evaluated inhibition of binding of His-tagged EcS1-RBD to monkey Vero cells (which are 261 known to express both HS proteoglycans (HSPGs) and the ACE2 protein receptor required for 262 SARS-CoV-2 attachment and cell invasion). Fixed cells were exposed to His-tagged S1 RBD for 263 lhr, in the presence or absence of additional compounds, with subsequent washing and detection 264 using a fluorescently-labelled anti-His tag antibody. A clear dose response was noted for both 265 pixatimod and heparin as a comparator compound (Fig 2E), with 32% and 51% inhibition achieved 266 at 100 µg/mL respectively. These data confirm that pixatimod can interfere with binding of S1-267 RBD to cells containing HSPGs and ACE2 protein receptors. 268

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# 270 Pixatimod directly inhibits S1-RBD binding to ACE2

To further evaluate the mechanism of action of pixatimod its direct effects on the interaction of S1-RBD with the ACE2 protein receptor was measured using a competitive ELISA assay. Inhibition of binding of mS1-RBD preincubated with various concentrations of inhibitor compounds was measured by detection with an anti-RBD antibody. A dose response was observed with pixatimod showing an IC<sub>50</sub> of 10.1  $\mu$ g/ml (**Fig 2F**). In comparison heparin also demonstrated inhibitory activity but with lower potency (24.6  $\mu$ g/ml; **Fig 2F**). Importantly, this data confirms a direct mechanism of action of pixatimod via inhibition of S1-RBD binding to the ACE2 protein receptor.

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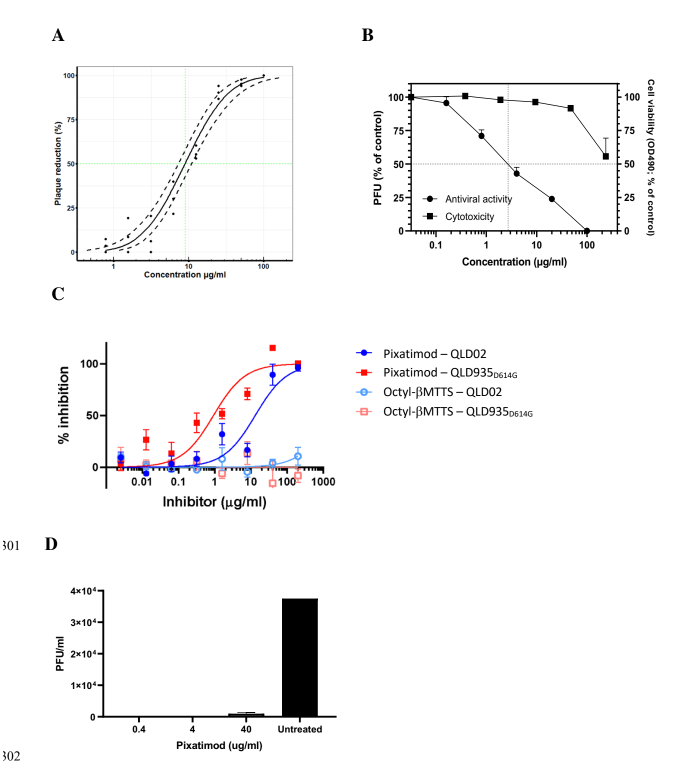
# 279 Pixatimod inhibits SARS-CoV-2 infection

The effect of pixatimod on SARS-CoV-2 infection of Vero cells was examined using a standard 280 plaque reduction neutralisation assay. Pixatimod was pre-incubated with the SARS-CoV-2 clinical 281 isolate from Victoria, Australia (VIC01) for 1 hr before infecting the cells. Significant decreases 282 were observed in the number of PFU after pixatimod treatment of SARS-CoV-2 (Fig 3A). Analysis 283 of multiple dose response curves yielded an  $EC_{50}$  for pixatimod in the range of 2.4-13.8 µg/mL 284 (mean 8.1  $\mu$ g/ml; n=3 assays) (**Table 1**). In comparison, an EC<sub>50</sub> of ~10  $\mu$ g/ml has been observed 285 for unfractionated heparin with a SARS-CoV-2 Italy UniSR1/2020 isolate (8) and 20-64 µg/ml for 286 the SARS-CoV-2 VIC01 isolate (21). 287

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To establish that these antiviral effects were relevant for wider clinical viral isolates, assays were 289 conducted with the isolate DE-Gbg20 from Sweden in a plaque reduction assay. Pixatimod 290 inhibited infectivity of the DE-Gbg20 isolate with an EC<sub>50</sub> value of 2.7 µg/mL (Fig 3B), similar to 291 that found in experiments with the VIC01 isolate. Analysis of pixatimod cytotoxicity for Vero cells 292 using a tetrazolium-based assay revealed that pixatimod decreased by 50% (CC<sub>50</sub>) the viability of 293 Vero cells at concentration >236  $\mu$ g/mL, i.e., well above the EC<sub>50</sub> values observed in the plaque 294 reduction assay (Fig. 3B; Table 1). Selectivity index (SI) values for pixatimod ranged from >17 to 295 >98 for these assays. 296

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303 Figure 3: Pixatimod inhibits infection of Vero and human bronchial epithelial cells with different SARS-CoV-2 304 virus isolates. Live virus infectivity assays were performed as described in Methods for 3 different SARS-CoV-2 305 isolates (representative data shown). A, Plaque reduction neutralization assay of Victoria isolate (VIC01) with a Probit mid-point analysis curve  $\pm 95\%$  confidence intervals (dashed lines) (EC<sub>50</sub> 8.9 µg/ml; 95% CI, 7-11; n=3). B, Plaque 306 reduction assay of DE isolate,  $EC_{50}$  2.7 µg/ml; n=3, +/- SD. C, Cytopathic assay of Queensland isolates,  $EC_{50}$  13.2 307 308 (QLD02) and 0.9 (QLD935 with D614G mutation) µg/ml n=6, +/- SEM. Representative examples are shown in each case. Results of pixatimod inhibition of SARS-CoV-2 infectivity are expressed as percent plaque reduction (A), plaque 309 310 number as a percent of control (B), or percent inhibition from cytopathic effect (C). Panel B also shows cytotoxicity 311 data for Vero cells for calculation of CC<sub>50</sub> value (>236  $\mu$ g/ml). In panel C, data is also shown for octyl  $\beta$ -maltotetraoside 312 tridecasulfate (Octyl-BMTTS; Supplementary Materials Fig S2), an analogue of pixatimod which lacks the steroid side-313 chain. D. Plaque assay to measure inhibitory effect of pixatimod on viral shedding in BCi-NS1.1 bronchial epithelial

- cells grown in an air liquid interface (ALI). ALI cultures were infected for 2h with SARS-CoV-2 (VIC01, MOI=0.2)
- previously preincubated for 1 hr at 37°C with 0.4, 4 and 40 µg/ml of pixatimod or HBSS for untreated. After 72 hours
- 316 an apical rinse was performed with 200 µl of HBSS and 100 µl of the wash were used in a plaque assay in Vero E6
- 317 cells. Values are expressed as plaque forming unit (PFU)/ml, n=2 + SD.
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# Table 1: Anti-SARS-CoV-2 activities of pixatimod in Vero cells.

