1	A bacteria-based assay to study SARS-CoV-2 protein-protein interactions
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3	Running title: Bacteria-based SARS-CoV-2 protein interactome
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## 13 Abstract

14 Methods for detecting and dissecting the interactions of virally encoded proteins are essential 15 for probing basic viral biology and providing a foundation for therapeutic advances. The dearth 16 of targeted therapeutics for the treatment of COVID-19, an ongoing global health crisis, 17 underscores the importance of gaining a deeper understanding of the interactions of SARS-18 CoV-2-encoded proteins. Here we describe the use of a convenient bacteria-based two-hybrid 19 (B2H) system to analyze the SARS-CoV-2 proteome. We identify sixteen distinct intraviral 20 protein-protein interactions (PPIs), involving sixteen proteins. We find that many of the identified 21 proteins interact with more than one partner. We further show how our system facilitates the 22 genetic dissection of these interactions, enabling the identification of selectively disruptive 23 mutations. We also describe a modified B2H system that permits the detection of disulfide bond-24 dependent PPIs in the normally reducing *Escherichia coli* cytoplasm and we use this system to 25 detect the interaction of the SARS-CoV-2 spike protein receptor-binding domain (RBD) with its 26 cognate cell surface receptor ACE2. We then examine how the RBD-ACE2 interaction is 27 perturbed by several RBD amino acid substitutions found in currently circulating SARS-CoV-2 28 variants. Our findings illustrate the utility of a genetically tractable bacterial system for probing 29 the interactions of viral proteins and investigating the effects of emerging mutations. In principle, 30 the system could also facilitate the identification of potential therapeutics that disrupt specific 31 interactions of virally encoded proteins. More generally, our findings establish the feasibility of 32 using a B2H system to detect and dissect disulfide bond-dependent interactions of eukaryotic 33 proteins.

## 34 Importance

35 Understanding how virally encoded proteins interact with one another is essential in elucidating 36 basic viral biology, providing a foundation for therapeutic discovery. Here we describe the use of 37 a versatile bacteria-based system to investigate the interactions of the protein set encoded by 38 SARS-CoV-2, the virus responsible for the current pandemic. We identify sixteen distinct 39 intraviral protein-protein interactions, involving sixteen proteins, many of which interact with 40 more than one partner. Our system facilitates the genetic dissection of these interactions, 41 enabling the identification of selectively disruptive mutations. We also describe a modified 42 version of our bacteria-based system that permits detection of the interaction between the 43 SARS-CoV-2 spike protein (specifically its receptor binding domain) and its cognate human cell 44 surface receptor ACE2 and we investigate the effects of spike mutations found in currently 45 circulating SARS-CoV-2 variants. Our findings illustrate the general utility of our system for 46 probing the interactions of virally encoded proteins.

## 47 Introduction

48 The causative agent of COVID-19, SARS-CoV-2, like SARS-CoV (hereafter SARS-CoV-49 1) and the Middle East respiratory syndrome coronavirus (MERS-CoV), is a zoonotic pathogen 50 that belongs to the genus of  $\beta$ -coronaviruses [1,2]. A ~30 kb single stranded (+)-sense RNA 51 virus, SARS-CoV-2 encodes 16 non-structural proteins (Nsp1-Nsp16), which are transcribed 52 from two major open reading frames (ORF1a and ORF1b) and later post-translationally 53 processed by proteases to give rise to the individual Nsps [3]. The main function of the Nsps is 54 to provide and maintain the replication and transcription complex (RTC), promoting viral RNA 55 synthesis by the RNA-dependent RNA polymerase Nsp12 [3]. However, the Nsps have also 56 been implicated in other viral processes such as host innate immune system evasion - for 57 example, by suppressing aspects of the interferon response [4]. The virus also encodes four 58 structural proteins, the membrane (M) protein, the nucleocapsid (N) protein, the envelope (E) 59 protein and the spike (S) glycoprotein, and at least six accessory proteins (ORF3a, ORF6, 60 ORF7a, ORF7b, ORF8 and ORF10) [5]. The main function of coronavirus structural proteins is 61 to mediate cell entry, virus particle assembly and release from the host cells by budding, though 62 like the Nsps, structural proteins also participate in immune evasion. By contrast, the accessory 63 proteins are non-conserved and highly variable among different coronavirus species; although their functional roles remain largely unknown, they too have been associated with immune 64 65 evasion and disease severity [3].

Given the ongoing global crisis caused by the SARS-CoV-2 pandemic and the continuing need for targeted therapeutics for the treatment of COVID-19, understanding the intraviral and viral-host protein-protein interactions (PPIs) of SARS-CoV-2 remains a priority. An extensive virus-virus and host-virus PPI study recently highlighted the importance of Nsp10 as a potential inducer of the so-called cytokine storm (a dysregulated and hyperactive immune response) [6], thought to be the main cause for severe disease outcome and death in COVID-19

patients [2]. Li *et al.* (2021) further identified Nsp8 as a SARS-CoV-2 PPI hub [6], promoting
interactions with other Nsps, accessory proteins, and one structural protein. Similar
observations were previously also obtained for SARS-CoV-1 Nsp8 [7]. These findings suggest
that Nsp8 and Nsp10 might provide particularly efficacious targets for drug development.

76 The SARS-CoV-2 spike protein, which is present on the viral surface as trimers, consists 77 of two functionally distinct subunits, S1 and S2 [8,9]. The membrane-distal S1 subunit uses its 78 receptor-binding domain (RBD) to initiate the process of viral entry into human host cells by 79 binding to the cell surface protein angiotensin-converting enzyme 2 (ACE2), which also serves 80 as the receptor for SARS-CoV-1 but not for the more distantly related MERS-CoV. Following 81 ACE2 binding, the membrane-localized host cell serine protease TMPRSS2 cleaves the spike 82 protein at a specific site, triggering a series of dramatic conformational changes in the S2 83 subunit, which in turn mediate fusion of the viral and host membranes, enabling viral entry [10]. 84 As well as being a critical determinant of viral tropism, the RBD is a major target for SARS-CoV-85 2-neutralizing antibodies, including those identified from convalescent patient peripheral blood 86 mononuclear cells and those elicited by current (spike-based) vaccines [9,11-18].

87 Compared with those of other RNA viruses, the mutation rate of SARS-CoV-2 is 88 considered low-moderate (6-9x10<sup>-4</sup> bases/genome/year) [19-21], although others have pointed 89 out that multiple identical mutation hotspot events occurring at different points in time could lead 90 to an underestimation of the overall mutation rate [22]. Nevertheless, the pandemic has given 91 rise to a proliferation of variant lineages, including those designated variants of concern (VOCs) 92 by the World Health Organization (WHO), based on one or more of the following criteria: 93 increase in transmissibility; increase in virulence; decrease in effectiveness of public health 94 measures, diagnostics, therapeutics or vaccines (www.who.int/en/activities/tracking-SARS-CoV-95 2-variants). All of the VOCs carry spike mutations, including one or more that localize to the

96 RBD, motivating efforts to gain a systematic understanding of the effects of RBD amino acid
97 substitutions on ACE2 binding [23].

98 Here we employ a bacterial two-hybrid (B2H) system [24,25] to study the PPIs of SARS-99 CoV-2 in a heterologous non-eukaryotic system. Using this system, we describe a bacteria-100 based intraviral interactome. We further demonstrate the utility of the bacterial system for 101 genetically dissecting the SARS-CoV-2 PPIs by identifying mutations that selectively affect one 102 or another interaction. In addition, we describe a modified B2H system that allows us to detect 103 disulfide bond-dependent PPIs in the otherwise reducing Escherichia coli cytoplasm. We use 104 this system to detect the spike RBD-ACE2 interaction and to investigate the effects of mutations 105 found in VOCs. Our findings set the stage for further investigations of viral PPIs in a convenient 106 and genetically tractable bacterial system, as well as establishing the feasibility of using our 107 modified system to detect and dissect disulfide bond-dependent PPIs of other eukaryotic 108 proteins.

109

## 110 Results

## 111 Bacterial two-hybrid system to detect interactions of SARS-CoV-2 proteome

112 Previous studies have used yeast two-hybrid (Y2H) systems, a mammalian two-hybrid system 113 and co-immunoprecipitation experiments (co-IPs) to investigate the SARS-CoV-1 and SARS-114 CoV-2 protein interactomes, identifying overlapping but also distinct interactions depending on 115 the employed system [6,7,26,27]. Compared with bacteria, yeast have a relatively slow growth 116 rate and are more difficult to culture and transform for labs that do not routinely work with yeast. 117 To provide a more accessible alternative to Y2H systems as well as the less commonly used 118 mammalian two-hybrid system, we here describe the successful use of a B2H system 119 developed in our lab (Fig. 1A) [24,25] to test for viral PPIs. We fused all NCBI-predicted E. coli 120 codon-optimized SARS-CoV-2 open reading frames (ORFs; listed in Fig. 1B, see also NCBI 121 accession #: NC\_045512.2) to the DNA binding protein CI of bacteriophage  $\lambda$  ( $\lambda$ CI) and to the 122 N-terminal domain of the  $\alpha$  subunit ( $\alpha$ NTD) of RNA polymerase (RNAP). We then tested each 123 SARS-CoV-2 ORF for interaction with the other SARS-CoV-2 ORFs and itself. Interaction 124 between two given ORFs (X and Y), fused to  $\alpha$ NTD and  $\lambda$ Cl, respectively, stabilizes the binding 125 of RNAP to the test promoter such that the magnitude of the *lacZ* reporter gene expression 126 correlates with the strength of the PPI (Fig. 1A).

