

18 Abstract

19 We integrate evolutionary predictions based on the neutral theory of molecular evolution with protein 20 dynamics to generate mechanistic insight into the molecular adaptations of the SARS-COV-2 Spike (S) 21 protein. With this approach, we first identified Candidate Adaptive Polymorphisms (CAPs) of the SARS-22 CoV-2 Spike protein and assessed the impact of these CAPs through dynamics analysis. Not only have we 23 found that CAPs frequently overlap with well-known functional sites, but also, using several different 24 dynamics-based metrics, we reveal the critical allosteric interplay between SARS-CoV-2 CAPs and the S 25 protein binding sites with the human ACE2 (hACE2) protein. CAPs interact far differently with the hACE2 26 binding site residues in the open conformation of S protein compared to the closed form. In particular, 27 the CAP sites control the dynamics binding residues in the open state, suggesting an allosteric control of 28 hACE2 binding. We also explored the characteristic mutations of different SARS-CoV-2 strains to find 29 dynamic hallmarks and potential effects of future mutations. Our analyses reveal that Delta strain-specific 30 variants have non-additive (i.e., epistatic) interactions with CAP sites, whereas the less pathogenic 31 Omicron strains have mostly compensatory variants. Finally, our dynamics-based analysis suggests that 32 the novel mutations observed in the Omicron strain epistatically interact with the CAP sites to help escape 33 antibody binding.

34 Introduction

35 Since 2019, the evolution of SARS-CoV-2 in humans has been characterized by the spread of mutations, 36 many notably found within the Spike (S) glycoprotein. The S protein is directly related to the human 37 immune response to COVID-19 and, as such, has been one of the most studied and targeted proteins in 38 the SARS-CoV-2 research (Shang et al. 2020; Harvey, Carabelli, Jackson, Gupta, Thomson, Harrison, 39 Ludden, Reeve, Rambaut, Consortium, et al. 2021; Jackson et al. 2022; Carabelli et al. 2023; Markov et al. 40 2023). Subsequently, research into the biophysical properties and mutational patterns associated with S

41 protein evolution not only remains critical to understanding the pandemic but also emerges as a useful 42 system to understand the mechanics of molecular adaptation within viruses.

43 For successful infection of a human host, the S glycoprotein of SARS-CoV-2 binds to the human ACE2 44 (hACE2) receptor through its receptor-binding domain (RBD). Evidence indicates that fine-tuning S protein 45 interactions with hACE2 significantly affects viral reproduction (Rehman et al. 2020; Saputri et al. 2020; 46 Rochman et al. 2021). Previous evolutionary studies show a complex network of interactions among 47 mutated residues (Changeux and Edelstein 2005; Doshi et al. 2016; O'Rourke et al. 2016; Mishra and 48 Jernigan 2018). Therefore, there has been a vast effort to uncover which mutations are important steps 49 of adaptation for the S protein (Cagliani et al. 2020; Damas et al. 2020; Singh and Yi 2021; Kistler et al. 50 2022; Maher et al. 2022; Neher 2022). In particular, a significant aspect of many such studies was a focus 51 on understanding adaptive mutations of SARS-CoV-2 that contributed to the leap to human hosts (Cagliani 52 et al. 2020; Damas et al. 2020; Singh and Yi 2021; Starr, Zepeda, et al. 2022). This is because SARS-CoV-2 53 has continuously mutated since its early detection (Kistler et al. 2022), causing the emergence of CDC-54 designated "variants of concern" (VOCs) that an accelerated substitution rate may drive (Tay et al. 2022).

55 Predicting how new mutations impact the biophysical properties of the S protein remains a challenge, let 56 alone explaining their complex interactions with one-another and how they might affect hACE2 binding 57 because many factors affect hACE2 interactions. Binding affinity with hACE2 can be enhanced directly 58 through stronger receptor interactions or mediated through changes in RBD opening (Teruel et al. 2021; 59 Zhang et al. 2021; Díaz-Salinas et al. 2022). The RBD exhibits both 'closed' and 'open' conformational 60 states. In the closed state, the RBD is shielded from receptor binding. In the open state, the RBD is 61 accessible for hACE2 binding (Kirchdoerfer et al. 2016; Gur et al. 2020; Henderson et al. 2020; Hoffmann 62 et al. 2020). While some mutations may affect the transition between these states (Henderson et al. 2020; 63 Yurkovetskiy et al. 2020; Gobeil, Janowska, McDowell, Mansouri, Parks, Manne, et al. 2021; Sztain et al. 64 2021; Zhang et al. 2021; Shoemark et al. 2022), the other mutations may allosterically regulate RBD 65 openings through Furin cleavage site (residue ID range: 681-695) interactions to regulate hACE2 binding 66 (Deng et al. 2021; Gobeil, Janowska, McDowell, Mansouri, Parks, Stalls, et al. 2021; Khan et al. 2021; 67 Laiton-Donato et al. 2021).