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SARS-CoV-2 isolate	Assay Method	EC <sub>50</sub> (µg/mL) <sup>a</sup>	CC50 (µg/mL)	Selectivity Index <sup>a</sup>
VIC01 isolate	Plaque reduction	$8.1 \pm 3.1 \ (2.4-13.8)^{a}$		29 (>17 to >98ª)
DE-Gbg20 isolate	Plaque reduction	2.7 <sup>b</sup>		>87
	Cytopathic effect	(0.8 – 11.6°)	>236 <sup>d</sup>	>20 to >295
QLD02 isolate	Cytopathic effect	13.2 (8.0 - 21.6)		>17
QLD935 isolate	Cytopathic effect	0.9 (0.4 – 1.9)		>200

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<sup>a</sup> Mean values and individual assay result ranges, and resulting selectivity index ranges, in brackets.

 $^{b}$ Mean EC<sub>50</sub> ±SE based on the data from three independent virus plaque reduction assays (PRNT<sub>50</sub> values).

<sup>c</sup> EC<sub>50</sub> computed by the Reed and Muench formula based on the cytopathic effect assay. Range indicates upper

328 (complete protection of cells) and lower (partial protection) limits of EC<sub>50</sub> estimation.

<sup>d</sup> Cytotoxicity in Vero cells (determined at University of Gothenburg).

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332 In addition to the plaque reduction assays pixatimod inhibition of SARS-CoV-2 infectivity was assessed using assays that measured the cytopathic effects of the virus as an endpoint. Using the 333 Swedish DE-Gbg20 isolate, and two Australian isolates from Queensland (QLD02 and QLD935), 334 the EC<sub>50</sub> values for pixatimod inhibition of SARS-CoV-2 infectivity were determined to be 0.8-335 11.6, 13.2 and 0.9 µg/mL, respectively (**Table 1**), values comparable with those observed for the 336 plaque reduction assays (**Table 1**). We also noted that a pixatimod analogue octyl β-maltotetraoside 337 tridecasulfate (without the steroid side-chain) (Fig S2) lacked efficacy for both OLD02 and 338 QLD935 isolates (Fig 3C), demonstrating the importance of the steroid side-chain for activity, and 339 supporting the notion of the sterol moiety promoting RBD switching position and enhancing 340 availability of the heparin binding site. Notably, both DE-Gbg20 and QLD935 isolates contain the 341 D614G mutation of the spike protein commonly present in recent isolates (Table S1) (22). The 342 343 QLD935 isolate exhibited lower cytopathicity, which could partially contribute to the observed lower EC<sub>50</sub> for pixatimod against this isolate. 344

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To provide evidence of antiviral properties of pixatimod in a more physiologically relevant model for infection of human cells, we used a bronchial airway epithelial *in vitro* system of SARS-CoV-2 infection and replication. The hTERT transformed bronchial epithelial cell line BCi-NS1.1 differentiates into airway multi-ciliated cells when grown in Transwells at the air-liquid interface (ALI) cultures which show robust infection with SARS-CoV-2 when harvested 72 h after inoculation (*23,24*). To validate this model we firstly infected ALI bronchial cultures by adding SARS-CoV-2 (VIC01, MOI=0.2) on the apical side of the culture system and measured up to 1000

fold increase in viral RNA between 2h and 72h post-infection, measured by RT-PCR of viral RNA 353 using primers for N2/N3 regions of the N gene of SARS-CoV-2 (Fig S3, panel A). To determine 354 if pixatimod has inhibitory effects on SARS-CoV-2 infection and replication in this model, the virus 355 was preincubated with 0.4, 4 or 40 µg/ml of pixatimod (or HBSS for untreated) for 1 hr at 37°C 356 before the inoculum was added to the apical side of the cells. We examined infectious virus in the 357 apical wash from BCI ALI cultures by plaque assay on Vero E6 cells. Viral shedding of ALI 358 359 cultures infected with SARS-CoV-2 in the presence of 0.4 and 4 µg/ml pixatimod was completely abolished in comparison to the cells infected with the untreated virus as a control (Fig 3D), whereas 360 at 40 µg/ml pixatimod resulted in >40 fold reduction in PFU/ml (Fig 3D). To corroborate these 361 findings viral shedding from the apical side of the culture and viral load in the infected cells was 362 also measured by RT-PCR. Incubation of SARS-CoV-2 with different concentrations of pixatimod 363 reduced SARS-CoV-2 shedding from ALI cells into the apical phase by > 98%, and reduced SARS-364 CoV-2 RNA within the cells by >97% at 72hr at all pixatimod doses, compared to untreated infected 365 cells (Fig S3, panels A & B). Notably, efficacy at 0.4 and 4 µg/ml was >99% in both cases 366 indicating high potency of pixatimod in inhibition of SARS-CoV-2 infection of human bronchial 367 cells. Overall, these data in a physiologically-relevant human bronchial cell model show that 368 pixatimod is a potent inhibitor of SARS-CoV-2 in terms of both lowering viral shedding in the 369 apical compartment and viral load in the cell layer. 370

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# 372 Pixatimod inhibits SARS-CoV-2 infection in vivo

373 The potential efficacy of pixatimod in vivo was explored with an established SARS-CoV-2 lethal infection model using hACE2 transgenic mice wherein hACE2 is expressed behind a keratin 18 374 promoter (25-28). Virus was inoculated in 50  $\mu$ l via the intranasal route into the lungs (25-27). A 375 single prophylactic treatment of 16 mg/kg of pixatimod was given to K18-hACE2 mice one day 376 before intranasal infection with SARS-CoV-2. Two control groups were included, K18-hACE2 377 mice that were treated with saline and infected with virus, and C57BL/6J mice given pixatimod but 378 no virus. Mice were weighed daily and euthanized on day 5 when weight loss approached 20%, 379 the ethically defined endpoint. As reported previously in other preclinical models of infectious 380 381 disease (18,19), pixatimod led to an initial bodyweight loss of 8%. Weight loss was thus presented 382 relative to day 1 post-infection in order to focus on infection-induced weight loss (Fig. 4A), which generally occurs after day 3 (26). The mean weight loss for saline-treated K18-hACE2 mice was 383 significantly higher than for pixatimod treated K18-ACE2 mice (p=0.043, repeat measures 384 ANOVA including data from days 4 and 5), with the latter showing no significant infection-385 associated weight loss (Fig. 4A). 386

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Tissue viral loads were assessed on day 5 post infection, with pixatimod-treated animals showing a significant 3.1  $\log_{10}$  fold reduction in viral titers in nasal turbinates (p=0.037, Kolmogorov Smirnov test, non-parametric data distribution), and a 4.6  $\log_{10}$  fold reduction in viral titers in brain, which approached significance (p=0.0588, t test, parametric data distribution). Viral titers in the lungs were not significantly affected by pixatimod treatment (**Fig 4B**). In view of the robust nature of this model, with rapid effects such as massive weight loss and high level of lethality, these data indicate a remarkable prophylactic anti-SARS-CoV-2 effect of pixatimod.

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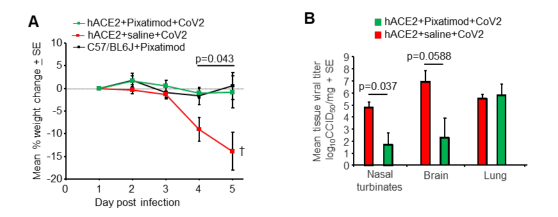


Fig 4: Pixatimod inhibits SARS-CoV-2 infection in K18 hACE2 transgenic mice. (A) Mean percentage weight
change relative to day 1 post infection. Statistics by repeat measures ANOVA for days 4 and 5. n=4 mice per group,
+/- SE. Mice were euthanized on day 5. (B) Mean tissue titres on day 5 post infection. Statistics by Kolmogorov
Smirnov test (Nasal turbinates) and t test (Brain). n=4 mice per group, +/- SE.