## 127 Identification of the SARS-CoV-2 interactome using a B2H system

128 Using our B2H system, we initially tested each SARS-CoV-2 ORF against each other SARS-129 CoV-2 ORF and itself in biological duplicate. Protein pairs with at least a 2-fold activation of lacZ 130 over background in one of the replicates were selected for further analysis. The list of interacting 131 proteins was further refined by performing repeat experiments with three biological replicates for 132 each initially identified potential PPI pair. This resulted in a final list of sixteen interacting SARS-133 CoV-2 protein pairs, including four self-interactions (Fig. 2). Some of these interactions were 134 identified only with a specific fusion partner combination (*i.e.*, protein X fused to  $\alpha$ NTD and 135 protein Y fused to  $\lambda$ Cl, or the other way around), while others were fusion partner-independent 136 (*i.e.*, interaction between proteins X and Y regardless of their fusion to  $\alpha$ NTD or  $\lambda$ Cl). Self-137 interacting proteins (Nsp7, Nsp9, ORF6 and ORF10) were by definition fusion partner-138 insensitive; however, four other pairs of proteins (Nsp7+Nsp8, Nsp10+Nsp14, Nsp10+Nsp16, 139 Nsp3+N, and Nsp8+ORF6) also interacted detectably regardless of the fusion partner 140 (Supplementary Fig. 1).

Among the identified interacting pairs, several particularly strong PPIs were observed, including the Nsp7 self-interaction, Nsp7+Nsp8, Nsp10+Nsp16, N+Nsp3 and Nsp9+Nsp11 (Fig. 3). In fact, the Nsp10+Nsp16 pair interacted significantly more strongly than our positive control,

144 representing one of the strongest interactions we have ever measured with our B2H assay. For 145 our B2H assays, we routinely consider an interaction to be reliable when we detect at least a 146 two-fold increase in *lacZ* reporter gene expression (measured as  $\beta$ -galactosidase activity) over 147 the background (obtained with the negative controls). Applying this cut-off to our experimental 148 data, we identified several medium-to-weak interactions (2- to 5-fold increase over the negative 149 controls; Supplementary Fig. 2). The interactions of Nsp8+ORF7b and ORF10+ORF10 closely 150 missed the 2-fold cutoff but were nonetheless included in the list because a previous SARS-151 CoV-2 interactome study also identified those interactions (based on co-IP data) [6].

152 Comparison of our SARS-CoV-2 B2H data with the previously reported SARS-CoV-2 153 Y2H and co-IP data [6] revealed four PPIs that were shared among the three assay systems, 154 providing strong support for their biological relevance (Supplementary Fig. 3). These included 155 Nsp7+Nsp8, Nsp8+ORF10, Nsp10+Nsp14 and ORF6+ORF6. Others were identified either in 156 only one of the assay systems (*i.e.*, B2H, Y2H or co-IP) or in two assay systems (B2H and Y2H, 157 B2H and co-IP, or Y2H and co-IP) (Supplementary Fig. 3). Furthermore, some of our identified 158 interactions are validated by co-crystal structures. These included Nsp7+Nsp8 (Protein Data 159 Bank (PDB) accession number 6YHU [28]), Nsp10+Nsp14 (PDB: 5NFY from SARS-CoV-1 [29] 160 or more recently 7DIY from SARS-CoV-2 [30]), Nsp10+Nsp16 (PDB: 6W4H [31]) and the Nsp9 161 self-interaction (PDB: 6W9Q [32]). Notably, no self-interaction of Nsp9 was identified in a 162 previous Y2H and co-IP analysis of the SARS-CoV-2 interactome [6], highlighting the 163 importance of employing several different interaction assays when studying the interactome of a 164 given protein set to avoid loss of information due to experimental system idiosyncrasies.

Similar to previous observations for SARS-CoV-1 [7], we identified Nsp8 as a major SARS-CoV-2 interaction hub, interacting with six other SARS-CoV-2 ORFs (Fig. 3, Supplementary Fig. 2), consistent with a critical role for Nsp8 in SARS coronavirus biology. Nonetheless, most of the interaction partners we identified for Nsp8 in SARS-CoV-2 are

169 different than those identified previously for SARS-CoV-1 [7,26,27] (Supplementary Fig. 4). 170 Overall, only six PPIs were identified in our SARS-CoV-2 B2H analysis and at least one of three 171 independent SARS-CoV-1 Y2H studies, including two involving Nsp8 (Supplementary Fig. 4). 172 Notably, there are considerable differences between the results of the three previous Y2H 173 studies [7,26,27] and only three PPIs (Nsp8+Nsp7, Nsp10+Nsp14, and Nsp10+Nsp16) were 174 independently identified in two SARS-CoV-1 two-hybrid assays and our SARS-CoV-2 B2H 175 assay (Supplementary Fig. 4). This could reflect significant differences between the PPI 176 networks in SARS-CoV-1 and SARS-CoV-2 and/or differences in the assays themselves 177 (procedures and background organism).

# Targeted mutational screens identify interaction partner-specific sites of protein-protein interaction in CoV-2 proteins with more than one interaction partner

180 As a genetic assay, the B2H system facilitates the dissection of specific PPIs through both 181 targeted and random mutagenesis. Having established the utility of the B2H assay in testing for 182 viral PPIs, we next sought to use this assay to dissect the interactions of selected viral proteins 183 through targeted mutational analysis. Specifically, we chose proteins that interacted with more 184 than one partner and sought to disrupt the interaction of such a protein with one of its partners 185 while preserving its interaction with another. We initially selected Nsp10 with two known 186 interaction partners, Nsp14 and Nsp16, and attempted to disrupt only its interaction with Nsp14. 187 To identify suitable targets for mutagenesis, we analyzed the crystal structures of Nsp10-Nsp14 188 (PDB ID: 5NFY [29]) and Nsp10-Nsp16 (PDB ID: 6W4H [31]) and their protein-protein interfaces 189 using PDBePISA [33]. Based on this approach, we selected three sets of amino acid 190 substitutions likely to affect the binding of Nsp10 to Nsp14, while leaving its interaction with 191 Nsp16 intact (assuming that the substitutions do not result in allosteric effects). While the Nsp10 192 F16A/F19A/V21A set targeted the hydrophobic region, the Nsp10 T5A/T12A/S15A and 193 S29A/S33A sets partially disrupted the hydrogen bond network of the Nsp10-Nsp14 interface

194 (Fig. 4A,C). Each of the three multiply substituted Nsp10 mutants lost the ability to interact detectably with Nsp14 while maintaining an approximately wild-type interaction with Nsp16 (Fig. 195 196 4B). We note that the close approach of amino acid side chains at a protein-protein interface as 197 revealed by X-ray crystallography does not necessarily indicate that they participate in a 198 functionally important interaction. However, the loss of a detectable interaction between each of 199 the three Nsp10 mutants and Nsp14 in our B2H assay suggests that at least a subset of the 200 selected residues make stabilizing contacts. Furthermore, although the Nsp10-Nsp16 201 interaction serves as a control, we also confirmed that the introduced amino acid substitutions 202 were not generally destabilizing (Supplementary Fig. 5).

203 We then focused on Nsp16 with two interaction partners, Nsp10 and Nsp15, targeting 204 the Nsp16-Nsp10 pair, which displayed a significantly higher B2H signal than that of the Nsp16-205 Nsp15 pair. Here we also utilized the available crystal structure for the Nsp16-Nsp10 complex; 206 however, as there is no structure for the Nsp16-Nsp15 complex, the substitutions introduced 207 into Nsp16 were based solely on their predicted effects on its interaction with Nsp10. 208 Endeavoring to disrupt the Nsp16-Nsp10 interaction, we created two Nsp16 triple substitution 209 mutants, targeting hydrophobic (I40A/M41A/V44A) or hydrophilic (K76A/Q87A/D106A) contacts, 210 and a mutant with the six substitutions combined (Fig. 4C). The data reveal drastic effects of 211 these substitutions on the binding of Nsp16 to Nsp10, resulting in near background or 212 background levels of reporter gene expression for each of the mutants (Fig. 4D). The effects of 213 the same substitutions on the binding of Nsp16 to Nsp15 were modest and not statistically 214 significant. Notably, even though Nsp16 interacts much more weakly with Nsp15 than with 215 Nsp10, reporter gene expression was lower for each of the Nsp16 mutants in combination with 216 Nsp10 than when tested in combination with Nsp15 (Fig. 4D). We also confirmed that these 217 effects are not the result of altered protein levels (Supplementary Fig. 6). Together, these data

illustrate a proof-of-principle approach that can be used to obtain functionally informativemutants within a PPI network.

