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A Multiparametric and High-Throughput Platform for Host-Virus Binding Screens

6 Authors

Title

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24 Abstract

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Speed is key during infectious disease outbreaks. It is essential, for example, to identify critical host 25 binding factors to the pathogens as fast as possible. The complexity of host plasma membrane is 26 often a limiting factor hindering fast and accurate determination of host binding factors as well as 27 high-throughput screening for neutralizing antimicrobial drug targets. Here we describe a multi-28 parametric and high-throughput platform tackling this bottleneck and enabling fast screens for host 29 binding factors as well as new antiviral drug targets. The sensitivity and robustness of our platform 30 was validated by blocking SARS-CoV-2 spike particles with nanobodies and IgGs from human 31 serum samples. 32

3334 Teaser

35 A fast screening platform tackling host-pathogen interactions.

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45 MAIN TEXT

46 Introduction

Emerging microbial pathogens, such as bacteria, fungi and viruses, tremendously challenge human 47 health and cause significant economical and societal burden worldwide. Therefore, tools facilitating 48 and improving pandemic preparedness are of uttermost importance to minimize these negative 49 50 effects. Current state-of-the-art methods, such as enzyme-linked immunosorbent assay (ELISA), reverse transcription-polymerase chain reaction (RT-PCR) and RT loop-mediated isothermal 51 amplification (RT-LAMP) usually rely on bulk measurements resulting in a single readout-value 52 (1). In addition, during the peaks of SARS-CoV-2 pandemic, RT-PCR instruments were used to 53 capacity slowing down pandemic surveillance and highlighting the need for additional readout-54 55 systems. Especially flow cytometry, enabling fast and high-throughput measurements of complex mixtures, is widely used in clinics for immunophenotyping and would be an attractive and broadly 56 available technique for such purposes (2). 57

To complement existing bulk measurement methods, we aimed to develop a fast and high-58 59 throughput platform to study host-pathogen interactions. The system should not only reconstitute host cell proteins, but also the lipid bilayer, which is mostly neglected in current state-of-the-art 60 methods but often hosts important attachment factors. However, the complexity of the mammalian 61 plasma membrane consisting of thousands of different lipids and proteins embedded between an 62 outer glycocalyx and inner cortical cytoskeleton is overwhelming. This complexity not only slows 63 64 down our efforts to identify important interaction partners but also obscures specific interactions between host and pathogen due to the plenitude of involved molecules and interactions. To 65 overcome this bottleneck and reduce complexity, bottom-up model membrane systems are 66 attractive alternatives which allow for precise control over composition and properties. Among 67 these, planar supported lipid bilayer systems (SLBs) were widely used (3) but do not account for 68 69 cells' three-dimensional nature. Three-dimensional model systems, such as large unilamellar vesicles (LUVs), giant unilamellar vesicles (GUVs), and cell-derived giant plasma membrane 70 vesicles (GPMVs) help to recreate cellular curvature but are challenging to use in high-throughput 71 flow cytometry because of their fragility and size-inhomogeneity. 72

73 For this reason, we coated cell-sized 5µm silica beads with a lipid bilayer consisting of 98 mole percent 1-palmitoyl-2-oleoyl-glycero-3-phosphocholine (POPC) doped with 2 mole percent of a 74 nickelated anchoring lipid (18:1 DGS-NTA(Ni)). Next, we attached His-tagged host-cell proteins 75 of interest to membrane-coated beads to generate functionalized bead-supported lipid bilayers 76 (fBSLBs) serving as minimal synthetic host-cells (Fig. 1A). In contrast to methods relying on 77 random surface-adsorption, fBSLBs ensure proper protein orientation, tightly controllable receptor 78 mobility and density as well as molecular interactions at the membrane plane. In addition, the 79 presence of a hydrophobic lipid bilayer more closely mimics the cellular environment and enables 80 to discriminate between binding preferences of pathogens to either host-cell proteins or lipids. For 81 example, surface proteins of several viruses can bind different host-cell lipids facilitating cellular 82 uptake and shaping viral tropism (4). 83

In this study, we show that fBSLBs carrying different host cell receptors, such as angiotensinconverting enzyme 2 (ACE2), can serve as highly diverse platform to screen for unknown proteinand lipid-binding molecules, drugs influencing host-pathogen interactions and the blocking efficiency of neutralizing antibodies present in human serum samples. Its fast implementation, easy adaptability of multiple parameters and high-throughput capability propel our method as an important platform to understand and tackle host-pathogen interactions.