# 403 **Discussion**

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The current COVID-19 pandemic illustrates the critical need to develop both effective vaccines and 104 therapeutics for emerging viruses; established antiviral agents appear to have limited utility against 405 SARS-CoV-2. Owing to their use as a means of cell attachment by many viruses, HS represents an 406 ideal broad-spectrum antiviral target (2). Binding of a viral protein to cell-surface HS is often the 407 first step in a cascade of interactions that is required for viral entry and the initiation of infection 408 (29). As HS and heparin contain the same saccharide building blocks and HS-binding proteins also 409 interact with heparin, this drug is gaining attention beyond its anticoagulant properties in COVID-410 19 treatment (29). Here we demonstrate a direct mechanism of action of pixatimod on attenuating 411 S1-RBD binding to ACE2. These data are supported by recent studies on heparin using native mass 412 spectrometry (30), and also reveal for the first time the ability of HS mimetics to inhibit S1-RBD 413 binding to ACE2. 414

415

Heparin has been shown to inhibit binding of SARS-CoV-2 spike protein to a human cell line (31), 416 and to inhibit entry into human cells of pseudovirus carrying the SARS-CoV-2 spike protein 417 (10,32). However, the question of whether therapeutic doses of heparins are effective for COVID-418 19 patients as an antiviral treatment awaits the outcome of clinical trials; bleeding complications 419 are possible (33), though non-anticoagulant heparin or HS preparations could be deployed that 420 reduce cell binding and infectivity without a risk of causing bleeding (9,10). However, HS mimetics 421 422 offer additional advantages in comparison to heparin beyond simply reducing anticoagulant activity (9), most notably their ready availability at scale via synthetic chemistry production that addresses 423 the well-known fragility of the heparin supply chain (11). As a clinical-stage HS mimetic, 124 pixatimod provides better control over structure, molecular weight diversity (a single molecular 425 entity), sulfation, purity and stability. Herein, we reveal a direct interaction of the clinical candidate 426 pixatimod with the S1 spike protein RBD, supported by molecular modelling data. Pixatimod also 427 inhibited the interaction of S1 RBD with Vero cells which express the ACE2 receptor. Moreover, 428 infectivity assays, of two types (plaque reduction and cytopathic effect, Table 1) confirm pixatimod 429 is a potent inhibitor of SARS-CoV-2 infection of Vero cells, at concentrations ranging from 0.8 to 430

13.8  $\mu$ g/mL which are well within its known therapeutic range. Interestingly, we noted that the lipophilic steroid side chain of pixatimod was critical for its potency and is predicted from modelling to interact with S1-RBD. This unique feature, making it an unusual amphiphilic HS mimetic, has also been shown to confer virucidal activity against Herpes Simplex virus by disruption of the viral lipid envelope (*15*).

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Pixatimod has only mild anti-coagulant activity, and has been administered i.v. to over 80 cancer 437 patients, being well tolerated with predictable pharmacokinetics (PK) and no reports of heparin-438 139 induced thrombocytopenia (12). Further, cytotoxicity in vitro is low; we observed a  $CC_{50} > 100$  $\mu$ M (>236  $\mu$ g/mL) in Vero cells, consistent with cytotoxicity data on human cells (34). Importantly, 140 the maximum plasma concentration (Cmax) of pixatimod following a single treatment of 100 mg 441 in cancer patients is 29.5 µg/mL with a Cmin of 2.7 µg/mL measured one week following treatment 142 (12), indicating that an equivalent dosing regimen should be sufficient to achieve antiviral activity 443 in human subjects. The low anticoagulant activity of pixatimod is an advantage since it could be 144 used as a direct antiviral agent in combination therapies with heparins, which are being used to treat 145 coagulopathies observed in COVID-19 patients (35). It was also encouraging that pixatimod 146 inhibition of multiple clinical isolates of SARS-CoV-2 was noted, demonstrating potential for 147 widespread effectiveness. The presence of multiple binding sites for pixatimod in the Spike protein 148 would suggest robustness against mutations that may arise later in pandemic and/or in the following 149 coronavirus outbreaks. While recent widespread isolates with D614G spike mutants appear to be 450 2-3 fold more sensitive to the antiviral activity of pixatimod, caution needs to be taken in 451 452 interpreting the data of the cytopathicity assay used to determine this activity as G614 isolates (at least QLD935) exhibited lower cytopathicity than D614 isolates. 453

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Since SARS-CoV-2 is predominantly a respiratory infection, our data from human bronchial cells 455 are directly supportive of high potency of pixatimod against infection of clinically relevant human 456 cells. Further, the potential efficacy in humans is supported by our initial data from the established 457 K18 hACE2 transgenic mouse model which demonstrated the ability to rescue the dramatic weight 458 loss phenotype observed. Given the half-life of pixatimod in mice is 38h (34), compared to 141h in 459 human (13), future studies should adopt repeat-dose schedules to identify the optimal schedule in 460 models of SARS-CoV-2 infection. We also suggest that direct intranasal delivery of pixatimod 461 would be worth investigation, in view of recent positive data on the delivery of nebulized heparin 462 for treatment of ARDS (36) which has significance for potential application of heparin in COVID-463 19 treatment (21). 164

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Why Pixatimod should have such a dramatic effect in nasal turbinates and brain, but not lungs in 466 the K18 hACE2 model is unclear but may reflect very high lung (supra-physiological) K18-driven 467 hACE2 receptor expression in the lungs in this model (27) and/or pharmacokinetics (34). 468 Nevertheless the results are encouraging, as the initial infection in humans occurs in the upper 469 respiratory tract, with ARDS developing only after infection has over time spread down into the 170 lungs (37). This progression is not recapitulated in the mouse model where fulminant lung infection 471 requires direct inoculation of virus into the lungs via the intranasal route. Infection of the brain 472 may occur via the olfactory epithelium (38) so the reduced infection of brain may reflect the reduced 473 infection of nasal turbinates. Although CNS involvement in COVID-19 in humans is now well 174

documented<sup>32</sup>, the fulminant brain infection seen in this mouse model does not seem to be a feature
 of human disease.