68 Moreover, as new mutations accumulate, culminating in the emergence of a new VOC, these mutations 69 must occur on varied sequence backgrounds containing neutral, nearly-neutral, and adaptive mutations. 70 While many studies have explored the impacts of individual mutations, VOCs result in a substantial 71 difference in protein function compared to their individual effects (Moulana et al. 2022a; Starr, Greaney, 72 Hannon, et al. 2022; Moulana et al. 2023; Witte et al. 2023). Here we integrate an evolutionary approach 73 with protein dynamics analysis to address the fundamental mechanisms of mutations dictating the VOCs 74 and the impact of their epistatic interaction on the function of the S protein. Many earlier studies have 75 combined phylogeny and evolutionary theory to identify adaptive mutations as well as to see how the 76 viral sequence has changed over time (Frost et al. 2018; Boni et al. 2020; Cagliani et al. 2020; Damas et al. 77 2020; Tang et al. 2020). Similarly, we first use a well-established Evolutionary Probability (EP) approach 78 (Liu et al. 2016) that utilizes phylogenetic trees in combination with the neutral theory of molecular 79 evolution to determine Candidate Adaptive Polymorphisms (CAPs) determined using the early Wuhan 80 sequence as a variant. CAPs are substitutions in SARS-CoV-2 that are rarely observed in other closely 81 related sequences (Figure 1A), which implies a degree of functional importance and makes them 82 candidates for adaptation (Liu et al. 2016). In support of this method, we find an overlap between our list 83 of sites containing CAPs and putative adaptive sites identified by others (Cagliani et al. 2020; Singh and Yi 84 2021; Starr, Zepeda, et al. 2022). Second, we use a suite of computational tools to analyze how CAPs that 85 arose in the early and late phases of the COVID-19 pandemic modulate the dynamics of the S protein. We 86 also explore the complex interactions between these sets of CAPs to gain mechanistic insight into the 87 behavior of molecular adaptation involving the S protein. In particular, we focused on how mutations 88 modulate protein dynamics, as we and others have previously found that rather than changing a protein's

89 structure solely, mutations modulate conformational dynamics leading to changes in biophysical 90 properties such as stability, flexibility, and allosteric dynamic coupling, any of which may affect protein 91 binding (Swint-Kruse et al. 1998; Keskin et al. 2000; Bhabha et al. 2013; Nussinov, R., Tsai, C.-J 2013; 92 Campbell, E. et al. 2016; Ma and Nussinov 2016: 201; Saavedra et al. 2018; Kuzmanic et al. 2020).

93 With this evolutionary-dynamics unified approach, we aim to answer the following questions about VOCs: 94 Are all the mutations in VOCs adaptive in nature? Are they coupled to one another or provide some 95 measure of biophysical, dynamical, or mechanical compensation? While many of these mutations are 96 found within the RBD domain, numerous others are located distal to this region; hence, we aim to 97 investigate the functional roles of distal mutations, particularly from a protein dynamics perspective. Our 98 integrated analysis revealed that protein dynamics play a significant role in the evolution of the S protein. 99 The flexibility of sites withing the S protein shows a strong, direct correlation with substitution rate, and 100 newly evolving CAPS are mostly compensatory (i.e., additive) mutations that modulate the dynamics of 101 the hACE2 binding site. Yet other CAPs, 346R, 486F, and 498Q, show highly epistatic (i.e., non-additive) 102 modulation of the hACE2 binding site and provide immune escape benefits.

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104 Results and Discussions

105 Candidate Adaptive mutations in the Spike protein are recognized via Evolutionary Probabilities.

106 SARS-CoV-2 is part of a family of coronaviruses, many of which infect mainly animals and are less capable 107 of infecting humans (Dicken et al. 2021). Therefore, to identify the most likely mutations responsible for 108 the infection of human hosts (i.e., putative adaptive mutations for humans), we estimated the (neutral) 109 evolutionary probability (EP) scores of mutations found within the S protein (Liu et al. 2016). EP scores of 110 the amino acid variants of S protein are obtained by constructing a maximum likelihood phylogenetic tree 111 (Goldman 1990) containing 19 orthologous coronavirus sequences, which were selected based on the

112 amount of divergence over evolutionary history to ensure that each amino acid position had ample time 113 to experience purifying selection (Patel et al. 2018) (Figure 1A). The likelihood of finding a particular amino 114 acid in the sequence is then determined using a Bayesian framework, with calculations carried out by 115 MEGA X software (Kumar et al. 2018). As apparent in the name, EP scores obtained for the amino acids in 116 the sequence provide information regarding the likelihood of finding them at their position, given the 117 history of the sequence. Amino acid residues receiving low EP scores (<0.05) at a position are less likely to 118 be found in a given position within the sequence because they are non-neutral. Generally, positions with 119 low EP amino acids are far less common than those containing mutations with high EP, a trend also 120 realized in the CoV-2 S protein (Figure 1B).

121 Of particular interest is an observed evolutionary change where an amino acid with high EP is replaced by 122 an amino acid residue with low EP. While amino acids with low EP should be harmful or deleterious to 123 viral fitness due to functional disruption or change, fixation of a low EP amino acid at a position suggests 124 an underlying mechanism for natural selection to operate. These fixed, low EP mutations are called 125 candidate adaptive polymorphisms (CAPs) as they are predicted to alter protein function, and adaptive 126 pressures may drive their prevalence (Patel et al. 2018). Indeed, there is an overlap between these CAPs 127 and the mutations suggested by other methods to be adaptive for the S protein (Cagliani et al. 2020; Singh 128 and Yi 2021; Starr, Zepeda, et al. 2022).