## 220 The B2H system as a tool to study circulating spike variants and their binding to ACE2

221 To further assess whether our B2H system can facilitate the study of emerging mutational 222 changes in viral populations, we next asked whether we could use our system to study the 223 interaction between the SARS-CoV-2 spike protein and ACE2. For this, we obtained an E. coli 224 codon-optimized gene fragment encoding the human ACE2 peptidase domain (aa 19-615, 225 hereafter ACE2). We inserted this gene fragment and a set of gene fragments encoding multiple 226 domains of the spike protein (including the RBD, aa 331-521) into our two-hybrid vectors, fusing 227 ACE2 and each of the spike domains to both  $\lambda$ CI and  $\alpha$ NTD. Initial experiments using our 228 standard E. coli B2H strain (FW102 OL2-62 termed B2H: Supplementary Table 1A) failed to 229 reveal an interaction of ACE2 with any of the selected spike domains (Supplementary Fig. 7; 230 data not shown). However, previous studies demonstrated that proper disulfide bond formation 231 is essential in order for spike and ACE2 to engage in a direct interaction [34,35]. Because the E. 232 coli cytoplasm is a reducing environment [36], we considered the possibility that the failure to 233 detect a spike-ACE2 interaction with our standard B2H strain might be due to a lack of proper 234 disulfide bond formation. To circumvent this obstacle, we modified a commercially available E. 235 coli strain (SHuffle from NEB, MA, USA) that permits the efficient expression and formation of 236 active full-length antibodies in the E. coli cytoplasm [37], adapting it for use with our two-hybrid 237 system (Methods). The SHuffle strain is deleted for two genes that encode cytoplasmic 238 reductases (trxB and gor) and also harbors the normally periplasmic disulfide bond isomerase 239 DsbC in the cytoplasm [38,39]. With this modified oxidizing strain (termed BLS148; 240 Supplementary Table 1A), we were able to detect an interaction of the spike RBD with ACE2 241 (Fig. 5A,B). Moreover, this interaction was abrogated when we mutated a pair of cysteine 242 residues (replacing them individually and in combination with serine residues) that engage in

disulfide bond formation within the RBD (C379 and C432) [34], consistent with the surmise that
the oxidizing strain permits detection of the RBD-ACE2 interaction by enabling appropriate
disulfide bond formation and correct folding of the interacting partners (Fig. 5B; Supplementary
Fig. 8).

247 Having adapted our B2H system for the study of disulfide bond-dependent PPIs, we 248 sought to test different spike (RBD) circulating variants for their abilities to bind ACE2. The RBD 249 amino acid substitutions included in our study are found in several SARS-CoV-2 variants that 250 were previously designated VOCs by the Centers for Disease Control and Prevention (USA; 251 https://www.cdc.gov/coronavirus/2019-ncov/variants/variant-info.html, initially accessed 252 05/30/2021). Specifically, we included the Alpha variant (B.1.1.7, first identified in the United 253 Kingdom) that carries the RBD N501Y substitution, the Beta variant (B.1.351, first identified in 254 South Africa) that carries the RBD K417N, E484K and N501Y substitutions, as well as the 255 Epsilon variant (B.1.429, first identified in California) that carries the RBD L452R substitution, 256 introducing the corresponding mutations into the spike RBD on our B2H vector (Fig. 5D). The 257 latter two variants have recently gained more attention as they are considered immune escape 258 variants, potentially resulting in a partial loss of immunity in previously infected or immunized 259 people [40-47]. In contrast, the Alpha variant is not characterized by a marked escape from 260 antibody neutralization [41,42,45–47]. Factors that are believed to contribute, potentially, to 261 immune escape include changes in the spike protein that: (i) enhance or stabilize its binding to 262 ACE2 or (ii) decrease the binding of specific anti-spike neutralizing antibodies [48-50].

As well as testing the Alpha, Beta and Epsilon RBDs for their abilities to bind ACE2, we included RBD mutants bearing component single and double substitutions from the Beta variant (Fig. 5C; Supplementary Fig. 9). We found that the N501Y substitution (in the context of the Alpha variant) had no observable effect on ACE2 binding. In contrast, the L452R substitution (Epsilon variant) resulted in a statistically significant increase in ACE2 binding. Substitutions

268 K417N, E484K and N501Y (Beta variant) together resulted in a significant reduction in ACE2 269 binding, as did the individual component substitutions K417N and E484K (with the E484K 270 substitution having the stronger effect). However, the effects of these two substitutions were 271 partially (E484K) or fully (K417N) abrogated when combined with the N501Y substitution. The 272 binding of the triply substituted variant was indistinguishable from that of the E484K/N501Y 273 double mutant, indicating that in this context the K417N substitution neither weakens or 274 strengthens the interaction. Together, these findings indicate that our modified B2H system 275 enables detection of disulfide bond-dependent PPIs and can be used to investigate the effects 276 of RBD variant substitutions on the RBD-ACE2 interaction.

277

#### 278 **Discussion**

## 279 Use of bacteria-based assay to investigate SARS-CoV-2 interactome

280 Here we use a versatile bacteria-based genetic tool for detecting and dissecting PPIs 281 [24,25] to screen the SARS-CoV-2 proteome for intraviral PPIs. We detected a total of sixteen 282 PPIs, including four self-interactions. Nine of these interactions were also detected in a previous 283 SARS-CoV-2 PPI study (Supplementary Fig. 3), as assessed by Y2H-based screens and/or 284 mammalian cell-based co-IP experiments [6]. Additionally, four of the interactions we detected 285 have been captured in co-crystal structures, including the Nsp9 self-interaction (PDB: 6W9Q; 286 [32]), which was not identified by either Y2H or co-IP analyses [6]. Of the six interactions we 287 detected that were not previously described in the context of SARS-CoV-2, three were 288 previously detected by Y2H analyses in the context of SARS CoV-1 (Supplementary Fig. 4). Of 289 the remaining three interactions, not previously described, two (Nsp9+Nsp11 and Nsp3+N) were 290 particularly strong as assessed in our B2H assay (Fig. 3).

291 Although the different assays that have been used to characterize the SARS-CoV-2 292 interactome have provided results that often corroborate one another, there are many examples 293 of interactions that have been detected with only one of the assays. These discrepancies 294 highlight the importance of employing multiple assay systems, each with its own inherent 295 limitations, to maximize the likelihood of obtaining a complete picture. Because of its 296 experimental accessibility, we expect that our B2H assay will be useful in evaluating other viral 297 proteomes, particularly by taking advantage of our oxidizing reporter strain that better 298 approximates the eukaryotic cell environment in allowing for proper disulfide bond formation in 299 the E. coli cytoplasm [51,52]. To further extend the spectrum of testable viral and eukaryotic 300 PPIs, the system could be augmented to enable the detection of phosphorylation-dependent 301 PPIs by introducing specific mammalian kinases into our reporter stain [53]. Given that many 302 mammalian (and presumably viral) proteins are constitutively phosphorylated in yeast [54-56], a 303 lack of properly phosphorylated proteins in our B2H system could explain, at least in principle, 304 why some SARS-CoV-2 PPIs were identified only in the Y2H screens [6] and not in our system 305 (Supplementary Fig. 3). We note, however, that a comprehensive phosphoproteomics analysis 306 of SARS-CoV-2-infected cells [57] suggests that other than interactions involving the N protein, 307 which was found to be phosphorylated at multiple sites, most of the viral PPIs that were 308 detected by Y2H analysis but not in our B2H system involve proteins that were not detectably 309 phosphorylated.

## 310 Genetic dissection of specific SARS-CoV-2 PPIs

A benefit of two-hybrid approaches for studying PPIs is that detected interactions can be readily dissected genetically, something that is particularly straightforward to do with our B2H system. As a proof-of-principle, we used a structure-based approach to investigate the effects of targeted mutations on specific SARS-CoV-2 PPIs, identifying substitutions that disrupt one interaction but not another. In addition to facilitating the evaluation of specific circulating or

316 targeted mutations, our B2H system can readily be adapted to screen for randomly generated 317 mutations that selectively affect one PPI and not another [58] when there is insufficient 318 information to make informed predictions from structural or other data. The identification of such 319 mutations could facilitate the functional analysis of particular PPIs and inform the choice of 320 potential drug targets for small molecule drug design. Furthermore, with a suitably modified 321 reporter strain to improve compound accessibility [59], compound or peptide libraries could be 322 screened to identify candidates that might target specific SARS-CoV-2 PPIs. It should also be 323 feasible to adapt our B2H reporter system for an *in vitro* cell-free protein expression system, 324 thereby facilitating compound screenings.

## 325 Use of oxidizing B2H reporter strain enables detection of RBD-ACE2 interaction

326 Based on previous studies, we anticipated a high-affinity interaction between the spike 327 RBD and ACE2 [9,23,34]. With our modified bacteria-based system, we found that the RBD-328 ACE2 interaction resulted in a roughly 3-fold increase in *lacZ* reporter gene expression over 329 background, a relatively modest effect. One possible explanation is that the  $\lambda$ CI-ACE2 fusion 330 protein is produced at relatively low levels compared with unfused  $\lambda CI$  and other  $\lambda CI$  fusion 331 proteins we have studied in the past (Supplementary Fig. 8), perhaps resulting in intracellular 332 concentrations insufficient to saturate the DNA-binding site on our *lacZ* reporter. Another 333 possible explanation (not mutually exclusive) lies in the fact that both the SARS-CoV-2 spike 334 protein and ACE2 are glycosylated in mammalian cells [9,60], with some studies suggesting that 335 glycan-side chain interactions may be important in stabilizing the RBD-ACE2 interaction [61,62]. 336 Thus, the interaction detected in our B2H system could be compromised by the lack of 337 mammalian-like N- and O-glycosylation in E. coli [63].