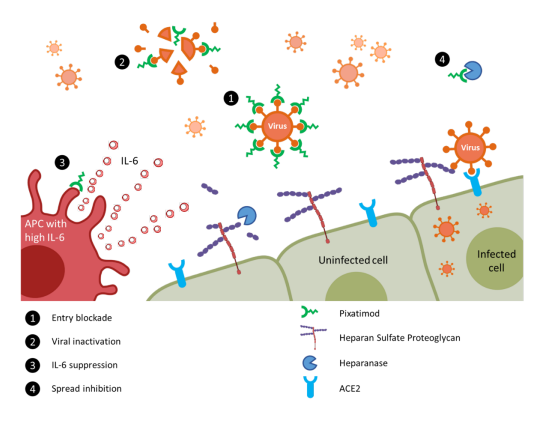
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It is notable that there are multiple potential mechanisms of action of pixatimod against SARS-178 179 CoV-2 (summarised in Fig 5), including direct inhibition of HS-Spike-ACE2 interactions via complex and multiple interactions with amino acids in RBD. Notably, the latter contrasts with the 480 more restricted epitopes typical of antibodies which are observed to be subject to mutational escape 481 potentially requiring vaccine redesign or use of multivalent strategies (39). In this respect we 482 speculate that heparin mimetics like pixatimod may prove important therapeutic tools for initial 483 responses to escape variants. Beyond this. pixatimod also has immunomodulatory effects which 184 may alleviate some of the immunopathologies associated with moderate-severe COVID-19 disease. 485 Pixatimod also inhibits the pro-inflammatory enzyme heparanase (34) and has been demonstrated 486 to suppress IL-6 in inflammatory (pancreatitis) and viral (Ross River virus) animal models (14, 18). 487 Moreover, it blocks the heparanase-dependent invasion of macrophages into tumours in mouse 488 cancer models (35) which may be relevant to invasion of monocytes and macrophages into the 189 lungs associated with severe COVID-19 disease (40). Vaccinia virus has recently been shown to 490 rely on host heparanase to degrade HS in order to spread to distant sites (41), revealing a role for 491 heparanase in the progression of disease that may also apply for SARS-CoV-2 in COVID-19. 192 Notably, increased plasma heparanase activity is associated with COVID-19 (42). Thus multiple 493 additional beneficial effects of pixatimod might be anticipated from its heparanase inhibitory 194 495 properties.

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Based on the data presented here, pixatimod has potent antiviral activity against SARS-CoV-2 at 497 therapeutically relevant concentrations both in vitro and in vivo, and against relevant human 198 bronchial cells. These activities are in addition to the known heparanase-inhibitory and 199 500 immunomodulatory properties of pixatimod which may further support the host response to COVID-19 infection. Collectively this provides a strong rationale to justify entry of pixatimod into 501 clinical trials for COVID-19. Furthermore, we have demonstrated the first proof-of-concept for 502 employing HS mimetics against SARS-CoV-2 with implications for wider future applications of 503 this class of broad-spectrum antivirals, potentially against SARS-CoV-2 escape variants, other HS-504 505 binding viruses and also those that may emerge as future global threats.

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**Figure 5: Proposed multi-modal mechanisms of pixatimod activity against SARS-CoV-2 and other viruses.** The principal mode of action demonstrated here is that pixatimod acts as a decoy receptor<sup>1</sup>, blocking S1-RBD binding to HS co-receptors and inhibiting viral attachment to host cells, thus blocking viral infection. Additional potential modes of action include: [2] virucidal activity of pixatimod, dependent upon the cholestanol moiety<sup>15</sup>, which may lead to degradation and permanent inactivation of SARS-CoV-2 virus particles; [3] suppression of IL-6 secretion by antigen presenting cells, primarily macrophages<sup>14</sup>; and [4] blocking viral escape from host cells by inhibiting heparanase which otherwise promotes viral escape by cleaving HS receptors<sup>41,42</sup>.

#### 518 519

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# 521 Materials and Methods

*Computational methods:* The crystal structure of the RBD-ACE2 complex (PDB ID: 6LZG) was 522 retrieved from the RCSB Protein Data Bank. Structures were stripped of water molecules, ACE2 523 chain and any cofactors and/or ligands present. UCSF Chimera was used to edit the structure and 524 for visualisation. Without prior knowledge of the pixatimod binding site, one molecule of the ligand 525 was placed in the simulation system containing the protein, solvent and ions and molecular 526 dynamics (MD) simulations were performed for 600 ns using the AMBER16 package. Such 527 unguided simulations, as reviewed before (43), have been used to predict the binding sites on a 528 protein's surface and drive the design of new ligands. All the MD simulations were carried out 529 using the pmemd.cuda module (44) of the AMBER 16 molecular dynamics package and the 530 analyses were performed using the cpptraj module of AmberTools16 (45). Simulation systems were 531 set up by placing the spike RBD domain at the centre of the octahedral simulation box (with an 532 extension of at least 12 Å from each side). Pixatimod was randomly placed in the box. This was 533 followed by addition of TIP3P water (46) and Na<sup>+</sup> ions for neutralising the charge of the system. 534 Proteins were parameterized using the Amber99SB-ildn force field (47) whereas Glycam-06 535

(version j) (48) and Lipid14 (49) force fields were used for the sulfated tetrasaccharide and cholestanol moieties of pixatimod, respectively. Four replicates of the unguided simulations were performed ( $4 \times 600$  ns). Periodic boundary conditions were applied, and the time step was set to 2 fs. The electrostatic energy was calculated with the particle mesh Ewald (PME) method. SHAKE constraints were applied on the bonds involving hydrogen. A cut-off of 12 Å was applied to the Lennard-Jones and direct space electrostatic interactions with a uniform density approximation included to correct for the long-range van der Waals interactions.

The system was first minimized without electrostatics for 500 steps, then with a restraint of 25 543 kcal/(mol  $Å^2$ ) applied on the protein and pixatimod. This minimization was followed by 100-ps 544 MD simulation with 25 kcal/(mol Å<sup>2</sup>) positional restraints applied on the protein and ligand, and 545 the temperature was slowly increased from 0 to 300 K. Then, followed by 500 steps of steepest 546 descent cycles followed by 500 steps of conjugate gradient minimization, and 50-ps equilibrations 547 with a restraint force constant of 5 kcal/(mol  $Å^2$ ) in the protein and ligand, followed by final 2 ns 548 equilibration without restraints to equilibrate the density. The first few steps were all carried out at 549 constant volume followed by at least 600 ns production MD simulation at 300 K (Langevin 550 dynamics, collision frequency: 5/ps) with 1-atm constant pressure. Trajectories were collected and 551 data analyses such as RMSD, RMSF and free energy of binding were performed on the last 30000 552 frames. The binding free energy and pairwise residue contributions (50) were calculated using the 553 Molecular mechanics-Generalized Born (GB) equation (MM/GBSA) procedure implemented in 554 AmberTools16. The details of this method have been extensively reviewed (51). The polar 555 solvation energy contribution was calculated by using  $GB^{OBC II}$  (igb= 5) (52). The value of the 556 implicit solvent dielectric constant and the solute dielectric constant for GB calculations was set to 557 80 and 1, respectively. The solvent probe radius was set to 1.4 Å as default. The entropy calculation 558 is computationally expensive and therefore not performed for the purposes of this study. 559

Expression of His-tagged recombinant SARS-CoV-2 S1 RBD in E. coli: Residues 330-583 of 560 the SARS-CoV-2 spike protein (GenBank: MN908947) were cloned upstream of a N-terminal 561 6XHisTag in the pRSETA expression vector and transformed into SHuffle® T7 Express Competent 562 E. coli (NEB, UK). Protein expression was carried out in MagicMediaTM E. coli Expression Media 563 (Invitrogen, UK) at 30°C for 24 hrs, 250 rpm. The bacterial pellet was suspended in 5 mL lysis 564 buffer (BugBuster Protein Extraction Reagent, Merck Millipore, UK; containing DNAse) and 565 incubated at room temperature for 30 mins. Protein was purified from inclusion bodies using IMAC 566 chromatography under denaturing conditions. On-column protein refolding was performed by 567 applying a gradient with decreasing concentrations of the denaturing agent (from 8M Urea). After 568 extensive washing, protein was eluted using 20 mM NaH<sub>2</sub>PO<sub>4</sub>, pH 8.0, 300 mM NaCl, 500 mM 569 imidazole. Fractions were pooled and buffer-exchanged to phosphate-buffered saline (PBS; 140 570 mM NaCl, 5 mM NaH<sub>2</sub>PO<sub>4</sub>, 5 mM Na<sub>2</sub>HPO<sub>4</sub>, pH 7.4; Lonza, UK) using Sephadex G-25 media (GE 571 Healthcare, UK). Recombinant protein (termed EcS1-RBD) was stored at -4°C until required. 572