130 Figure 1. (A) The evolutionary probabilities (EP) of each amino acid in the S protein sequence are 131 calculated by taking the multiple sequence alignment of the S proteins through their evolutionary tree 132 and using Bayesian inferences to determine the likelihood of finding a particular residue at a particular 133 location within a given sequence. Simply, if the residue is found at a location 'x' in closely related 134 sequences, it will have a higher EP at location 'x' in the target sequence. Residues with an EP <0.05 in the 135 target sequence are CAPs (Red). (B) The distribution of EP scores of the wild-type residues in the S protein. 136 Here, lower EP scores are shown in red, and higher EP scores in blue. While the vast majority of the wild-137 type (reference) protein consists of high EP residues, a few residues have low EP. (C) The CAPs are also 138 highlighted as red spheres in the open configuration of the S protein, with the open chain in a darker 139 shade. We observe that a majority of the CAP positions reside at the receptor binding domain (RBD) and 140 the Furin cleavage site (676-689; (Wrobel et al. 2020)) shown as transparent light gray spheres.

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141 Interestingly, most of the CAP residues are in functionally critical sites, including the receptor binding 142 domain (RBD) and the Furin cleavage site (Figure 1C). As mentioned earlier, the RBD plays a key role in 143 initiating the infection of a healthy cell by binding it with the host organism's ACE2 protein. Before ACE2

144 binding, one chain of the homotrimer comprising the S protein must open to expose the RBD (Kirchdoerfer 145 et al. 2016; Henderson et al. 2020; Hoffmann et al. 2020; Sztain et al. 2021). The Furin cleavage site plays 146 a key role in the opening process as it aids in the cleavage of the S protein into two domains: S1 and S2 147 (Wrobel et al. 2020: 13). Similar cleavage sites have been found in related coronaviruses, including HKU1 148 and Middle East respiratory syndrome coronavirus (MERS-CoV), which infect humans (Chan et al. 2008: 149 1; Millet and Whittaker 2014; Millet and Whittaker 2015), and the acquisition of similar cleavage sites is 150 associated with increased pathogenicity in other viruses such as the influenza virus (Steinhauer 1999). 151 Interestingly, however, CAPs do not display such an overwhelming tendency to occur at well-known 152 critical sites within human proteins studied with similar methods (Ose, Campitelli, et al. 2022), yet 153 mutations at those sites are associated with disease, indicating their critical role in inducing functional 154 change. Therefore, the identified CAPs in the S protein, which are signs of recent evolution, can provide 155 mechanistic insights regarding the molecular adaptation of the virus. In particular, we aimed to analyze 156 how these CAP positions in the S protein modulate the interaction with hACE2 using our protein dynamics-157 based analysis (Gerek and Ozkan 2011; Nevin Gerek, Z., Kumar, S., Banu Ozkan, S. 2013; Larrimore et al. 158 2017; Kumar, A., Glembo, T.J., Ozkan, S.B. 2015b).

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160 Asymmetry in communications among the network of interactions in Spike describes how CAPs regulate 161 the dynamics of the Spike protein.

162 A mutation at a given amino acid position inevitably not only alters local interactions, but this change 163 cascades through the residue-residue interaction network, which gives rise to a variation in native 164 ensemble dynamics to modulate function (Dror et al. 2012; Labbadia and Morimoto 2015; Sekhar and Kay 165 2019; Campitelli et al.). Thus, we analyze the internal dynamics of the system to understand the functional 166 role of CAPs in S proteins. This analysis allows us to gain a mechanistic understanding of the relationship

167 between CAP mutations and biophysical outcomes (Teruel et al. 2021). First, we implement the Dynamic 168 Coupling Index (DCI) approach to study long-distance coupling between the CAPs and the hACE2 binding 169 sites emerging from the 3D network of interactions across the S protein system. The DCI parameter 170 combines Perturbation Response Scanning and Linear Response Theory to capture the strength of a 171 displacement response for position *i* upon perturbation of position *j*, relative to the average fluctuation 172 response of position *i* to all other positions in the protein. It represents the strength of dynamic coupling 173 between positions i and j upon perturbation to j .

174 Further, asymmetry can be captured in the DCI values, as dynamic coupling is not necessarily due to an 175 anisotropic network. That is, each amino acid has a set of positions to which it is highly coupled, and this 176 anisotropy in connections gives rise to unique differences in coupling between a given *i*, *j* pair of amino 177 acids which do not have direct interactions. By calculating the coupling of the hACE2 binding interface in 178 the RBD with respect to the CAP residue positions and vice versa, we can generate DCI_{asym} (Figure 2A) as 179 the difference between the normalized displacement response of position *j* upon a perturbation to 180 position i (DCI_{ii}) and the normalized displacement response of position i upon a perturbation to position j 181 (DCI_{ii}) (See Methods). If the DCI_{asym} values significantly differ from zero, it shows asymmetry in coupling 182 and presents a cause-effect relationship between the i , j pair in terms of force/signal propagation. This 183 metric has been used previously in a variety of systems to analyze the unique behavior of positions within 184 a protein and a given position's propensity to effect biophysical changes upon mutation, particularly at 185 long distances (Modi and Ozkan 2018; Campitelli and Ozkan 2020; Kolbaba-Kartchner et al. 2021; Ose, 186 Butler, et al. 2022; Kazan et al. 2023; Campitelli et al. 2020b).