338 Our B2H system enabled us to assess the effects of specific RBD amino acid 339 substitutions that have been identified in globally circulating SARS-CoV-2 variants. We focused

specifically on three VOCs, as designated by the CDC at the time we initiated our study: the Alpha variant (B.1.1.7), the Beta variant (B.1.351) and the Epsilon variant (B.1.429), carrying RBD substitutions N501Y, N501Y/K417N/E484K, and L452R, respectively (see Fig. 5D) [41]. We note that as of September 21, 2021 each of these variants has been deescalated from a VOC to a variant being monitored (VBM) by the CDC. We also note that the highly contagious and rapidly proliferating Delta variant, currently designated as a VOC, harbors the L452R substitution in the RBD, together with a second substitution (T478K) [64].

347 Mutated in both the Alpha and the Beta variants, residue N501 is localized at the binding 348 interface with ACE2 [34,65] and many reports have suggested that the N501Y substitution 349 increases the affinity of the RBD for ACE2 [23,48,66-70] (but see [71] for a discrepant 350 prediction), potentially explaining the elevated infectivity of the Alpha variant. In a study in which 351 the effects of all possible RBD amino acid substitutions were examined using a yeast-surface-352 display platform, Starr et al. identified N501Y as one of the substitutions causing the highest 353 gain in ACE2-binding affinity [23]. In contrast, our B2H assay did not reveal any significant effect 354 of the N501Y substitution on the strength of the RBD-ACE2 interaction. Possibly this 355 discrepancy is due to the lack of glycosylation in the bacterial system; in fact, an ACE2 glycan 356 (N322) that has been reported to enhance RBD-ACE2 binding is part of the same binding patch 357 that includes N501 [62]. Nonetheless, we did observe a binding enhancing, compensatory effect 358 of the N501Y substitution when tested in the context of the Beta variant. That is, we found that 359 Beta-associated substitutions K417N and E484K both reduced ACE2 binding when tested 360 individually, and that the N501Y substitution compensated for these effects, partially in the case 361 of E484K and fully in the case of K417N. Our results thus suggest that substitutions K417N and 362 E484K, which have been implicated in significant immune escape [43-45,72-74], may impose a 363 cost on ACE2 binding that is compensated by the N501Y substitution (see also [70]). Consistent 364 with our findings, the K417N substitution has been previously reported to weaken ACE2 binding

365 [72,75,76]; however, in contrast with our results, Starr *et al.* [23] found that the E484K
366 substitution had a small positive effect on ACE2 binding.

367 In the case of the L452R substitution, which is present in the Epsilon variant and also in 368 the Delta variant [64], we observed a modest enhancement of ACE2 binding. Residue L452 is 369 positioned at the edge of the binding interface with ACE2 and although this residue does not 370 make direct contact with ACE2 [34,77], evidence suggests that substitution L452R enhances 371 viral infectivity significantly [77,78]. Furthermore, it has been suggested that the L452R mutation 372 is responsible for the dramatic clonal expansion of lineages carrying this mutation [79], possibly 373 due to a decrease in the potency of antibody neutralization or through other immune escape 374 characteristics [44,46,64,77,78,80]. Whether or not an effect of the L452R substitution on ACE2 375 binding, apparently modest, is a contributing factor in the rapid spread of variants carrying this 376 mutation remains to be determined.

## 377 Summary

378 Taken together, our results illustrate the utility of a B2H system as an accessible and 379 economical genetic tool to complement other methods for studying viral PPIs. To the best of our 380 knowledge, we provide the first bacteria-based viral interactome, describing sixteen different 381 intraviral PPIs from SARS-CoV-2. As a non-eukaryotic system, the B2H assay is unlikely to 382 contain bridging factors that can complicate the interpretation of positive results. At the same 383 time, the bacterial system lacks the machinery for enabling potentially relevant post-translational 384 modifications such as protein phosphorylation (which could however be engineered into the 385 system; [53]) and protein glycosylation. Although generally a limitation, the lack of protein 386 glycosylation could in certain situations be informative, enabling a comparison between systems 387 that do and do not support this modification. The new oxidizing B2H reporter strain that we 388 describe enabled us to detect the SARS-CoV-2 spike RBD-ACE2 interaction and characterize

the effects of several RBD substitutions present in circulating variants. This strain provides a means to test newly arising coronavirus lineages for binding to ACE2 or other human cell surface receptors in the future, as well as extending the reach of the B2H system to include disulfide bond-dependent PPIs in general.

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## 706 Figure Legends

707 Fig. 1: Bacterial two-hybrid assay used to study the SARS-CoV-2 interactome. (A) (top) 708 Schematic depiction of the employed transcription-based bacterial two-hybrid system. 709 Interaction between protein moieties X (purple) and Y (slate blue), which are fused to the N-710 terminal domain of the  $\alpha$  subunit of *E. coli* RNAP ( $\alpha$ NTD) and the  $\lambda$ Cl protein, respectively, 711 stabilizes the binding of RNAP to test promoter placO<sub>L</sub>2–62, thereby activating transcription of 712 the *lacZ* reporter gene. The test promoter bears the  $\lambda$  operator O<sub>1</sub>2 centered at position -62 713 upstream of the transcription start site. (bottom) E. coli cell containing genetic elements that are 714 involved in the bacterial two-hybrid system. The chromosomal *lacZ* locus is deleted and the test 715 promoter and fused *lacZ* reporter gene are encoded on an F' episome. The  $\lambda$ CI-Y and  $\alpha$ NTD-X 716 fusion proteins are encoded on compatible plasmids and produced under the control of IPTG-717 inducible promoters. (B) List of all tested SARS-CoV-2 ORFs as predicted by the NCBI 718 reference genome (Accession #: NC 045512.2). The respective nucleotide range for each ORF 719 based on the NCBI reference sequence is indicated, together with the resulting amino acid 720 sequence length. Except for the spike protein, all ORFs were cloned as full-length genes. For 721 spike, we chose to test the interaction of its ectodomain (aa 16-1213) to avoid complications 722 due to its N-terminal signal peptide and C-terminal transmembrane domain.

723

Fig. 2: Detection of protein-protein interactions by the bacterial two-hybrid system. Interaction matrix of all tested ORFs. Positive interactions, regardless of the fusion partner, are indicated with purple squares and self-interactions are indicated by orange-framed squares. Detailed information about fusion constructs for which positive interactions were identified is given in Supplementary Fig. 1. To avoid data duplication, only one half of the matrix is shown while the other is shaded in grey.

730

731 Fig. 3: Strong SARS-CoV-2 protein-protein interactions identified by B2H assays. Shown 732 are two-hybrid data for strong interactions (arbitrarily defined as Miller units > 500). Indicated 733 ORFs are fused either to the  $\alpha$ NTD (indicated as  $\alpha$ ) or to full-length  $\lambda$ Cl (indicated as Cl).  $\alpha$  and 734  $\lambda$ Cl negative controls express full-length  $\alpha$  and full-length  $\lambda$ Cl, respectively. The interaction of domain 4 of the RNAP  $\sigma^{70}$  subunit (fused to the  $\alpha$ NTD) with the flap domain of the RNAP  $\beta$ 735 736 subunit (fused to  $\lambda$ Cl) served as a positive control (pos) [81,82]. Bar graphs show the averages 737 of three biological replicates (n=3) and  $\beta$ -galactosidase activities are given in Miller units. Error 738 bars indicate the standard deviation. Values indicated with asterisks are significantly different 739 from the negative control. \*\*\*\*: P<0.0001 (One-way ANOVA with Turkey's multiple comparison 740 test).

## 741 Fig. 4: Selective disruption of protein interfaces for protein with two interaction partners.

742 (A) Depiction of crystal structure (PDB ID: 5NFY [29]) of SARS-CoV-1 Nsp10 (pale cyan) in 743 complex with Nsp14 (pale pink). Zoom-in shows amino acids (sticks) chosen for mutational 744 analysis of Nsp10 (orange, olive, and burgundy) and their corresponding main interaction 745 partners in Nsp14 (pale pink). (B) B2H results showing effects of Nsp10 substitutions on its 746 interactions with Nsp14 and with Nsp16. Amino acid substitutions introduced into Nsp10 are 747 given in the box. (C) Depiction of crystal structure of the SARS-CoV-2 Nsp16-Nsp10 protein 748 complex (PDB ID: 6W4H [31]) colored respectively in pale yellow and pale cyan. Additional Nterminal Nsp10<sup>7-22</sup> region is included and was obtained from superimposed Nsp10 structure 749 750 from PDB ID: 5NFY (green). Zoom-in shows amino acids (sticks) chosen for mutational analysis 751 of Nsp16 (orange and burgundy) and their corresponding main interaction partners in Nsp10 752 (pale cyan). (D) B2H results showing effects of Nsp16 substitutions on its interactions with 753 Nsp10 and with Nsp15. Amino acid substitutions introduced into Nsp16 are given in the box. 754 (**B**,**D**) Indicated ORFs are fused either to the  $\alpha$ NTD (indicated as  $\alpha$ ) or to full-length  $\lambda$ Cl.  $\alpha$  and 755  $\lambda$ CI negative controls express full-length  $\alpha$  and full length  $\lambda$ CI, respectively. Bar graphs show 756 the averages of three biological replicates (n=3) and  $\beta$ -galactosidase activities are given in Miller 757 units. Error bars indicate the standard deviation. Values indicated with asterisks are significantly 758 different from the WT. ns: not significant; \*: P<0.05; \*\*: P<0.01; \*\*\*\*: P<0.001 (One-way 759 ANOVA with Dunnett's multiple comparison test). Black dashed lines in A and C represent 760 hydrogen bonds.