573 *Expression of S1-RBD in mammalian cells:* Secreted RBD-SD1 (termed mS1-RBD) was 574 transiently produced in suspension HEK293-6E cells. A plasmid encoding RBD-SD1, residues 575 319–591 of SARS-CoV-2 S were cloned upstream of a C-terminal HRV3C protease cleavage site, 576 a monomeric Fc tag and an His<sub>8x</sub> Tag were a gift from Jason S. McLellan, University of Texas at Austin. Briefly, 100 mL of HEK293-6E cells were seeded at a cell density of  $0.5 \times 106$  cells/ml 24hr before transfection with polyethyleneimine (PEI). For transfection, 100 µg of the ACE2 plasmid and 300 µg of PEI (1:3 ratio) were incubated for 15 min at room temperature. Transfected cells were cultured for 48 hr and fed with 100 mL fresh media for additional 48 hr before harvest. RBD-SD1was purified by HiTrap Protein G HP column (GE Healthcare, US) pre-equilibrated in PBS and eluted with 0.1 M glycine (pH 2.7). Purity of proteins was evaluated by Coomassie staining of SDS-PAGE gels, and proteins were quantified by BCA Protein Assay Kit (Thermo Scientific).

Secondary structure determination of SARS-CoV-2 S1 RBD by circular dichroism spectroscopy: 584 The circular dichroism (CD) spectrum of the SARS-CoV-2 S1 RBD in PBS was recorded using a 585 586 J-1500 Jasco CD spectrometer (Jasco, UK), Spectral Manager II software (JASCO, UK) using a 0.2 mm path length, quartz cuvette (Hellma, USA). All spectra were obtained using a scanning of 587 100 nm/min, with 1 nm resolution throughout the range  $\lambda = 190$  - 260 nm and are presented as the 588 the mean of five independent scans, following instrument calibration with camphorsulfonic acid. 589 SARS-CoV-2 S1 RBD was buffer-exchanged (prior to spectral analysis) using a 10 kDa Vivaspin 590 centrifugal filter (Sartorius, Germany) at 12,000 g, thrice and CD spectra were collected using 21 591 µL of a 0.6 mg/mL solution in PBS, pH 7.4. Spectra of heparin (unfractionated porcine mucosal 592 heparin, Celsus) were collected in the same buffer at approximately comparable concentrations, 593 since this is a polydisperse material. Collected data were analysed with Spectral Manager II 594 595 software prior to processing with GraphPad Prism 7, using second order polynomial smoothing through 21 neighbours. Secondary structural prediction was calculated using the BeStSel analysis 596 server (53). 597

To ensure that the CD spectral change of SARS-CoV-2 S1 RBD in the presence of pixatimod did not arise from the addition of the compound alone, a difference spectrum was analysed. The theoretical CD spectrum that resulted from the arithmetic addition of the CD spectrum of the SARS-CoV-2 S1 RBD and that of pixatimod differed from the observed experimental CD spectrum of SARS-CoV-2 S1 RBD mixed with compound alone. This demonstrates that the change in the CD spectrum arose from a conformational change following binding to pixatimod (Supplementary Materials, Fig S3).

Differential scanning fluorimetry: Differential scanning fluorimetry (DSF) was conducted on 505 mammalian expressed mS1-RBD (1 µg) in PBS pH 7.6 and 1.25 X Sypro Orange (Invitrogen) to 506 a total well volume of 40 µL in 96-well qPCR plates (AB Biosystems). Unfractionated porcine 507 mucosal heparin (Celsus) or pixatimod (10 µg) were introduced to determine the effect on the 508 thermal stability of, mS1-RBD using an AB biosystems StepOne plus qPCR machine, employing 509 the TAMRA filter setting. Melt curve experiments were performed following a 2-minute initial 510 incubation at 25 °C, with succeeding 0.5 °C increments every 30 s up to a final temperature of 511 90°C. Control wells containing H<sub>2</sub>O, heparin or pixatimod (10 µg) without mS1-RBD (1 µg) also 512 employed to ensure a change in the melt curve was solely a result of protein-ligand interactions and 513 interactions with Sypro Orange. Smoothed first derivative plots (9 neighbours, 2nd-order 514 polynomial, Savitxky-Golay) were constructed using Prism 8 (GraphPad). T<sub>m</sub> values were 515 calculated using MatLab softaware (R20018a, MathWorks) and  $\Delta T_m$  values determined from the 516 difference between the T<sub>m</sub> of RBD alone or in the presence of heparin or pixatimod. 517

Cell binding of S1 RBD: African green monkey Vero kidney epithelial cells (Vero E6) were 518 purchased from ATCC. Cells were maintained at 50-75% confluence in DMEM supplemented 519 with 10% foetal bovine serum, 20 mM L-glutamine, 100 U/mL penicillin-G and 100 U/mL 520 streptomycin sulfate (all purchased from Gibco/ThermoFisher, UK). Cells were maintained at 37 521 °C, in 5% CO<sub>2</sub> and plated into 96-well cell culture plates at 1000 cells/well in 100 µL of 522 maintenance medium. Cells were allowed to adhere overnight. Medium was aspirated and wells 523 524 were washed 3x with 200 µL calcium, magnesium-free PBS (CMF-PBS, Lonza, UK). Cells were fixed with 100 µL 10% neutral buffered Formalin (Thermofisher, UK) for 10 minutes at room 525 temperature, then washed 3x with 200 µL CMF-PBS. 100 µL CMF-PBS was added to each well 526 and plates were stored at 4 °C until use. Before use, wells were blocked with 200 µL CMF-PBS + 527 1% BSA (Sigma-Roche, UK) for 1 hour at room temperature, and washed 3x with 200 µL CMF-528 PBS + 0.1% Tween-20 (PBST, Sigma-Roche, UK) followed by 2x with 200 µL CMF-PBS. 529

His-tagged S1-RBD (50 µg/mL) and compounds at indicated concentrations were added to each 530 well in 25  $\mu$ L PBST + 0.1% BSA as indicated. Wells were incubated for 1 hour at room temperature 531 with rocking. Wells were washed 3x with 200 µL PBST and 2x with 200 µL CMF-PBS. Binding 532 of His-tagged S1-RBD was detected with Alexa Fluor 488 anti-his tag antibody (clone J095G46, 533 Biolegend, UK) 1:5000 in 25 µL PBST + 0.1% BSA per well. Wells were incubated in the dark 534 for 1 hour at room temperature with rocking. Wells were washed 3x with 200 µL PBST and 2x 535 with 200 µL CMF-PBS. Fluorescence was read at Ex. 485:Em 535 on a Tecan Infinite M200Pro 536 plate reader. Results are presented as normalized mean (where 0 is the fluorescence without added 537 S1-RBD, and 1 is the fluorescence with 50  $\mu$ g/mL S1-RBD;  $\pm$  %CV, n=3). 538