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188 Figure 2. (A) Schematic representation of DCI asymmetry. (B) DCI asymmetry of CAP residue positions 189 with the binding interface of RBD in the open chain. Residues in the closed chains with a low EP amino 190 acid in the reference sequence dominate the binding site interface of RBD in the open chain. There is a 191 significant difference between the asymmetry profiles of the closed (M = -0.06, SD = 0.33) and open (M = 192 -1.68 , SD = 0.89) conformations (p < .001).

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194 Recent work from our group has shown an enhancement in cross-chain communication within the main 195 protease of SARS COV-2 compared to SARS COV-1 (Campitelli et al. 2022). Furthermore, previous studies 196 have shown that allosteric inter-chain communication is important to S protein function (Zhou et al. 2020; 197 Spinello et al. 2021; Tan et al. 2022; Xue et al. 2022). In support of these findings, we observe through 198 DCI_{asym} that when the S protein is in its pre-fusion conformation with one chain open, the CAPs in the 199 closed chains have negative coupling asymmetry with respect to the hACE2 binding site interface in the 200 RBD-open chain. This indicates an allosteric control where the hACE2 binding site is dominated by the 201 dynamics of the CAPs in closed chains (Figure 2B, yellow bars). As this open-state RBD is critical for the 202 viral infection of host cells (Kirchdoerfer et al. 2016), our results suggest that this type of closed-to-open 203 cross-chain interaction is important for viral proliferation. Our prior studies on DCI_{asym} show a similar trend 204 in Lactose Inhibitor (LacI), a protein with a functional role in gene expression through binding DNA. The 205 allosteric mutations (i.e., mutations on the sites that are far from the DNA binding sites) that alter DNA

206 binding affinity not only exhibited unique asymmetry profiles with the DNA binding sites of LacI, but also 207 regulated the dynamics of these binding sites (Campitelli et al. 2020b).

208 Similarly, it is possible that mutations to such residue positions within the S protein can be used to regulate 209 the dynamics of the hACE2 bindings sites of the open RBD state. We, therefore, propose that the residue 210 positions with CAP substitutions hold the potential for mutations in the spike sequence which can alter 211 the opening and closing dynamics of the RBD domain. This hypothesis is further supported by many 212 mutations already observed at these residue positions which alter the infection rate (Brister et al. 2015). 213 Interestingly, residues responsible for extremely low asymmetry values (< -4) lie overwhelmingly in the 214 region 476–486. These same residues were suggested to stabilize S protein dynamics and prime it for host 215 Furin proteolysis (Raghuvamsi et al. 2021).

216 Moreover, as a control, we performed the same analysis on the S protein with the RBD domains of all 217 chains in the closed configuration. In this case, we observed that the DCI_{asym} of the CAPs residue positions 218 with respect to the hACE2 interface in the other chains yields a largely symmetric distribution about 0 219 (Figure 2B, green bars). This verifies that the asymmetry in the coupling of CAPs with the exposed binding 220 site interface in pre-fusion configuration results from one of the RBDs opening up and further suggests 221 the allosteric role played by CAPs in locking the S protein in the RBD open state.

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223 Dynamic analysis shows that rigid sites tend to be more highly conserved than flexible sites.

224 CAPs represent important S protein amino acid changes between related coronaviruses across multiple 225 species and the Wuhan-Hu-1 reference sequence (MN908947). Since SARS-CoV-2 first spread to humans, 226 it has continued to mutate and evolve rapidly, particularly regarding the S protein (Amicone et al. 2022; 227 Liu et al. 2022; Tay et al. 2022). Like the mutations leading to the Wuhan strain caused an increase in 228 binding affinity to hACE2, continued evolution in human hosts has resulted in further altered binding

229 affinities as well as different phenotypic outcomes for those infected (Ali et al. 2021; Barton et al. 2021;