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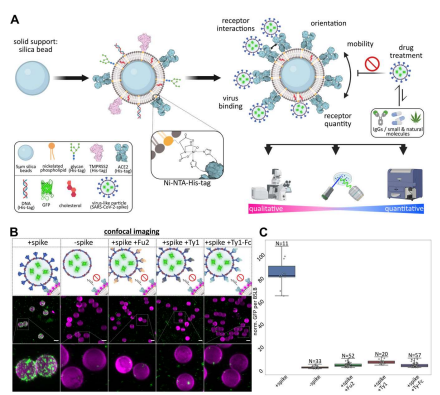
91 **Results**

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93 **fBSLBs enable fast and qualitative host-pathogen interaction studies**

94 Upon coating of 5µm silica beads with POPC:DGS-NTA(Ni) 98:2 mole percent of liposome solution, we verified proper bilayer formation by measuring diffusion of a fluorescent lipid 95 analogue using fluorescence correlation spectroscopy (FCS) (fig. S1, A and B), which matched 96 with previous data (5, 6). We first generated fBSLBs carrying ACE2 and studied their interaction 97 with SARS-CoV-2 spike expressing virus-like particles (+S-VLPs) using confocal microscopy 98 (Fig.1, B and C). To quantify VLP-binding per bead we developed an automated image analysis 99 workflow using Fiji (7) (fig. S2). While there was strong interaction between ACE2-fBSLBs and 100 +S-VLPs, it was absent in VLPs with no spike (-S-VLPs) and +S-VLPs pre-treated with SARS-101 CoV-2 neutralizing spike nanobodies which were shown to be potent tools to neutralize SARS-102 CoV-2 by blocking the interaction between spike receptor-binding domain (RBD) and its host 103 104 receptor ACE2 (8, 9). Thus, fBSLBs can serve as powerful screening platform to identify efficient inhibitors with therapeutic potential. 105





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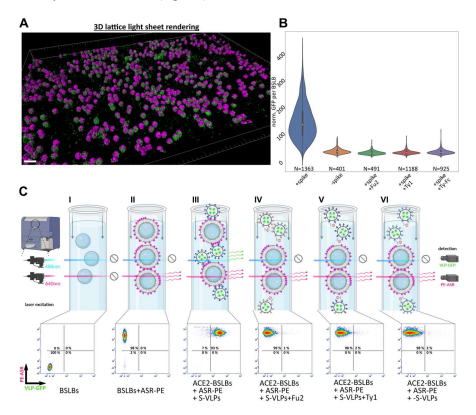
109 Fig. 1. Design and characterization of our multi-parametric and high-throughput platform based on fBSLBs to 110 study host-virus interactions. (A) Scheme depicting the bottom-up assembly of fBSLBs and available readout 111 techniques. (B) LSM maximum-intensity projections of BSLBs (magenta) and VLPs (green) showing specific 112 interaction between SARS-CoV-2 spike VLPs (+S-VLPs) and ACE2-fBSLBs. (C) Quantification of viral GFP-signal 113 per fBSLBs of each condition from (b) shows specific attachment of +S-VLPs to ACE2-fBSLBs (median=84.05, N=11) and no interaction between -S-VLPs and ACE2-BSLBs (median=2.88, N=33) and nanobody-pretreated +S-114 115 VLPs and ACE2-BSLBs (Fu2: median=4.70, N=52; Ty1: median=7.77, N=20; Ty-Fc: median=4.41, N=57). Boxplot 116 with overlay of individual data points, median as black center line, box showing the quartiles and whiskers from 117 minimum to maximum value. Illustrations were created using Biorender.com and Inkscape. 118