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Competition ELISA for S1 RBD binding to ACE2: High binding 96 well plates (Greiner) were 540 coated with 3  $\mu$ g/mL streptavidin (Fisher) in 50 mM sodium carbonate buffer pH 9.6 (50  $\mu$ L/ well) 541 for 1 hour at 37 °C. Plates were washed 3 times with 300 µL PBS, 0.2% Brij35 (PBSB) and blocked 542 with 300 µL PBSB + 1% casein for 1 hour at 37 °C. Plates were washed a further 3 times with 300 543 544 µL PBSB prior to the addition of 100 ng/mL BiotinylatedACE2 (Sino Biological) in PBSB + 1% casein (50 µL/ well) and incubated for 1 hour at 37 °C. Plates were again washed 3 times with 300 545 µL PBSB prior to the addition of 50 µL/well mS1-RBD (5µg/mL) in PBSB + 1% casein, which had 546 been pre-incubated for 30 minutes at room temperature with or without varying concentrations of 547 heparin or pixatimod (100- 0.7 µg/mL) in separate tubes. Plates were incubated for 1 hour at 37 °C 548 to allow for mS1-RBD-ACE2 binding and were subsequently washed with 300 µL/well PBSB. 549 Bound mS1-RBD was detected by incubation with 0.5 µg/mL Rabbit-SARS-CoV-2 (2019-nCoV) 550 Spike RBD Antibody (Stratech) in PBSB + 1% casein (50 µL/well) for 1 hour at 37 °C. Following 551 a further 3 washes with PBSB plates were incubated for 30 minutes at 37 °C with horseradish 552 peroxidase conjugated Donkey anti-Rabbit IgG diluted 1:1000, v/v in PBSB + 1% casein 553 (Bioledgend). Plates were washed a final 5 times with 300 µL PBSB before being developed for 10 554 minutes with 3,3',5,5'- tetramethylbenzidine prepared according to the manufacturer's instructions 555 (Fisher). Reactions were stopped by the addition of 20  $\mu$ L 2M H<sub>s</sub>SO<sub>4</sub> and plates were read at  $\lambda$  =450 556 nm using a Tecan Infinate M200 Pro mulit-well plate reader (Tecan Group). Control wells 557 containing no biotinylated ACE2 were employed to ensure binding was specific. 558

Live SARS-CoV-2 virus assays: SARS-CoV-2 Victoria isolate (GISAID accession, 560 EPI ISL 406844): a plaque reduction assay was performed with the SARS-CoV-2 561 Victoria/01/2020 (passage 3) isolate, generously provided by The Doherty Institute, Melbourne, 562 Australia at P1, was diluted to a concentration of 1.4 x 10<sup>3</sup> pfu/mL (70 pfu/50µL) in minimal 563 essential media (MEM) (Life Technologies, California, USA) containing 1% (v/v) foetal calf serum 564 (FCS) (Life Technologies) and 25 mM HEPES buffer (Sigma) and mixed 50:50 with pixatimod 565 dilutions, in a 96-well V-bottomed plate. The plate was incubated at 37 °C in a humidified box for 566 1 hour to allow the virus to be exposed to pixatimod. The virus-compound mixture was transferred 567 onto the wells of a washed 24-well plate that had been seeded with Vero E6 cells 568 [ECACC 85020206] the previous day at 1.5 x 10<sup>5</sup> cells/well. The virus-compound mixture was left 569 to adsorb for an hour at 37°C, then plaque assay overlay media was applied (MEM containing 1.5% 570 carboxymethylcellulose (Sigma, Dorset, UK), 4% (v/v) FCS and 25 mM HEPES buffer). After 571 incubation at 37 °C in a humidified box, for 5 days, plates were fixed overnight with 20% (v/v) 572 formalin/PBS, washed with tap water and then stained with methyl crystal violet solution (0.2% 573 v/v) (Sigma) and plaques were counted. Compound dilutions were performed in either duplicate or 574 quadruplicate. Compound dilutions and cells only were run in duplicate, to determine if there was 575 any cell cytotoxicity. A mid-point probit analysis (written in R programming language for statistical 576 computing and graphics) was used to determine the amount (µg/mL) required to reduce SARS-577 CoV-2 viral plaques by 50% (PRNT50) compared with the virus only control (n=5). An internal 578 positive control for the PRNT assay was run in triplicate using a sample of heat-inactivated human 579 MERS convalescent serum known to neutralise SARS-CoV-2 (National Institute for Biological 580 581 Standards and Control, UK).

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SARS-CoV-2 DE-Gbg20 isolate (GISAID accession under application): Plaque reduction assay 583 for SARS-CoV-2 clinical isolate DE-Gbg20 from Sweden was performed in a similar manner, 584 except for the virus and the pixatimod (fivefold decreasing concentrations at a range 100-0.16 585 µg/ml) were diluted in DMEM supplemented with 2% heat-inactivated FCS, and 100 U of penicillin 586 and 60 µg/ml of streptomycin (DMEM-S). The virus (100 PFU) and pixatimod (fivefold decreasing 587 concentrations at a range  $100 - 0.16 \,\mu\text{g/ml}$ ) were mixed and incubated for 30 min in humidified 588 atmosphere comprising 5% CO<sub>2</sub> (CO<sub>2</sub> incubator). The mixtures were then transferred to Vero cells 589 (ATCC CCL-81) and following incubation with cells for 90 min in the CO<sub>2</sub> incubator, the 590 methylcellulose overlay was added. Three separate experiments each with duplicates were 591 performed. 592

A cytopathic effect assay was performed with the SARS-CoV-2 DE-Gbg20 isolate and Vero cells 593 (ATCC) plated at 2 x 10<sup>4</sup> per well in 96-well plates the day prior to the experiment. Serial fivefold 594 dilutions of pixatimod in DMEM supplemented with 2% heat-inactivated FCS, and 100 U of 595 penicillin and 60 µg/mL of streptomycin (DMEM-S) were incubated with 100 TCID50 of SARS-596 CoV-2 isolate DE for 20 min in humidified atmosphere comprising 5% CO<sub>2</sub> (CO<sub>2</sub> incubator). The 597 final concentrations of pixatimod were in a range 0.075 µg/mL to 47.2 µg/mL. The cells were 598 rinsed once with 50 µL of DMEM-S, and then 200 µL of the virus-pixatimod mixtures were added 599 to each well with cells (in quadruplicates). After incubation of the virus-pixatimod mixtures with 700 cells for 3 days in the CO<sub>2</sub> incubator, the cells were inspected under a microscope for the presence 701 of virus induced cytopathic effect where complete protection of cells were denoted as "+" while a 702

partial protection ( $^{50\%}$  of cells showing no cytopathic effect) was recorded as "+/-". The 50% endpoint (EC<sub>50</sub>) was computed by the Reed and Muench method.