230 Ozono et al. 2021).

231 We explore whether protein dynamics has played a role in the selection of mutational sites during the 232 evolution of the S protein since 2019. Our previous work has indicated that rate of evolution per positional 233 site exhibits a positive correlation with positional flexibility; generally, positions that exhibit higher 234 flexibility are also sites that experience a higher number of amino acid substitutions (Liu and Bahar 2012; 235 Maguid et al. 2008; Maguid et al. 2006; Mikulska-Ruminska et al. 2019; Nevin Gerek, Z., Kumar, S., Banu 236 Ozkan, S. 2013). To confirm these findings for the evolution of the S protein using the sequenced variants 237 of infected humans, we analyze the site-specific amino acid flexibility using the Dynamic Flexibility Index 238 (DFI). Using the same mathematical foundation as DCI, DFI evaluates each position's displacement 239 response to random force perturbations at other locations in the protein (Gerek and Ozkan 2011; Nevin 240 Gerek, Z., Kumar, S., Banu Ozkan, S. 2013), and it can be considered a measure of a given position's ability 241 to explore its local conformational space. We found that the covid-19 S protein shows the expected high 242 correlation between the occurrence of mutations and site flexibility (Figure 3) when we compare %DFI 243 (DFI ranked by percentile) to the average number of variants per position found within a given %DFI bin. 244 Previous studies have indicated that rigid residues are critical for functional dynamics, thus more likely to 245 impact function if mutated and, generally, can lead to a loss of function and thus more conserved (Kim, 246 H. et al. 2015; Butler et al. 2018; Modi, Risso, et al. 2021; Modi, Campitelli, et al. 2021; Kazan et al. 2022; 247 Ose, Butler, et al. 2022; Stevens et al. 2022; Campitelli et al. 2020a; Kumar, A., Glembo, T.J., Ozkan, S.B. 248 2015b). This analysis also agrees with these previous studies and highlights the power of negative 249 selection, in line with the neutral theory of molecular evolution, stating that deleterious mutations (i.e., 250 those on the rigid positions) should be eliminated and therefore not observed (Kimura, Motoo 1983).

252 Figure 3. The average number of variants observed among residues of different flexibility. Residues were 253 sorted into one of five bins based on flexibility. After that, the average number of variants for residues 254 within that bin was calculated. Here, the number of variants is defined as the number of different amino 255 acid varieties found at that site. Mutational data was calculated across approximately 24,000 SARS-CoV-256 2 S protein sequences from the NCBI Datasets Project (Brister et al. 2015). Residue flexibility, as reported 257 here via %DFI, was computed using structure id 6vsb from the Protein DataBank (Berman, H.M. et al. 258 2000). More rigid residues tend to have fewer variants. $(r = 0.94)$.

259 Continued mutations within human hosts have resulted in a multitude of variants. Indeed, by fitting 260 various molecular clock models to genome sequence data, VOC emergence is punctuated by an episodic 261 period of rapid evolution, with a substitution rate of up to 4-fold greater than the background substitution 262 rate (Kumar et al. 2021; Tay et al. 2022). With such an aggressive evolutionary rate, we are finding VOCs 263 to consist of a number of different characteristic mutations, almost all of which are CAPs.

264 To explore the dynamic effects of the evolution of the Spike in humans, we examine asymmetry with these 265 new potentially adaptive sites, namely the low EP (CAP) characteristic mutation sites observed in the Delta 266 variant, the widely dominant variant from December 2021 to January 2022 (Thye et al. 2021), and the 267 Omicron variant, a highly transmissible variant whose lineages have remained dominant since January 268 2022) (Kim et al. 2021) (Figure 4). This analysis revealed a mechanism similar to that for the CAPs in the 269 reference protein (Figure 2), as the open-chain binding interface is also allosterically controlled by these 270 potentially new adaptive sites. Regarding this, we see that the asymmetry is much more pronounced in 271 observed mutations of Omicron variants suggesting that these new mutations have a stronger power in 272 controlling the dynamics of open chain hACE2 binding interface compared to those observed in Delta 273 variants. We can speculate that the difference in virulence and infection rates between Omicron and Delta 274 (Earnest et al. 2022; Bager et al. 2021; Sheikh et al. 2021; Twohig et al. 2022;(Houhamdi et al. 2022; Menni 275 et al. 2022) might be due to these specific CAPs within each variant and their differences in allosterically 276 controlling the dynamics of open RBD binding sites as observed in the DCI $_{\text{asym}}$ analysis.

278 Figure 4. (A) DCI asymmetry with low EP characteristic mutation sites of Delta or Omicron strains in the 279 closed chains and the binding interface of RBD in the open chain. Delta displays a second peak closer to 280 zero, suggesting that Delta mutation sites (M = -0.98, SD = 0.80) have less allosteric control over the 281 hACE2 binding sites than Omicron mutation sites (M = -1.74, SD = 1.00) (p < .001). However, both sets 282 of sites have far more control over hACE2 binding sites than expected, based on a random control group 283 (M = 0.03, SD = 0.85) (p < .001). (B) S protein structure showing binding interface sites (transparent gray), 284 Delta mutation sites (magenta), Omicron mutation sites (cyan), and sites mutated in both Omicron and 285 Delta (Blue).

287 Experimental results motivate the use of EpiScore within the SARS-CoV-2 Spike protein.

288 The fact that the identified CAPs in the reference protein and the more recently evolved CAPs of Delta 289 and Omicron variants both show a high degree of control over the active sites begs the question: what is 290 the complex interaction between these previous and new CAP sites? Motivated by this concept, we 291 explore the interplay of mutational pairs to understand the effects of the specific amino acid backgrounds 292 associated with these two predominant variants. Some CAP sites in Delta and Omicron have already been 293 considered adaptive (Kemp et al. 2021; Kistler et al. 2022; Maher et al. 2022; Neher 2022).

294 It is well understood that the impact of even a single mutation to a protein sequence can sometimes 295 dramatically alter the biophysical behavior of the system. However, the mechanistic impact of point 296 mutations can only be fully understood when the sequence background upon which it is made is 297 accounted for. This means that, in the case of strains with multiple mutations, the interplay between 298 mutated positions will ultimately impact a protein as an aggregate behavior, where the presence of 299 previous mutations may strongly (or weakly) influence some mutations. This concept of non-additivity is 300 known as epistasis. In fact, studies of evolutionary pathways of mutations have suggested that a majority 301 of the mutations have a second or a higher order epistasis among them (Bershtein et al. 2006). Nature 302 exploits this higher order complex relationship between the mutations to evolve their function.