119 fBSLBs enable quantitative high-throughput screening

To increase number of data points and decrease acquisition time, we performed fast, quantitative,
3D lattice light-sheet microscopy (LLSM) and quantified viral loads per fBSLB (Fig. 2, A and B)
which confirmed confocal microscopy data. To screen several tens of thousands of fBSLBs within

minutes, fast and high-throughput flow cytometry can be used thanks to the firm nature of fBSLBs. 123 Individual fBSLBs were easily detected by their specific scattering signal and presence of the lipid 124 bilayer confirmed by 1.2-dioleoyl-sn-glycero-3-phosphoethanolamine Abberior STAR RED (ASR-125 PE) labelling while VLPs were labelled with eGFP. Upon addition of ASR-PE and +S-VLPs to 126 ACE2-fBSLBs, we observed a strong increase of fluorescence intensity per bead both in virus 127 (green) and in membrane (red) channels (Fig. 2C). Moreover, virus signal decreased significantly 128 upon nanobody treatment, confirming the neutralizing ability of nanobodies. Hence, fBSLBs enable 129 to study host-virus interactions using quantitative high-throughput flow cytometry which is usually 130 not feasible due to the small size of viral particles. Moreover, it serves as a powerful platform to 131 study concentration-dependent effects of molecules on the binding between viruses and host-cell 132 receptors. To show this, we determined optimal concentrations of ACE2 on the fBSLBs and the 133 amount +S-VLPs by titration series (fig. S3). 134

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138 Fig. 2. High-throughput measurements using fBSLBs. (A) Fast and quantitative LLSM enabling big-volume renderings of ACE2-fBSLBs (magenta) interacting with +S-VLPs (green). (B) Quantification of VLP-GFP signal per 139 bead proves specific interaction between +S-VLPs and ACE2-fBSLBs (N>400). Violin plot with miniature boxplot 140 141 showing quartiles and median as white dot. +S-VLPs show significant increased binding to ACE2-fBSLBs as compared 142 to the other groups (p-value<0.0001). (C) Fast high-throughput screening of interaction between VLPs and ACE2-143 fBSLBs using flow cytometry. Strong signal of the fluorescent lipid ASR-PE (y-axis) confirms functional bilayer formation and interaction of VLPs with fBSLBs can be followed by intensity changes in the VLP-GFP channel (x-axis) 144 145 (N>8500 per condition). Illustrations were created using Biorender.com and Inkscape.

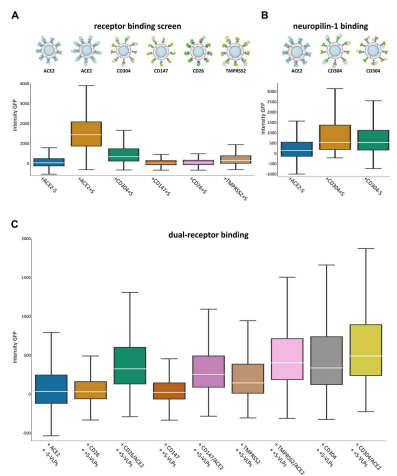
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147 Screening receptors using fBSLBs

Besides ACE2, other receptors have been described to contribute to SARS-CoV-2 binding to the host-cell surface and subsequent infection. For this reason, we tested interaction of +S-VLPs with reported host-cell receptors Neuropilin-1 (CD304) (*10*, *11*), Basigin (CD147) (*12*), DPP4 (CD26) (*13*, *14*) and TMPRSS2 (*15*, *16*) using fBSLBs in combination with flow cytometry. As expected, +S-VLPs showed strongest interaction with ACE2-fBSLBs (Fig. 3A). Interestingly, +S-VLPs also interacted with CD304-fBSLBs and TMPRSS2-fBSLBs, confirming that these two proteins act as

host binding factors, but neither interaction was as strong as for ACE2-fBSLBs. No binding was 154 observed for CD147-fBSLBs or CD26-fBSLBs, suggesting that these proteins cannot act as host 155 binding factors alone and might require additional host-cell binding elements. Notably, CD304-156 fBSLBs binding to VLPs was independent of spike-protein on their surface, e.g., -S-VLPs also 157 bound to CD304-fBSLBs effectively while they did not bind any other proteins we tested (Fig. 3B). 158 This suggests the presence of another interaction partner on the viral particles to this receptor. To 159 check this hypothesis, we performed dual-receptor screens with each individual receptor in absence 160 or presence of same molar concentration of ACE2 (Fig. 3C). The presence of ACE2 always 161 significantly increased the interaction of +S-VLPs with fBSLBs, but the overall strongest binding 162 was observed in the simultaneous presence of CD304 and ACE2, supporting the idea of two 163 different additive binding mechanisms. 164

fBSLBs allow tight control not only on the composition of surface proteins but also of lipid 165 composition. We made use of this and screened for reported lipid co-receptors for spike, such as 166 GM1 gangliosides (17). Despite varying GM1 concentrations in fBSLBs, we could not observe any 167 concentration-dependent binding of VLPs pseudotyped with spike, beta-spike, delta-spike, Ebola 168 169 virus glycoprotein (GP) or without any viral protein (fig. S4, A and B). These results highlight the need for additional high-affinity host-cell binding factors for efficient virus-host interaction. 170