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SARS-CoV-2 QLD02 (GISAID accession EPI ISL 407896) and QLD935 (GISAID accession 706 EPI ISL 436097) clinical isolates from Australia: A cytopathic effect assay was carried out as 707 described above for the DE-Gbg20 isolate, with 10 ffu/well and 3 days incubation. In this assay, 708 Vero E6 cells were plated at 2 x 10<sup>4</sup> per well in 96-well plates the day prior to experiment. Serial 709 five-fold dilutions of pixatimod in DMEM supplemented with 2% heat-inactivated FCS, and 100 710 711 U of penicillin and 60 µg/mL of streptomycin (DMEM-S) were incubated with 10 foci forming units of SARS-CoV-2 OLD02 or OLD935 isolate and incubated for 30 min in humidified 712 atmosphere comprising 5%  $CO_2$  (CO<sub>2</sub> incubator). The cells were rinsed once with 50  $\mu$ L of DMEM-713 S, and then 200 µL of the virus-pixatimod mixtures were added to each well with cells (in 714 triplicates). After incubation of the virus-pixatimod mixtures with cells for 3 days in the CO<sub>2</sub> 715 incubator, the cells were fixed with 4% PFA and then stained with crystal violet. Then crystal violet 716 was released by methanol and OD at 595nm was measured to quantify cell viability (protection 717 from infection). The EC<sub>50</sub> was then calculated using GraphPad Prism. 718

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Cytotoxicity assays: The assay was performed as described previously (17). Briefly, Vero cells 720 (ATCC, 2 x  $10^4$  cells/well) were seeded in 96 well cluster plates to become nearly confluent at the 721 day of the experiment. The cell growth medium was then removed and 100 µL of serial fivefold 722 dilutions of pixatimod in DMEM-S (ranging from 0.09 to 236 µg/mL) were added to cells. 723 724 Following incubation of cells with pixatimod for 3 days in the CO<sub>2</sub> incubator, 20 µL of the MTS salt containing CellTiter 96 Aqueous One Solution reagent (Promega, Madison, WI) was added and 725 incubated for further 1-2 h at 37 °C. The absorbance was recorded at 490 nm against a background 726 of 650 nm. Two separate experiments each in duplicates were performed and the results are 727 expressed as percentage of absorbance value detected with pixatimod relative to control cells. 728

Human bronchial cell infection assays: The hTERT transformed bronchial epithelial cell line BCi-730 NS1.1 (23,24) expanded in PneumaCult-Ex Plus Basal Medium supplemented with Pneumacult Ex 731 Plus supplements, hydrocortisone, nystatin and penicillin-streptomycin. BCi-NS1.1 cells were 732 grown at an Air Liquid Interface (ALI) in PneumaCult-ALI Basal Medium supplemented with 733 Pneumacult ALI supplement hydrocortisone, PneumaCult ALI maintenance supplement, heparin, 734 nystatin and penicillin-streptomycin (all from Stemcell Technologies) at 37°C in 5% CO<sub>2</sub>. BCi-735 NS1.1 cell ciliation was observed by microscopy and cells were differentiated and ciliated after 4 736 to 5 weeks at ALI. Transepithelial electrical resistance (TER) was monitored weekly using an 737 EVOM Voltohmmeter (World Precision Instruments) and cells with a TER  $\geq 1000 \text{ W}\Omega\text{m}^2$  were 738 739 used.

540 SARS-CoV-2 strain BetaCoV/ Australia/VIC01/2020 was used for infection: 50,000 plaqueforming units (MOI=0.1-0.2, obtained from Public Health England and propagated in Vero E6 cells for no more than two passages before use) were preincubated with either 4, 40 and 400  $\mu$ g/ml of

Wockhardt heparin or 0.4, 4 and 40  $\mu$ ug/ml of pixatimod for 1 hr at 37°C in a total volume of 200  $\mu$ l (or an equivalent volume of saline was added apically in the untreated controls). BCi-NS1.1 cells (4 to 8 weeks after ALI) after being washed three times with HBSS were apically infected with the

preincubated virus and compounds. After 2h the virus-compound solution was removed from the

apical side and washed twice each time with 200ul HBSS. The last wash was collected for plaque

assays and 50 µl stored in Qiazol. Similarly at 72 h a further 200 µl HBSS wash was performed and
collected as at 2h. Cells were also harvested with QIAzol (QIAGEN) at 2 h after the HBSS wash
and at 72 h post-infection for RNA extraction.

Plaque assay was performed to quantify extracellular virus released in the supernatant of infected 751 ALI. Vero E6 cells were seeded at 2.5 x 10<sup>5</sup> cells/well in a 12-well plate and left for a period of 24 752 hours in DMEM supplemented with 10% foetal bovine serum (FBS), glutamine and 50 U/ml 753 penicillin-streptomycin at 37°C in 5% CO<sub>2</sub>. Cells were washed once with infection medium (serum-754 free DMEM supplemented with 25 mM HEPES) and 100 µl from washes of infected ALI cultures 755 added to wells in 10 fold serial dilutions. After a 1 hour incubation at 37°C in 5% CO2, infectious 756 supernatants were removed and a 1.5ml overlay of 1 x DMEM supplemented with 4% FBS, 25 mM 757 HEPES and 0.6% (w/v) cellulose (Sigma; cat no 435244) was added. Plates were incubated at 37°C 758 and 5% CO2 for 72 hours before removing the overlay, fixing with 8% formaldehyde in PBS, and 759 staining with 0.1% (w/v) crystal violet in a solution of 20% (v/v) ethanol. 760

RNA was isolated from cell lysates using standard phenol–chloroform extraction, and reverse
transcribed to cDNA using a High Capacity cDNA Reverse Transcription kit (Thermo Fisher
Scientific) following the manufacturer's instructions. Taqman gene expression assays for N2 and
N3 regions of the SARS-CoV-2 N gene were made (ThermoFisher) following CDC specifications
(Division of Viral Diseases, National Center for Immunization and Respiratory Diseases, Centers
for Disease Control and Prevention, Atlanta, GA, USA, 20 January 2020 copy).

To measure viral shedding in the apical side, gene expression was quantified using the  $2^{-\Delta Ct}$  method.

For determination of relative changes of RNA within the cells the average Ct values of both SARS-

<sup>769</sup> CoV-2 N2 and N3 gene RNA were normalised against the house keeping gene B2M and shown as <sup>770</sup> fold changes compared to the untreated cells using the  $2^{-\Delta\Delta Ct}$  method.

771

K18-hACE2 mouse experiments: Mouse experiments are approved by the QIMR Berghofer MRI 772 Biosafety Committee and Animal Ethics Committee (project P3600), and are conducted in 773 accordance with the "Australian Code for the care and use of animals for scientific purposes" as 774 defined by the National Health and Medical Research Council of Australia. Work was conducted 775 in a dedicated suite in a biosafety level-3 (PC3) facility at the OIMR Berghofer MRI (Australian 776 Department of Agriculture, Water and the Environment certification Q2326 and Office of the Gene 777 Technology Regulator certification 3445). Heterozygous K18 hACE2-transgenic mice (The 778 779 Jackson Laboratory, Bar Harbour, ME, USA) were bred in-house by crossing with C57BL/6J mice (Animal Resources Center, Canning Vale, WA, Australia). DNA from tail tips was isolated using 780 Extract-N-Amp Tissue PCR kit (Sigma) and PCR genotyping undertaken as described (The Jackson 781 Laboratory. Genotyping protocols database. B6.Cg-Tg(K18-ACE2)2Prlmn/J. Stock No: 034860. 782 Protocol 38276), except using primers Forward - 5'-CTTGGTGATATGTGGGGTAGA-3', 783 reverse 5' CGCTTCATCTCCCACCACTT-3' (recommended by NIOBIOHN, Osaka, Japan). 784