303 To computationally capture and interpret the pairwise effects of mutations, we have developed an in-304 house computational tool called EpiScore (Figure 5A). Here, we evaluate how a given position pair *i* j may 305 affect other critical positions k of the protein. EpiScore is the relative coupling strength to a position k 306 when positions *i* and *j* are perturbed *simultaneously* compared to the average dynamic coupling strength 307 of i to k and j to k. EpiScore has previously been used successfully to capture overarching trends in GB1 308 deep mutational scan data as well as specific instances of the development of antibiotic resistance in 309 various enzymatic systems (Campitelli and Ozkan 2020). An EpiScore of 1 indicates perfect coupling 310 additivity, and deviations from this value represent non-additive behavior between position pairs and

311 functionally important sites. Prior EpiScore work has shown a difference in EpiScore between the sites of 312 compensatory and non-compensatory mutations, where both yield distributions with peaks around 1, but 313 non-compensatory mutations show higher deviation in their EpiScore distribution (Ose, Campitelli, et al. 314 2022).

315 Many studies have confirmed epistasis between residues within the S protein (Moulana et al. 2022a; Starr, 316 Greaney, Hannon, et al. 2022; Moulana et al. 2023; Witte et al. 2023). These epistatic residues can have 317 various effects on hACE2 or antibody binding. To further motivate our use of EpiScore within SARS-CoV-318 2, we calculate the EpiScore (Figure 5B) of a set of mutation pairs used by Moulana et al. (Nature comm 319 2022) and compare our results to quantified epistatic effects determined by the experimental hACE2 320 binding affinity of "first-order" single mutation variants compared to "second-order" mutation pair. Our 321 EpiScore results and the experimentally determined epistasis have a reasonable similarity. Both methods 322 captured highly epistatic behavior among residues 493, 496, 498, 501, and 505, as well as a lack of epistatic

323 behavior for residues 339, 371, 373, 324 and 375.

325 **Figure 5**. (A) Schematic
EpiScore: $\%$ DCI_{FIII} 22 representation of cross-chain $\frac{1}{2}$ (%DCI_{ik} + %DCI_{ik}) EpiScore, describing i, j, in chain B and 328 C respectively and its impact in RBD 329 binding position k in the open RBD 330 conformer chain A. (B) Colors indicate 331 EpiScore values for given mutation 332 pairs, averaged over hACE2 binding 333 sites. Cross-chain residue pairs in the 334 upper right tend to be highly 335 epistatic, similar to pairwise second-336 order interaction coefficients from 337 Moulana et al. (Nature comm 2022).

338 Episcore highlights the epistatic relationship between the recent adaptive mutations in VOCs and the

339 CAPs of the Wuhan reference.

340 Seeking further to understand the role of epistasis within S protein variants, we explored the possibility 341 of epistatic relationships between the CAPs of the Wuhan variant and the new CAPs in VOCs. Thus, we 342 computed the Episcore of these CAP positions in the closed RBD domains (i.e., chain B and C) with respect 343 to functional hACE2 binding interface sites of the open RBD domain chain (chain A) (Figure 5B) and 344 obtained Episcore distributions.

347 **Figure 6.** EpiScores with I = low EP Delta mutation sites (magenta), low EP Omicron mutation sites (Cyan), 348 and a random selection of sites (gray), $j = low$ EP sites in the Wuhan variant, and $k = the$ binding interface 349 of the open chain. EpiScores using sites of either variant (Delta: M = 0.70, SD = 0.50, Omicron M = 0.86, 350 SD = 0.46) are significantly different (p<.001) from a set of EpiScores using random sites (M = 0.84, SD = 351 0.41), but the distribution for Delta variants differs much more from the other two. EpiScores for other 352 variants can be found in Supplementary Figure S1.

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354 To contrast these variants, Omicron (Figure 6, cyan) shows a high proportion of additive, potentially 355 compensatory, mutations compared to the Delta variant (Figure 6, magenta), with a peak centered on 1. 356 The comparatively more pathogenic Delta variant exhibited many non-additive, suspected non-357 compensatory mutations with EpiScores below one. This again suggests that the cross-communication 358 between the open and closed chain of the S protein is important for regulating the function. Four out of 359 seven low EP Delta mutation sites used in this analysis often resulted in EpiScores below 1. Each of those 360 is found in the N-terminal domain (NTD) on or near the N3 loop and is implicated in antibody escape in 361 recent studies (Chi et al. 2020; Weisblum et al. 2020; Harvey, Carabelli, Jackson, Gupta, Thomson, 362 Harrison, Ludden, Reeve, Rambaut, COVID-19 Genomics UK (COG-UK) Consortium, et al. 2021; Klinakis et 363 al. 2021; Cantoni et al. 2022). The low EpiScores of NTD mutations suggest that they dampen the control 364 of Wuhan variant CAPs over the active site in addition to their effects on antibody binding. It is possible 365 that what the Delta variant gained in transmission rate also came with being more harmfully pathogenic 366 due in part to negatively epistatic interactions. It follows that the mutations leading to the development 367 of the Omicron strain were compensatory in nature, possibly leading to a lower pathogenicity and higher 368 effective immune escape, resulting in a higher transmission rate. It is worth noting that other variants 369 contain NTD mutations which result in low EpiScores, however the proportion of these mutations within 370 the set is considerably less than in Delta (Supplementary Figure S2).