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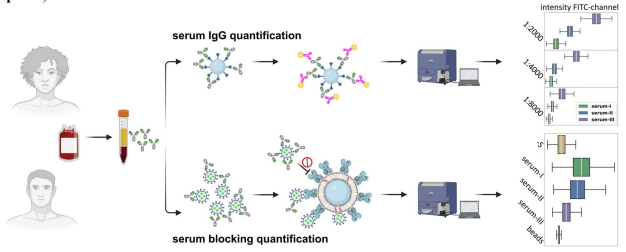
172 173 Fig. 3. Receptor screening using fBSLBs. (A) Application of fBSLBs to study interaction with different host-174 receptors. Scheme depicting fBSLBs with different His-tagged host-receptors. Box-and-whisker plots showing the 175 distribution of +S-VLP-GFP signal of 40000 BSLBs analyzed by flow cytometry. Besides ACE2 (median: 1440), 176 specific but less pronounced binding was also observed for CD304 (median: 337) and TMPRSS2 (median: 145). Except 177 for the dataset-pair +ACE2-S (blue) and +CD26+S (magenta) all populations are significantly different from each other 178 (p-value<0.0001). (B) Interaction of VLPs with CD304-fBSLBs in the absence of spike protein (N=20000). All 179 populations are significantly different from each other (p-value<0.0001). (C) Dual receptor screen using fBSLBs and 180 flow cytometry. No interaction between +S-VLPs and the host-cell receptors CD26 (median: 31) and CD147 (median:

181 23) was detected, respectively. Increased interaction with TMPRSS2 (median: 145) and CD304 (Neuropilin-1, median: 182 337) fBSLBs was observed, respectively. Upon coating BSLBs with 1:1 molar ratio of ACE2 and different host-cell 183 receptors all interactions were further increased while the receptor pairs TMPRSS2/ACE2 (median: 403) and 184 CD304/ACE2 (median: 491) showed strongest binding of +S-VLPs (N=40000 per condition). All populations are 185 significantly different from each other (p-value<0.0001). Box plots show inter-quartile range with white median line 186 and whiskers extend to 1.5 inter-quartile range.

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188 Surveillance of human serum samples using fBSLBs

Key for pandemic containment is surveillance of convalescent serum samples and their ability to 189 block the interaction between virus and host cell receptors. Virus-specific antibody levels in human 190 serum are usually proportional to neutralization of the virus and can be used to predict disease-191 192 outcome or the need for additional booster vaccinations (18, 19). Moreover, it is very important to understand whether anti-viral IgGs in prevalent serum samples still protect from upcoming new 193 variants to decide for vaccine-adjustments and therapeutic treatment options. To show the potential 194 of our method to answer these questions, we first determined the amount of spike-IgGs in three 195 human serum samples using a bead-based assay in combination with flow cytometry (Fig. 4, upper 196 197 panel). Glass beads were coated with recombinant spike receptor binding domain (RBD), incubated with serum samples, and anti-spike IgGs detected by labelling with secondary dye-conjugated anti-198 human antibodies. After we determined the relative levels of anti-spike IgGs in the three serum 199 samples, we blocked +S-VLPs with the different serum samples and studied the interaction with 200 ACE2-fBSLBs. The amount of anti-spike IgGs perfectly correlated with the blocking efficiency, 201 highlighting the ability of this method as powerful tool for pandemic surveillance (Fig. 4, lower 202 203 panel).



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206 Fig. 4. Application of fBSLBs to study blocking efficacy of neutralizing antibodies in human blood serum 207 samples. Scheme showing the processing of human serum samples to quantify amount of anti-spike IgGs and their capacity to block the interaction between +S-VLPs and ACE2-fBSLBs. Upper box plots illustrating the amount of anti-208 209 spike IgGs in three different serum samples at different dilutions (N=10000). Lower box plots showing the efficiency 210 to block the interaction between +S-VLPs and ACE2-BSLBs for the three different serum samples of each population 211 (N=10000). Note that the amount of anti-spike-IgGs in the serum samples correlates with blocking efficiency. All 212 populations are significantly different from each other (p-value<0.0001). Box plots show inter-quartile range with white 213 median line and whiskers extend to 1.5 inter-quartile range. Illustrations were created using Biorender.com and 214 Inkscape.