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762 Fig. 5: Interaction of spike RBD and ACE2 in an oxidizing E. coli strain. (A) Bacterial two-763 hybrid assays of (A) spike domains (as listed in Supplementary Fig. 7) tested against ACE2 in 764 BLS148, (B) indicated spike RBD cysteine mutants tested against ACE2 in BLS148 and B2H or 765 (C) indicated spike RBD circulating variants tested against ACE2 in BLS148. FL: full-length, 766 NTD: N-terminal domain; RBD: receptor binding domain; CTD: C-terminal domain (with or 767 without transmembrane domain (TMD)), Ecto: Ectodomain starting either at aa 13 or 16. (D) 768 Schematic depicting amino acid substitutions present in each of three RBD variants tested. The 769 measured effect of each substitution on ACE2 binding is indicated with a dash (no effect), a 770 downward pointing arrow (weakened binding) or an upward pointing arrow (strengthened 771 binding). (A-C) spike domains or RBD mutant variants were fused to the  $\alpha$ NTD (indicated as  $\alpha$ ) 772 and ACE2 was fused to full-length  $\lambda$ CI.  $\alpha$  and  $\lambda$ CI negative controls express full-length  $\alpha$  and 773 full-length  $\lambda$ Cl, respectively. Bar graphs show (A) one biological replicate or (B,C) the averages 774 of three biological replicates (n=3) and  $\beta$ -galactosidase activities are given in Miller units. Note: 775 results depicted in (C) have been confirmed in a total of seven independent experiments, one of 776 which is shown here. Error bars indicate the standard deviation. Values indicated with asterisks 777 are significantly different from the negative control. ns: not significant; \*: P<0.05; \*\*: P<0.01; \*\*\*\*: 778 P<0.0001 (Two-way ANOVA with Turkey's multiple comparison test). Western blot analysis 779 indicated that the spike RBD mutants used in (B) and (C) are present at intracellular levels 780 comparable to the wild-type RBD, ruling out protein instability as a cause for the observed 781 effects (Supplementary Figs. 8 and 9).

## 782 Supplementary Figure Legends

Supplementary Fig. 1: Fusion constructs of positive B2H interactions. List of all B2Hidentified SARS-CoV-2 PPIs. Green-shaded rectangles indicate the plasmid from which Protein A (first-mentioned protein in the left column) is produced to elicit a positive interaction with Protein B encoded by the other plasmid. When an interaction of Protein A and Protein B was identified regardless of the encoding plasmid, rectangles are shaded green in the column "both".

788 Supplementary Fig. 2: Medium-to-weak SARS-CoV-2 protein-protein interactions 789 identified B2H assays. Shown are two-hybrid data for medium-to-weak interactions (Miller unit 790 values between 2- and 5-fold above the negative control Miller unit value). Indicated ORFs are 791 fused either to the  $\alpha$ NTD (indicated as  $\alpha$ ) or to full-length  $\lambda$ CI (indicated as CI).  $\alpha$  and  $\lambda$ CI 792 negative controls express full-length  $\alpha$  and full-length  $\lambda$ CI, respectively. The interaction of 793 domain 4 of the RNAP  $\sigma^{70}$  subunit (fused to the  $\alpha$ NTD) with the flap domain of the RNAP  $\beta$ 794 subunit (fused to  $\lambda$ Cl) served as a positive control (pos) [81,82]. Bar graphs show the averages 795 of three biological replicates (n=3) and  $\beta$ -galactosidase activities are given in Miller units. Error 796 bars indicate the standard deviation. Values indicated with asterisks are significantly different 797 from the negative control. \*\*\*: P<0.001; \*\*\*\*: P<0.0001 (One-way ANOVA with Turkey's multiple 798 comparison test).

**Supplementary Fig. 3: Comparison of B2H with Y2H and co-IP data.** Interaction matrix showing all SARS-CoV-2 PPIs identified with the B2H assay (this study) or with the Y2H and co-IP assays [6]. Squares designating PPIs identified solely by the B2H assay are colored purple and contain a white disc, whereas squares designating PPIs identified solely by the Y2H assay or by co-IP experiments are colored red and blue, respectively. Squares designating interactions that were identified by two or three of the assays contain the respective colors as indicated in the key at the left side of the matrix.

Supplementary Fig. 4: Comparison of SARS-CoV-2 B2H data with SARS-CoV-1 two-806 807 hybrid data. Interaction matrix showing the sixteen PPIs identified with the B2H assay (this 808 study) and indicating which of these PPIs were also identified in at least one of three previous 809 SARS-CoV-1 two hybrid studies, including two Y2H analyses [7,27] and one mammalian two-810 hybrid analysis [26]. Squares designating PPIs identified solely by the B2H assay are colored 811 purple and contain a white disc. Coloring within the central disc designates a PPI that was also 812 identified in one or two of the SARS-CoV-1 two-hybrid analyses (von Brunn et al. [7], orange; 813 Imbert et al. [27], yellow; Pan et al. [26], cyan). Note that none of the PPIs were identified in all 814 three SARS-CoV-1 analyses.

## 815 Supplementary Fig. 5: Western blot analysis showing intracellular levels of Nsp10, Nsp14

- and Nsp16 fusion proteins. (top) Anti- $\alpha$ NTD and (bottom) anti- $\lambda$ Cl western blot of cell lysates
- taken from over-night cultures of cells used for B2H analysis shown in **Fig. 4B**. Samples in
- 818 lanes 1-5 are from cells producing the indicated  $\lambda$ CI-Nsp10 fusion protein or  $\lambda$ CI (see key) and
- 819 the  $\alpha$ -Nsp16 fusion protein, whereas samples in lanes 10-14 are from cells producing the
- indicated λCI-Nsp10 fusion protein or λCI and the α-Nsp14 fusion protein. Samples in lanes 6-9
- are from cells producing the indicated  $\lambda$ CI-Nsp10 fusion protein and full-length  $\alpha$ . We note that substitutions D29A+S33A appeared to be mildly destabilizing, whereas substitutions
- 823 F16A+F19A+V21A appeared to be mildly stabilizing.
- 824 Supplementary Fig. 6: Western blot analysis showing intracellular levels of Nsp10, Nsp15
- and Nsp16 fusion proteins. (top) Anti- $\alpha$ NTD and (bottom) anti- $\lambda$ Cl western blot of cell lysates taken from over-night cultures of cells used for the subsequent B2H analysis shown in **Fig. 4D**. Samples in lanes 1-5 are from cells producing the indicated  $\alpha$ -Nsp16 fusion protein or  $\alpha$  (see
- key) and the  $\lambda$ CI-Nsp10 fusion protein, whereas samples in lanes 10-14 are from cells
- producing the indicated α-Nsp16 fusion protein or α and the  $\lambda$ Cl-Nsp15 fusion protein. Samples
- in lanes 6-9 are from cells producing the indicated  $\alpha$ -Nsp16 fusion protein and full-length  $\lambda$ Cl. Lanes indicated with "X" are not discussed in the current manuscript but were not cut from the
- blot to avoid excessive manipulations of the original image. Note: Native, chromosomally
- 833 encoded full-length  $\alpha$  is detected in all samples.
- 834 Supplementary Fig. 7: Domains of the SARS-CoV-2 spike pr
- **Supplementary Fig. 7: Domains of the SARS-CoV-2 spike protein**. Depiction of SARS-CoV-2 spike domains, including the signal peptide (SP, predicted from aa 1-13 or 1-16), the Nterminal domain (NTD, aa 1-330), the receptor binding domain (RBD, aa 331-521), the Cterminal domain (CTD, aa 522-1273) and the transmembrane domain (TMD, aa 1202-1273). The table lists the spike domains that were produced in *E. coli* (as B2H fusion proteins) and tested for interaction with ACE2, their precise corresponding loci on the SARS-CoV-2 genome, and the amino acids encoded by each test domain.
- 841 Supplementary Fig. 8: Western blot analysis showing intracellular levels of spike RBD
- 842 fusion proteins in oxidizing vs. reducing *E. coli* test strains. (top) Anti- $\alpha$ NTD and (bottom) 843 anti-\lambdaCI Western blot of cell lysates taken from over-night cultures of BLS148 or B2H cells used 844 for the subsequent two-hybrid analyses shown in Fig. 5B. Samples in lanes 1-5 and 10-14 are 845 from BLS148 cells and B2H cells, respectively, producing the indicated  $\alpha$ -RBD fusion protein or 846  $\alpha$  (see key) and the  $\lambda$ CI-ACE2 fusion protein, whereas samples in lanes 6-9 and 15-18 are from 847 BLS148 cells and B2H cells, respectively, producing the indicated  $\alpha$ -RBD fusion protein and full-848 length  $\lambda$ Cl. Lanes indicated with "X" are not discussed in the current manuscript but were not 849 cut from the blot to avoid excessive manipulations of the original image. White dotted line 850 indicates stitched image where extraneous material was removed. Note: Native, chromosomally 851 encoded full-length  $\alpha$  is detected in all samples.