371 One of the more notable similarities of generated EpiScore distributions is a tail of EpiScore values upward 372 of 2.0, indicating highly epistatic behavior. Interestingly, these tails are largely due to three different CAPs: 373 346R, 486F, and 498Q. Those residues are nearby one another within the protein structure and have been 374 reported to play a role in antibody binding, either being known antibody binding sites (346R and 486F) or 375 having received very high antibody accessibility scores (498Q) (Harvey, Carabelli, Jackson, Gupta, 376 Thomson, Harrison, Ludden, Reeve, Rambaut, COVID-19 Genomics UK (COG-UK) Consortium, et al. 2021; 377 Raghuvamsi et al. 2021). These observed high Episcore values also support the other studies that the 378 epistatic interactions between these CAPs and the mutations of the VOCs within the S protein are crucial 379 for maintaining binding affinity of hACE2 whilst evading immunity (Hong et al. 2022; Moulana et al. 2022b; 380 Starr, Greaney, Stewart, et al. 2022).

381 Viewing EpiScores of Delta and Omicron potentially adaptive mutation sites with only CAP site 486F (a 382 binding site for both hACE2 and antibodies) (Huang et al. 2020; Ali et al. 2021; Harvey, Carabelli, Jackson,

383 Gupta, Thomson, Harrison, Ludden, Reeve, Rambaut, COVID-19 Genomics UK (COG-UK) Consortium, et 384 al. 2021; Raghuvamsi et al. 2021) shows highly epistatic interactions at other hACE2 binding sites (Figure 385 6). However, within a recent and rapidly spreading subvariant of Omicron, XBB 1.5, we see a mutation of 386 F to S, a rare double nucleotide mutation, at site 486. This new variant has unprecedented immune escape 387 capabilities, resisting neutralizing antibodies almost entirely (Qu et al. 2023). EpiScores of other XBB 1.5 388 specific mutation sites with 486S are almost entirely greater than 1, showing an even higher degree of 389 epistasis with the binding sites of RBD (Figure 7). These results present a threefold importance for the 390 F486S mutation: Not only does this residue alter antibody (i.e., immune escape) and hACE2 binding by 391 directly modifying a binding site, but it may also be responsible for modifying hACE2 binding via epistatic 392 cooperation with other co-occurring mutations.

Figure 7. EpiScores with $i =$ low EP Delta mutation sites (magenta), low EP Omicron mutation sites (Cyan), and a random selection of sites (gray), $i =$ site 486, and $k =$ the binding interface of the open chain. CAP and hACE2 and antibody binding site 486 displays epistasis with 2.0 almost all XBB 1.5 variant

406 binding site (M = 1.40, SD = 0.46) and presents a significantly different profile from other variant sites (p 407 <.01). EpiScores involving 486 for other variants can be found in Supplementary Figure S1.

409 Change in flexibility of RBD binding site correlates with experimental binding affinities for Omicron and

410 Omicron XBB variants.

411 Experimental studies have tracked hACE2 binding for different variants since the virus first spread (Ali et 412 al. 2021; Barton et al. 2021; Ozono et al. 2021; Wu et al. 2022). Within the Omicron variant, for example, 413 characteristic mutations on the RBD are shown to increase the overall binding affinity of the virus to the 414 ACE2 receptor, which is suspected to allow it to spread more easily (Kim et al. 2021). Furthermore, the 415 new Omicron XBB and Omicron XBB 1.5 variants contain additional mutations in the RBD and antibody 416 binding residues, which may further impact their dynamics and interactions with the host. 417

419 Figure 8. The %DFI calculations for variants Omicron, XBB, and XBB 1.5. (A) %DFI profile of the variants 420 are plotted in the same panel. The grey shaded areas and dashed lines indicate the ACE2 binding regions, 421 whereas the red dashed lines show the antibody binding residues. (B) The sum of %DFI values of RBD-422 ACE2 interface residues. The trend of total %DFI with the log of K_d values overlaps with the one seen with 423 the experiments (R=0.97). (C) The RBD antibody binding residues are used to calculate the sum of %DFI. 424 The ranking captured with the total %DFI agrees with the log of IC50 values from the experiments.

425 To gain deeper insights into the impact of dynamics on the binding affinity of hACE2 and antibodies with 426 the recent Omicron XBB variants, we conducted molecular dynamics (MD) simulations. By analyzing the 427 resulting trajectories, we investigated how these mutations influence the flexibility and rigidity of the RBD 428 and antibody binding residues, consequently affecting their binding affinity and potential for immune 429 evasion (Figure 8). To understand the overall flexibility changes, we measured the sum of DFI of the ACE2 430 binding residues, as well as the sum of DFI of the antibody binding residues, calculated from the MD 431 trajectories and compared then with experimental viral binding (disassociation constants) and immunity 432 evasion antibody IC50 values (Yue et al. 2023).