216 **Discussion**

217 Quick response to pandemic outbreaks is of uttermost importance for disease and damage control. 218 Our platform relies on material and molecules which are available from early pandemic onset, such 219 as the sequence of viral structural proteins and potential interaction partners. Exploiting highly 220 specific Ni-NTA-His-tag conjugation makes the platform highly versatile and accessible, since this

chemistry is widely used for protein purification and His-tagged proteins are available from a 221 myriad of commercial resources. Screening of potential host-cell receptors and co-receptors, 222 including lipids, can be done within a few hours using qualitative and high-throughput quantitative 223 readout platforms. In contrast to other methods, our method enables tight control of multiple cellular 224 parameters such as lipid composition, receptor mobility, receptor orientation, receptor-receptor 225 interactions, and local receptor densities. The platform allows to determine serum-virus 226 neutralization capacity in a safe laboratory environment within hours. Moreover, the presence of a 227 lipid bilayer more closely mimics the cellular environment and can help to entangle the complex 228 interplay between virus-receptor and virus-bilayer interactions which are often difficult to 229 discriminate. Due to its highly defined bottom-up assembly, fBSLBs are not prone to cellular 230 heterogeneity, e.g. due to differences in cell-cycle states, transcription and translation, which can 231 complicate drug screens. 232

However, this cellular heterogeneity could fine-tune host-pathogen interactions which is challenging to reproduce with our platform. Recent advances on coating beads with native cellular membranes would be an opportunity to recreate this complexity (20, 21). Another limitation of fBSLBs is its inability to initially detect cellular toxic compounds. This can also be advantageous since substances showing both cellular toxicity and binding inhibition are identified and not directly discarded. Further efforts in reducing cellular toxicity while maintaining inhibitory effects of these molecules would be an exciting way to find new drug targets.

Our broadly accessible platform enables to perform fast and high-throughput drug screens and to discriminate whether drugs act on the virus particles or on the host-cell receptors. Due to its bottomup design, our method should be readily extensible to other biomolecules (e.g. glycocalyx, DNA, RNA) and pathogens (bacteria, fungi) making it a valuable tool for future pandemic preparedness.

- 245 Materials and Methods
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247 **fBSLB** preparation

fBSLBs were prepared similar as described previously (6). For preparation of one batch of fBSLBs, 248 1×10^7 5µm silica beads (Bangs Laboratories) were vortexed thoroughly and washed three times 249 with PBS using 1500xg and 30 seconds centrifugation steps. Beads were coated with lipid bilayers 250 of defined compositions by incubation with 100µl 0.5mg/ml liposomes shaking at 1400rpm for 30 251 minutes. Liposomes were formed by mixing lipids dissolved in chloroform, solvent evaporation 252 under a steam of nitrogen, re-hydration, and tip-sonication (Branson Sonifier 250). To prepare 253 fBSLBs with His-tagged proteins, a lipid mixture consisting of 98mol% 16:0-18:1 POPC and 254 2mol% 18:1 DGS-Ni:NTA (Avanti Polar Lipids) was used. After bilayer formation beads were 255 washed two more times with PBS and 5pmol of His-tagged proteins added (Sino Biological: ACE2-256 His 10108-H08H, Neuropilin-1-His 10011-H08H, CD147-His 10186-H08H, CD26-His 10688-257 H08H). After 20 minutes on a rotary shaker the bilayer of fBSLBs was optionally directly labelled 258 with a fluorescent lipid analogue followed by 2 washing steps with PBS. Final fBSLBs were diluted 259 in 500µl PBS and used the same day. To study host-virus interactions, 20µl of fBSLBs were mixed 260 with 15µl of GFP-tagged pseudotyped VLPs and incubated for 30 minutes on a rotary shaker at 261 room temperature and directly used for microscopy or flow cytometry. Optionally, VLPs were pre-262 treated for 20 minutes on ice with 2µM Ty1, Ty1-Fc or Fu2 nanobodies4,5. 263