## 852 Supplementary Fig. 9: Western blot analysis showing intracellular levels of spike RBD

853 **mutants.** (top) Anti- $\alpha$ -NTD and (bottom) anti- $\lambda$ Cl western blot of cell lysates taken from over-854 night cultures of BLS148 cells used for the subsequent two-hybrid analysis shown in **Fig. 5C**.

- Samples in lanes 1-9 are from cells producing the indicated  $\alpha$ -RBD fusion protein or  $\alpha$  (see key)
- and the  $\lambda$ CI-ACE2 fusion protein, whereas samples in lanes 10-17 are from cells producing the
- 857 indicated  $\alpha$ -RBD fusion protein and full-length  $\lambda$ CI. Note: Native, chromosomally encoded full-
- 858 length  $\alpha$  in all samples.

## 859 Supplementary Table Legends

860 **Supplementary Table 1:** List of (**A**) strains and plasmids, and (**B**) oligonucleotide primers used 861 in this study.

## 862 Acknowledgments

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R35GM136247) awarded to AH.

#### 868 Author contribution

- AH and BLS designed the study. BLS performed the experimental work with support from PD.
- 870 GG analyzed crystal structure data and provided predictions for the mutational screens. BLS
- and AH drafted the manuscript with contributions from all coauthors.

### 872 **Competing interests**

873 The authors declare no competing interests.

### 874 Data availability

- 875 All data generated during and/or analyzed during the current study are either provided within the
- 876 manuscript or are available from the corresponding authors upon reasonable request.

## 877 Material and Methods

## 878 Bacterial strains and growth conditions

879 *E. coli* strains MAX Efficiency<sup>™</sup> DH5αF'IQ (Invitrogen) and NEB® 5-alpha F'IQ (New England 880 Biolabs, NEB) were used for routine cloning procedures and chemically competent E. coli were 881 transformed with plasmid DNA by the standard heat shock procedure. FW102 O<sub>1</sub>2-62 and 882 BLS148 strains were used for bacterial two-hybrid assays. All strains listed in Supplementary 883 Table 1A were grown in LB medium containing the appropriate antibiotics at standard 884 concentrations. BLS148 was created by P1 phage transduction of *AlaclZYA::kmR* from strain 885 TB12 (P1 phage lysates were a gift from Thomas Bernhardt, Harvard Medical School) to 886 SHuffle<sup>®</sup> Express (NEB) according to a protocol established by Robert T. Sauer (Massachusetts 887 Institute of Technology; protocol available at: 888 https://openwetware.org/wiki/Sauer:P1vir\_phage\_transduction), generating BLS128. Deletion of 889 lacZ in BLS128 was verified by colony PCR (using primers oBLS107+oBLS108 targeting lacZ to 890 test for absence of *lacZ* and primers oBLS109+oBLS110 targeting *motA* and primers 891 oBLS138+oBLS139 targeting cyaA as control reactions). Next, TSS competent BLS128 cells 892 were created according to [83], transformed with pCP20 (encoding the yeast Flp recombinase 893 gene to flip out the kanamycin resistance gene) and grown over night at 30 °C on LB plates 894 containing carbenicillin (100 µg/ml; Carb100). The next day, 10 colonies were picked and re-895 streaked on LB plates without antibiotics and then grown over night at 42 °C. From each of 896 those strains a single colony was picked and re-streaked on LB plates containing either 897 Carb100, kanamycin (20 µg/ml; Km20) or spectinomycin (50 µg/ml, Sp50) and grown over night 898 at 30 °C. A single Sp50 resistant but Carb100 and Km20 sensitive colony was picked and re-899 verified by streaking on the same growth plates. This strain that had lost the kmR resistance 900 cassette was then designated BLS133. Finally, the  $\beta$ -galactosidase reporter present on the F' 901 was introduced into BLS133 by mating with strain FW102 O<sub>L</sub>2-62 [84]. For this, both BLS133

and FW102 O<sub>L</sub>2–62 were grown over night at 37 °C in LB Sp50 or Km20, respectively, and then streaked on top of each other on the same LB plate. After about 8 hours at 37 °C cells were resuspended in LB and plated in serial dilutions on LB plates containing Sp50, Km20 and X-gal (40  $\mu$ g/ml; X-gal40) and then grown over night at 37 °C. A blue colony was picked and reverified by streaking again on a LB plate containing Sp50, Km20 and X-gal40, creating BLS148, a bacterial two-hybrid-compatible SHuffle<sup>®</sup> Express strain.

#### 908 Plasmid construction

All plasmids generated in this study (see Supplementary Table 1A) were either constructed using standard restriction enzyme-based cloning procedures or by Gibson assembly. Gibson assembly was performed for 1 h at 50 °C by default. Primers employed for plasmid construction are listed in Supplementary Table 1B. Plasmid sequence integrity was verified by Sanger sequencing from Genewiz or Quintarabio (both Boston, MA, USA). Unless otherwise stated, all sequence templates, except for Nsp11, were ordered as *E. coli* codon-optimized gene fragments from Twist Bioscience (San Francisco, CA, USA).

916 Except for spike, Nsp2, Nsp3, RNA-Polymerase (Nsp12) and helicase (Nsp13), all full-length 917 codon-optimized gene fragments were digested with NotI+BamHI, purified by DNA Clean & 918 Concentrator kit (Zymo Research) and then ligated into 50 ng NotI+BamHI-digested pBR $\alpha$  or 919 pAC $\lambda$ CI using T4 ligase (NEB) according to standard protocols generating the plasmids listed in 920 Supplementary Table 1A.

921 <u>Spike:</u> For pS63, the spike full-length (FL) sequence was amplified from *E. coli* codon-optimized 922 gene fragments by SARS\_67+SARS\_68 and cloned into NotI+BamHI-digested pBR $\alpha$  by Gibson 923 assembly. For pS64, the NTD sequence was amplified from pS63 by SARS\_67+SARS\_69 and 924 then cloned into NotI+BamHI-digested pBR $\alpha$  by Gibson assembly. For pS65, the RBD 925 sequence was amplified from pS63 by SARS\_70+SARS\_71 and then cloned into NotI+BamHI-

926 digested pBR $\alpha$  by Gibson assembly. For pS66, the CTD sequence was amplified from pS63 by 927 SARS 68+SARS 72 and then cloned into Notl+BamHI-digested pBR $\alpha$  by Gibson assembly. 928 For pS67, the Ectodomain (aa 13-1213) was amplified from pS63 by SARS\_73+SARS\_74 and 929 then cloned into NotI+BamHI-digested pBR $\alpha$  by Gibson assembly. For pS68, the Ectodomain 930 (aa 16-1213) was amplified from pS63 by SARS 74+SARS 75 and then cloned into 931 Notl+BamHI-digested pBR $\alpha$  by Gibson assembly. pS70 was generated by site-directed 932 mutagenesis (SDM; see below) using primers SARS 17+SARS 76 and pS63 as a template. 933 For pS72, FL spike was amplified from pS63 by SARS\_77+SARS\_78 and then cloned into 934 NotI+BamHI-digested pAC $\lambda$ CI by Gibson assembly. For pS73, the NTD sequence was amplified 935 from pS63 by SARS 77+SARS 79 and then cloned into NotI+BamHI-digested pACλCI by 936 Gibson assembly. For pS74, the RBD sequence was amplified from pS63 by 937 SARS 80+SARS 81 and then cloned into NotI+BamHI-digested pACλCI by Gibson assembly. 938 For pS75, the CTD sequence was amplified from pS63 by SARS 78+SARS-82 and then cloned 939 into NotI+BamHI-digested pACλCI by Gibson assembly. For pS76, the Ectodomain (aa 13-940 1213) was amplified from pS63 by SARS\_83+SARS\_84 and then cloned into NotI+BamHI-941 digested pAC\lambda Cl by Gibson assembly. For pS77, the Ectodomain (aa 16-1213) was amplified 942 from pS63 by SARS 84+SARS 85 and then cloned into NotI+BamHI-digested pACλCI by 943 Gibson assembly. pS79 was generated by SDM using primers SARS 17+SARS 86 and pS63 944 as a template.

945 <u>Nsp2</u>: The Nsp2 sequence was ordered as two single gene fragments, which were further 946 amplified by PCR using primers SARS\_109+SARS\_110 or SARS\_111+SARS\_112 and then 947 cloned into NotI+BamHI-digested pBR $\omega$ GP by Gibson assembly, creating pS85. Next, the whole 948 Nsp2 open reading frame (ORF) was cut from pS85 by NotI+BamHI and inserted into 50 ng

949 NotI+BamHI-digested pBRα or pACλCI using T4 ligase (NEB) according to standard protocols,
950 creating pS179 and pS180, respectively.

951 Nsp3: The Nsp3 sequence was ordered as four single gene fragments, which were further 952 amplified PCR using primers SARS 115+SARS 116, SARS 117+SARS 118, by 953 SARS\_119+SARS\_120 or SARS\_121+SARS\_122 and then cloned into NotI+BamHI-digested 954 pBR@GP by Gibson assembly, creating pS89. Next, the whole Nsp3 open reading frame (ORF) 955 was cut from pS89 by NotI+BamHI and inserted into 50 ng NotI+BamHI-digested pBR $\alpha$  or 956 pAC<sub>\lambda</sub>Cl using T4 ligase (NEB) according to standard protocols, creating pS181 and pS182. 957 respectively.