785

Female 6-8 week old K18 hACE2 mice (n=4 per group) were treated once with 16 mg/kg of pixatimod in 200  $\mu$ l saline i.p. on the day before infection. Control K18 hACE2 mice received 200  $\mu$ l saline on the day before infection. Age and gender matched C57BL/6J mice (n=4) were included as a drug toxicity control and received 16 mg/kg of pixatimod in 200  $\mu$ l saline i.p., but were not infected with virus. 791 hCoV-19/Australia/QLD02/2020 (QLD02) (GISAID Accession ID; EPI ISL 407896) virus was used to inoculate K18 hACE2 mice (n=4 per group) intranasally with 10<sup>4</sup> CCID<sub>50</sub> of virus in 50 µl 792 medium while under light anaesthesia; 3% isoflurane (Piramal Enterprises Ltd., Andhra Pradesh, 793 India) delivered using The Stinger, Rodent Anesthesia System (Advanced Anaesthesia 794 795 Specialists/Darvall, Gladesville, NSW, Australia). Mice were weighed daily and euthanized on day 5 post-infection using  $CO_2$ . Tissues were fixed for histology or were weighed and then 796 homogenised using four ceramic beads at 6000 rpm twice for 15 s (Precellys 24 Homogeniser, 797 Bertin Instruments, Montigny-le-Bretonneux, France). After centrifugation for 30 mins, 2000 g at 798 4°C, virus titres in supernatants were determined by CCID<sub>50</sub> assays using Vero E6 cells. 799

300 Statistical analysis: Experimental data are presented as means  $\pm$  SD, SEM or CV as noted. Statistical analyses were performed using analysis of a two-tailed Student's t test with GraphPad 301 Prism (GraphPad Software) unless otherwise noted. Differences were considered statistically 302 significant if the *P* value was less than 0.05. Statistical analysis for the K18-hACE2 mouse work 303 was performed using IBM SPSS Statistics for Windows, Version 19.0 (IBM Corp., Armonk, NY, 304 USA). The t-test was used when the data was deemed parametric, i.e. difference in variances was 305 <4, skewness was >2 and kurtosis was <2. Otherwise the non-parametric Kolmogorov-Smirnov 306 test was used. 307

#### 308

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<del>)</del>89

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Author contributions: J.E.T., V.F., M.A.S. and K.D. conceived the project. S.E.G., C.J.M-W.,
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J.D.J.S., M.C., D.W., P.R.Y., A.A.K., M.A.L., E.A.Y., R.K., R.L.M., Y-H.C., Z.Y., E.T.,M.A.S.,
T.M.A.W. and A.S. designed and conducted the experiments and undertook analyses. V.F., E.H.,
K.D. M.W.C., J.S., T.B., M.A.S. and J.E.T. analyzed results and prepared the manuscript.

)11

Competing Interests: E.H. and K.D. are employees of Zucero Therapeutics. V.F., E.H. and K.D.
 are inventors on pixatimod patents.

Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

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# **Supplementary Materials**

# The clinical-stage heparan sulfate mimetic pixatimod (PG545) potently inhibits SARS-CoV 2 virus via disruption of the Spike-ACE2 interaction

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Edward Hammond, Keith Dredge, Miles W. Carroll, Edward Trybala, Tomas Bergström, Vito
Ferro, Mark A. Skidmore and Jeremy E. Turnbull

)46

# )47 **The file includes:**

Table S1

Here Figs. S1 to S4

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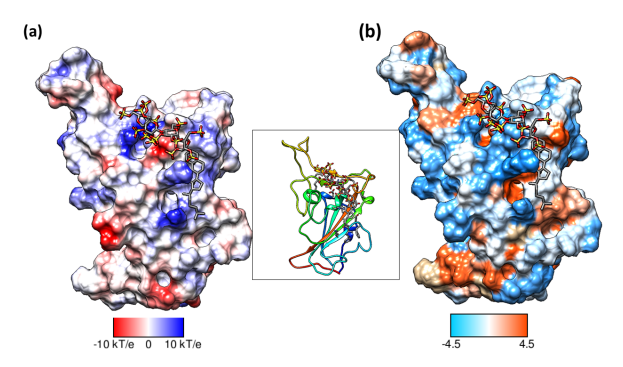
)55

# Table S1: Amino acids at position 614 in Spike protein of SARS-CoV-2 isolates

Isolate	Amino acids at position 614 in Spike	
VIC01	D	
QLD02	D	
QLD935	G	
DE-Gbg20	G	

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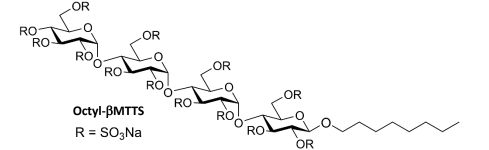
> )58 )59 )60 )61 )62 )63 )64 )65 )66



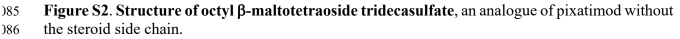
)67 )68

Figure S1. An alternate binding mode of pixatimod on the S1 RBD presented an unfavourable )69 total binding free energy. Surfaces are oriented in the same direction as shown in the ribbon )70 diagram in the inset. (a) Coulombic Surface Coloring defaults:  $\varepsilon = 4r$ , thresholds  $\pm 10$  kcal/mol·e )71 were used. Blue indicates surface with basic region whereas red indicates negatively charged )72 surface. (b) The hydrophobic surface was coloured using the Kyte-Doolittle scale wherein blue, )73 white and orange red colour indicates most hydrophilic, neutral and hydrophobic region, )74 respectively. UCSF Chimera was used for creating surfaces and rendering the images. Hydrogens )75 are not shown for clarity. )76

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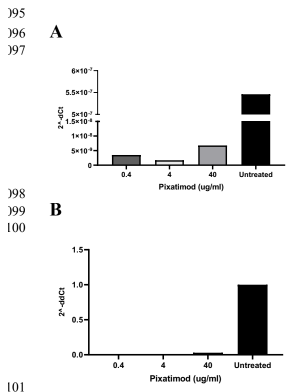
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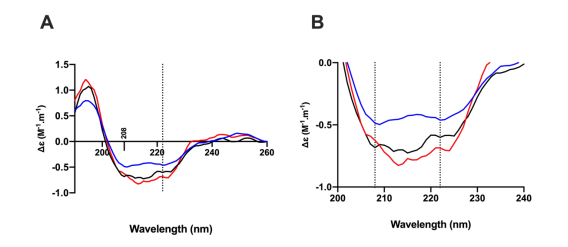


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**Figure S3: Pixatimod inhibits infection of human bronchial epithelial cells with SARS-CoV-2 virus.** Live virus infectivity assays of bronchial airway epithelial cells BCi-NS1.1, grown in an air liquid interface (ALI), were performed as described in Methods for the SARS-CoV-2 isolate VIC01. Viral shedding in the apical side (panel A) and viral load in infected cells (panel B) was measured by RT-PCR of viral RNA, as described in Methods. (representative data shown).

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Figure S4: The conformational change of the SARS-CoV-2 S1 RBD observed in the presence of pixatimod by circular dichroism (CD) spectroscopy. (A). Circular dichroism spectra (190 -260 nm) of SARS-CoV-2 S1 RBD alone (black solid line) and pixatimod (blue solid line) in PBS, pH 7.4. The red line represents the sum of the two individual spectra. Vertical dotted line indicates 222 nm (B) Details of the same spectra expanded between 200 and 240 nm. Vertical dotted lines

indicate 222 nm and 208 nm.