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266 VLP preparation

Mycoplasma-free HEK293T cells were cultured in DMEM supplemented with 10% FCS and grown
to ~70% confluency in T75 cell culture flasks. To produce VLPs, cells were co-transfected using
Lipofectamine 3000 and 15µg of DNA encoding for viral protein (pCMV14-3X-Flag-SARS-CoV2 S was a gift from Zhaohui Qian - Addgene plasmid # 145780; delta/beta spike expression plasmid

kindly provided by Benjamin Murrell; Ebola GP expression plasmid kindly provided by Jochen
Bodem), 7.5µg DNA encoding for HIV Vpr-GFP (NIH HIV Reagent Program, Division of AIDS,
NIAID, NIH: pEGFP-Vpr, ARP-11386, contributed by Dr. Warner C. Greene), and 7.5µg encoding
for a lentiviral packaging plasmid (psPAX2 was a gift from Didier Trono - Addgene plasmid #
12260). Media was exchanged after 12 hours and VLPs harvested after 24 and 48 hours and
enriched fiftyfold using LentiX concentrator according to the protocol provided by the
manufacturer (Takara).

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279 Microscopy and Quantification

After incubation with pseudotyped VLPs, fBSLBs were put into chambered glass coverslips 280 (IBIDI: 81817) and imaging performed in PBS. Confocal microscopy was performed using a C-281 Apochromat 40x/1.20 water immersion objective of the Zeiss LSM780 microscope. Viral GFP was 282 excited using 488nm argon laser and membrane-inserted ASR-PE was excited using a 633nm 283 helium neon laser, while emission was collected from 498-552nm and 641-695nm, respectively. 284 Full surface of 5µm fBSLBs was recorded by acquiring z-stacks with 24 slices each 0.3µm and 285 VLP-GFP signal per bead quantified using ImageJ following the provided macro and automated 286 workflow of Suppl. Fig. 02. To acquire fast, gentle, and big 3D volumes we used LLSM (Zeiss 287 Lattice Lightsheet 7) with 488nm and 640nm laser excitation for viral GFP and ASR-PE, 288 respectively. The general analysis workflow followed the one for confocal data, but parameters 289 were adjusted for differences in signal intensity. 290

292 Flow Cytometry

Upon interaction of VLPs with fBSLBs the mixture was diluted in 500µl PBS and transferred into flow tubes. Flow cytometry was performed using a BD Fortessa system acquired at low speed and 488nm (FITC) or 640nm (APC) excitation/emission settings used for VLP-GFP and ASR-PE, respectively. 10 000 to 20 000 events were acquired and analysed using FCS Express 7 and Python (FCSParser). Gating was only performed for data shown in Fig. 1f on singlet bead population clearly visible in the forward- versus side-scatter plot. This population was always at least 85% of the total bead population.

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301 Serum Blocking

Human blood from healthy donors was obtained from blood transfusion station of Karolinska 302 Hospital and serum prepared by centrifugation. The serum was aliquoted and frozen for further later 303 use. To determine the amount of anti-spike IgGs in serum samples, 1x107 5µm silica beads (Bangs 304 Laboratories) were washed three times with PBS and coated for 30 minutes with 47pmol SARS-305 CoV-2 RBD (BioSite: 40592-V08H) on a rotary shaker. After two washing steps with PBS beads 306 were resuspended in 500ul PBS supplemented with 4mg/ml BSA to block non-specific interaction 307 sites. 20µl of beads were incubated with stated serum dilutions over night at 4°C on a rotary shaker 308 to enable interaction of anti-spike IgGs with coated beads. After two washing steps, anti-spike IgGs 309 were labelled by incubation with 4µg/ml secondary anti-human IgG Alexa Fluor 488 antibodies 310 (ThermoFischer: A11013) for one hour at room temperature on a rotary shaker in the dark. Labelled 311 beads were washed and signal intensity of at least 9000 beads determined by flow cytometry. To 312 test serum blocking efficiency, VLPs were pre-treated with stated serum concentrations over night 313 at 4°C on a rotary shaker before incubated with ACE2-fBSLBs as described above. 314

316 Statistical Analysis

Visualization and statistical analysis of the data was performed using Python (Anaconda Navigator 2.3.2, JupyterLab 3.2.9) and Kruskal-Wallis H-test with post hoc pairwise test for multiple comparisons (Dunn's test with Bonferroni one-step correction). Standard error of the median was estimated by multiplying the standard error of the mean with the constant 1.253.

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434435 Competing interests:

- 436 Authors declare that they have no competing interests
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438 Data and materials availability:

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