958 <u>RNA-Polymerase (Nsp12)</u>: The RNA-Polymerase sequence was ordered as three single gene 959 fragments, which were further amplified by PCR using primers SARS\_131+SARS\_132, 960 SARS\_133+SARS\_134 or SARS\_135+SARS\_136 and then cloned into NotI+BamHI-digested 961 pBR $\omega$ GP by Gibson assembly, creating pS173. Next, the whole Nsp3 open reading frame 962 (ORF) was cut from pS89 by NotI+BamHI and inserted into 50 ng NotI+BamHI-digested pBR $\alpha$ 963 or pAC $\lambda$ CI using T4 ligase (NEB) according to standard protocols, creating pS181 and pS182, 964 respectively.

helicase (Nsp13): The helicase sequence was ordered as two single gene fragments, which were further amplified by PCR using primers SARS\_125+SARS\_126 or SARS\_127+SARS\_128 and then cloned into NotI+BamHI-digested pBR $\omega$ GP by Gibson assembly, creating pS169. Next, the whole Nsp2 open reading frame (ORF) was cut from pS85 by NotI+BamHI and inserted into 50 ng NotI+BamHI-digested pBR $\alpha$  or pAC $\lambda$ CI using T4 ligase (NEB) according to standard protocols, creating pS221 and pS222, respectively.

971 <u>ACE2</u>: The ACE2 N-terminal peptidase domain (aa 19-615) was ordered as a single gene 972 fragment and then further amplified by PCR using primers SARS\_264+SARS\_265 or 973 SARS\_266+SARS\_267 and then cloned into 50 ng NotI+BamHI-digested pBR $\alpha$  or pAC $\lambda$ CI by 974 Gibson assembly, creating pS260 and pS261, respectively.

975 <u>Nsp11</u> was cloned into pBR $\alpha$  or pAC $\lambda$ Cl as annealed primers. For this, 10 µl of 100 µM 976 SARS\_139 and SARS\_140 primers were mixed with 1 µl T4 Polynucleotide kinase (PNK; NEB) 977 in 1x PNK reaction buffer (NEB). The reaction mix was placed in a BioRad T100 Thermal cycler, 978 incubated for 30 min at 37 °C, inactivated for 5 min at 95 °C and then cooled to 4 °C at a 0.1 979 °C/s ramp rate. The annealed oligos were diluted 1:50 and then ligated into 50 ng NotI+BamHI-980 digested pBR $\alpha$  or pAC $\lambda$ Cl using T4 ligase (NEB), generating pS197 and pS198, respectively.

### 981 Plasmid mutagenesis

Plasmid mutagenesis to create SARS-CoV-2 mutant genes was achieved using the Q5<sup>®</sup> Site-Directed Mutagenesis (SDM) Kit according to the manufacturer's instructions (NEB) or by using Gibson assembly with mutations introduced into the complementary overhang regions of the primer sequences. For the Gibson assembly, plasmids were amplified with the indicated primer pairs and 1  $\mu$ l of the resulting PCR reaction was then ligated by Gibson assembly in a 10  $\mu$ l reaction volume.

988 pS254 was generated by SDM using primers SARS\_253+SARS\_254 and plasmid pS215 as a 989 template. pS256 was generated by SDM using primers SARS\_256+SARS\_257 and plasmid 990 pS215 as a template. pS257 was generated by SDM using primers SARS\_256+SARS\_257 and 991 plasmid pS254 as a template. pS262 was generated by Gibson assembly using primers 992 SARS\_268+SARS\_269 and plasmid pS196 as a template. pS263 was generated by Gibson 993 assembly using primers SARS\_270+SARS\_271 and plasmid pS196 as a template. pS264 was 994 generated by Gibson assembly using primers SARS\_272+SARS\_273 and plasmid pS196 as a

995 template. pS267 was generated by Gibson assembly using primers SARS 280+SARS 281 and 996 plasmid pS65 as a template. pS271 was generated by Gibson assembly using primers 997 SARS 287+SARS 288 and plasmid pS65 as a template. pS272 was generated by Gibson 998 assembly using primers SARS 289+SARS 290 and plasmid pS65 as a template. pS273 was 999 generated by Gibson assembly using primers SARS 289+SARS 290 and plasmid pS271 as a 1000 template. pS275 was generated by Gibson assembly using primers SARS 295+SARS 296 and 1001 plasmid pS65 as a template. pS276 was generated by Gibson assembly using primers 1002 SARS 291+SARS 292 and plasmid pS65 as a template. pS277 was generated by Gibson 1003 assembly using primers SARS\_293+SARS\_294 and plasmid pS65 as a template. pS278 was 1004 generated by Gibson assembly using primers SARS 291+SARS 292 and plasmid pS267 as a 1005 template. pS279 was generated by Gibson assembly using primers SARS 293+SARS 294 and 1006 plasmid pS267 as a template. pS280 was generated by Gibson assembly using primers 1007 SARS 293+SARS 294 and plasmid pS278 as a template.

## 1008 $\beta$ -galactosidase assays

1009  $\beta$ -galactosidase assays to study the SARS-CoV-2 interactome were performed essentially as 1010 described previously [85]. In particular, pBR $\alpha$  and pAC $\lambda$ CI plasmids containing the indicated 1011 inserts were co-transformed into FW102 O<sub>1</sub>2-62 by the heat shock procedure. Briefly, 2 µl of 1012 each plasmid (1:10 dil.) were mixed with 20 µl chemically competent FW102 OL2-62 cells, 1013 incubated on ice in 96-well PCR plates (VWR) for 30 min and then heat-shocked for 1 min at 42 1014 °C in a BioRad T100 Thermal cycler. Cells were placed on ice for 5 min, recovered in 80 µl 1015 fresh LB medium and then incubated at 37 °C for 1 h (Please note: we found that commercially 1016 available premixed LB drastically reduces transformation efficiency and also subsequent over 1017 night culture growth; we thus recommend using non-premixed LB medium instead). The 96-well 1018 plates were sealed with Rayon Films (VWR) to allow proper aeration and prevent 1019 contamination. Afterwards, 50 µl of transformed cells were transferred to 2 ml deep well plates

1020 containing 500 µl LB Carb100, chloramphenicol (25 µg/ml; Cm25), Km20 and 5 µM IPTG and 1021 grown over night at 37 °C, 800 rpm. The next day, 4 µl over-night culture was transferred to 96-1022 well flat bottom microtiter plates containing 200 µl LB Carb100, Cm25, Km20 and 20 µM IPTG 1023 and grown until approx.  $OD_{600}$  0.15-0.2 (measured in a VERSA Max microplate reader, 1024 Molecule Devices, San Jose, CA, USA). Then, 20 µl lysis solution (for one 96-well plate mix: 1.2 ml PopCulture<sup>®</sup> Reagent (MilliporeSigma, MA, USA), 2.5 µl 400 U/µl rLysozyme<sup>™</sup> 1025 (MilliporeSigma, MA, USA) and 1.25 µl Benzonase<sup>®</sup> Nuclease (MilliporeSigma, MA, USA)) was 1026 added to the cells and incubated for at least 30 min at 37 °C and 800 rpm (longer incubation 1027 1028 times were found to not negatively affect the experimental results). Afterwards, 30 µl lysed cell 1029 suspension was added to a fresh 96-well flat-bottom microtiter plate containing 150 µl Z-1030 buffer/ONPG solution (60 mM Na<sub>2</sub>HPO<sub>4</sub>, 40 mM NaH<sub>2</sub>PO<sub>4</sub>, 10 mM KCl, 1 mM MgSO<sub>4</sub>, 1 mg/ml 1031 ortho-Nitrophenyl-β-galactoside (ONPG)) and OD<sub>420</sub> values were recorded in a VERSA Max 1032 microplate reader (Molecule Devices, San Jose, CA, USA). β-galactosidase activity in Miller 1033 units was then calculated as described previously [85].

1034 β-galactosidase assays to study the RBD-ACE2 interaction were performed as follows. Strain 1035 BLS148 was transformed with the appropriate plasmids, as described in the preceding 1036 paragraph. Upon recovery of the transformed cells for 1 h at 37 °C, 50 µl cells were then 1037 transferred to 500 µl LB Carb100, Km20, Cm25 and 50 µM IPTG and grown for approx. 20 h at 1038 30 °C and 800 rpm. Subsequently, 15 µl cells were transferred to 185 µl LB medium in 96-well 1039 microtiter plates, combined with 20 µl lysis solution (for one 96-well plate: 1.2 ml PopCulture® Reagent (MilliporeSigma, MA, USA), 5.0 µl 400U/µl rLysozyme<sup>™</sup> (MilliporeSigma, MA, USA) 1040 and 2.5 µI Benzonase<sup>®</sup> Nuclease (MilliporeSigma, MA, USA)) and incubated for at least 30 min 1041 1042 at 30 °C, 800 rpm. All subsequent steps were then performed as described above.

1043

## 1044 Western blot analysis

1045 To verify the production of the respective fusion proteins, western blots of cell lysates from over-1046 night cultures were performed. For this, co-transformed cells were grown in the indicated IPTG 1047 concentration over-night in 550 µl total volume in 2 ml deep well plates at 30 or 37 °C, 800 rpm. 1048 The next day,  $OD_{600}$  values were recorded and 500 µl cells were pelleted by centrifugation (1 1049 min, 21,000 x g, room temperature (RT)) and either stored at -80 °C or directly processed. Cell 1050 pellets were then resuspended in lysis buffer (BugBuster® Protein Extraction Reagent 1051 (MilliporeSigma, MA, USA) supplemented with 1x cOmplete™, EDTA-free Protease Inhibitor Cocktail (MilliporeSigma, MA, USA), 1 U/µl rLysozyme<sup>™</sup> (MilliporeSigma, MA, USA; final 1052 1053 concentration) and 0.5 U/µI Benzonase<sup>®</sup> Nuclease (MilliporeSigma, MA, USA, final 1054 concentration)). The amount of lysis buffer for each cell pellet was calculated as follows: µI lysis 1055 buffer =  $OD_{600}$  x ml of culture pelleted x 60. Cells were lysed for 30 min at RT in an overhead 1056 shaker. Next, lysed cells were mixed 1:5 in PBS (10.14 mM Na<sub>2</sub>HPO<sub>4</sub>, 1.76 mM NaH<sub>2</sub>PO<sub>4</sub>, 2.7 1057 mM KCl, 137 mM NaCl; pH 7.4, Boston Bioproducts, MA, USA) and then incubated in 1 x 1058 Laemmli SDS sample buffer (Boston Bioproducts, MA, USA) at 95 °C for 10 min. 10 µl of the 1059 resulting solution was then applied to either 4–12% Criterion™ XT Bis-Tris Protein Gels 1060 (BioRad, Hercules, CA, USA) or NuPAGE<sup>™</sup> 4 - 12% Bis-Tris Mini Protein Gels (Thermo Fisher 1061 Scientific, MA, USA). Upon gel separation, proteins were transferred to Amersham Protran 0.45 1062 NC nitrocellulose membranes (Cytiva, MA, USA) using a Trans-Blot Turbo Transfer System 1063 (BioRad Hercules, CA, USA), blocked in blocking buffer (TBST: 50 mM Tris-HCl, 150 mM NaCl, 1064 pH 7.4, 0.1 % Tween-20 supplemented with 5% non-fat dry milk) for 30 min at RT and then 1065 incubated with mouse anti-a-NTD and rabbit anti-CI primary antibodies (both 1:3.000 dil.) in 1066 blocking buffer for 1 h at RT. After washing with TBST, blots were incubated with IRDye® 1067 680RD goat anti-mouse and IRDye® 800CW goat anti-rabbit (both 1:10,000 dil.; LI-COR Biosciences, NE, USA) in blocking buffer for 1 h at RT in the dark. After washing with TBST,

- 1069 proteins were then detected using a ChemiDoc MP system (BioRad. Hercules, CA, USA).
- 1070 Protein crystal structure analysis

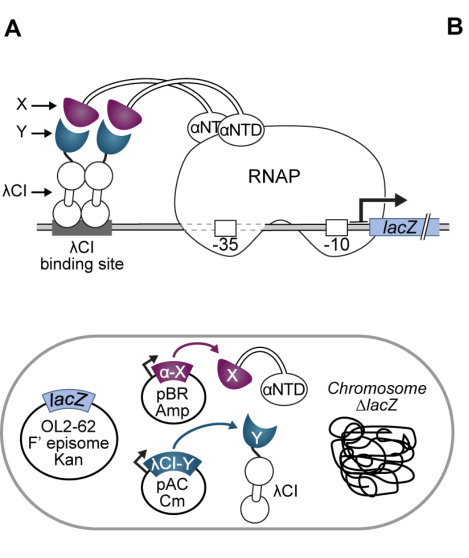
1071 Interfaces of two protein complexes, SARS-CoV-2 Nsp16-Nsp10 (PDB ID: 6W4H [31]) and
1072 SARS-CoV-1 Nsp10-Nsp14 (PDB ID: 5NFY [29]), were analyzed using PDBePISA software
1073 (*insert ref 20*). Amino acids involved in hydrogen bond formation or substantially contributing to
1074 hydrophobic contacts in each complex were subjected to alanine mutagenesis and tested in
1075 B2H assays. Structural images were prepared using PyMOL software (Schrodinger, LLC. 2010.
1076 The PyMOL Molecular Graphics System, Version 2.4.1).

1077 Statistical analysis

1078 Presentation of bacterial two-hybrid data and statistical analysis using one-way or two-way

1079 ANOVA with Tukey's or Dunnett's multiple comparison test was done using GraphPad Prism (v.

1080 9.1.2; San Diego, CA, USA).

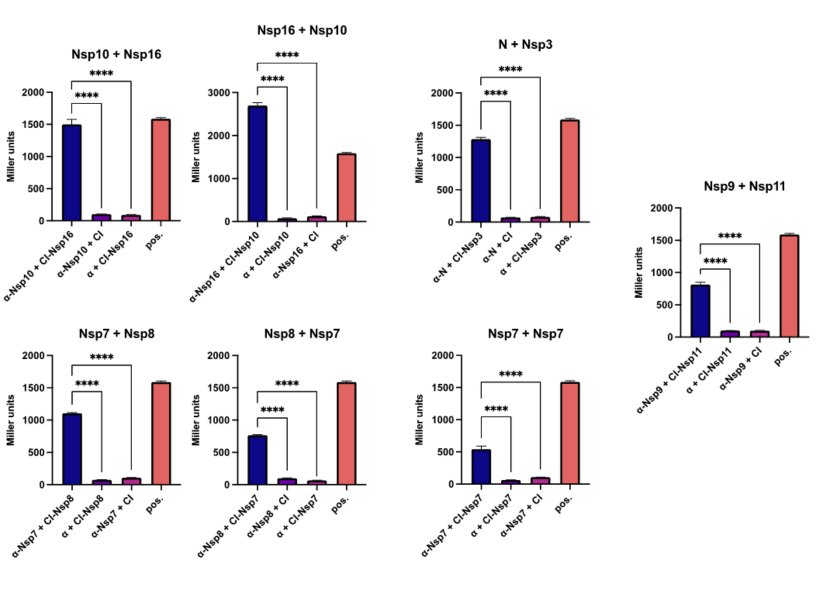


ORF	Genomic locus (nt range)	AA
Nsp1	266-805	188
Nsp2	806-2719	636
Nsp3	2720-8554	1,945
Nsp4	8555-10054	500
Nsp5	10055-10972	306
Nsp6	10973-11842	290
Nsp7	11843-12091	83
Nsp8	12092-12685	198
Nsp9	12686-13024	113
Nsp10	13025-13441	139
Nsp11	13442-13483	13
Nsp12 (RNA-Polymerase)	13442-16236	932
Nsp13 (Helicase)	16237-18039	601
Nsp14 (3'-5' Exonuclease)	18040-19620	527
Nsp15 (endoRNase)	19621-20658	346
Nsp16 (Methyltransferase)	20659-21555	298
Spike (S) ectodomain	21607-25201	1,201
3a	25393-26220	275
E	26245-26472	75
M	26525-27191	222
6	27202-27387	61
7a	27397-27759	121
7b	27756-27887	43
8	27894-28259	121
N	28274-29533	419
10	29558-29674	38

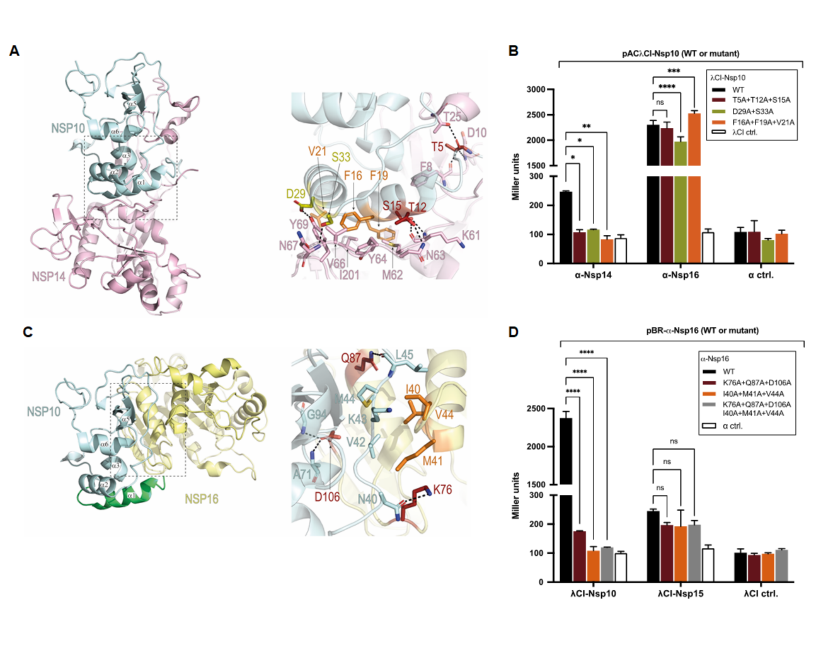
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	Nsp1	Nsp2	Nsp3	Nsp4	Nsp5	Nsp6	Nsp7	Ns p8	Nsp9	Nsp10	Nsp11	Nsp12	Nsp13	Nsp14	Nsp15	Nsp16	3a	6	7a	7b	80	10	s	W	z	ш
Nsp1																										
Nsp2																										
Nsp3																										
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