433 This investigation elucidated the impact of mutations in the receptor-binding domain (RBD) and antibody 434 binding residues on the binding affinity of the S protein and immune evasion by modulating their flexibility 435 and rigidity (Figure 8A). The Omicron XBB variant exhibits heightened flexibility in hACE2 and antibody 436 binding residues, reducing infectivity and enhancing immune evasion. Conversely, the Omicron XBB 1.5 437 variant induces distinct dynamics in these regions, rendering the RBD-ACE2 interface more rigid while 438 increasing flexibility in antibody binding residues. These effects indicate that Omicron XBB1.5 retains its 439 antibody escape capabilities while regaining ACE2 binding affinity comparable to previous Omicron 440 variants, in accordance with experimental findings (Yue et al. 2023). These findings suggest that mutations 441 in the RBD and antibody binding residues can have complex effects on the dynamics of the protein and, 442 ultimately, on the virus's ability to infect and evade the host immune system through an alteration of 443 biding site dynamics.

444

445 Conclusion:

446 We analyzed the evolutionary trajectory of the CoV-2 S protein in humans to understand the dynamic 447 and epistatic interactions of the mutations defining specific VOCs within the S protein. We first obtain 448 the phylogenetic tree of the COV-2 S protein and identify the sites of certain recent mutations known as

473 Long-ranged interactions between different sites within a given protein is critically important for protein 474 function (Peters and Lively 1999; Bershtein et al. 2006; Collins et al. 2006; Ekeberg et al. 2013; Levy et al. 475 2017; Harrigan et al. 2018; Otten et al. 2018; Rojas Echenique et al. 2019; Shimagaki and Weigt 2019; de 476 la Paz et al. 2020; Rizzato et al. 2020; Yang et al. 2020; Bisardi et al. 2022) and for the CoV-2 S protein in 477 particular (Zeng et al. 2020; Castiglione et al. 2021; Dong et al. 2021; Garvin et al. 2021; Nielsen et al. 478 2022; Ramarao-Milne et al. 2022; Rochman et al. 2022; Rodriguez-Rivas et al. 2022). By showing dynamic 479 differences between the interactions of CAPs, which have likely played a major role in allowing the virus 480 to infect human hosts, the binding site, and the characteristic mutations of dominant Delta and Omicron 481 strains, we see a "fine-tuning" of protein behavior. As variants continue to evolve, Omicron sub-variants 482 are of growing concern due in large part to further increased immune evasion (Callaway 2022; Wang, Guo, 483 et al. 2022; Wang, Iketani, et al. 2022), and we observe that the new mutations observed in antibody 484 binding sites yield more epistatic interaction with the CAPs. In addition to supporting previous dynamic 485 research on the S protein, this analysis provides the insight that CAP sites are of continued importance to 486 protein function and should be given special attention when considering the impact of future mutations.

487

488 Methods

489

490 Dynamic Flexibility and Dynamic Coupling

491 The Dynamic Flexibility Index utilizes a Perturbation Response Scanning technique that combines the Elas-492 tic Network Model (ENM) and Linear Response Theory (LRT) (Gerek and Ozkan 2011; Nevin Gerek, Z., 493 Kumar, S., Banu Ozkan, S. 2013). In ENM, the protein is considered as a network of beads at $C\alpha$ positions 494 interacting with each other via a harmonic spring potential. Using LRT, ∆R is calculated as the fluctuation 495 response vector of residue *i* due to unit force's **F** perturbation on residue *i*, averaged over multiple unit 496 force directions to simulate an isotropic perturbation.

516 position in the structure.

519
$$
DFI_{i} = \frac{\sum_{j=1}^{N} |\Delta R^{j}|_{i}}{\sum_{i=1}^{N} \sum_{j=1}^{N} |\Delta R^{j}|_{i}}
$$

 518 (4)

520 It is also often useful to quantify position flexibility relative to the flexibility ranges unique to individual 521 structures. To that end, DFI can be presented as a percentile rank, %DFI. All %DFI calculations present in 522 this work used the DFI value of every residue of the full spike structure for ranking. The DFI parameter can 523 be considered a measure of a given amino acid position's ability to explore its local conformational space.

524

525 Dynamic Coupling Index

526 Similar to DFI, the dynamic coupling index (DCI) (Larrimore et al. 2017; Kumar, A., Glembo, T.J., Ozkan, 527 S.B. 2015b) also utilizes Perturbation Response Scanning with the Elastic Network Model and Linear 528 Response Theory. DCI captures the strength of displacement response of a given position *i* upon 529 perturbation to a single functionally important position (or subset of positions) j, relative to the average 530 fluctuation response of position *i* when all of the positions within a structure are perturbed.

531

534
$$
DCI_{ji} = \frac{\sum_{j}^{N_{\text{functional}}} |\Delta R^{j}|_{i}/N_{\text{functional}}}{\sum_{j=1}^{N} |\Delta R^{j}|_{i}/N}
$$

 532 (5)

533 When only positional pairs are concerned, this expression reduces to:

537
$$
DCI_{ji} = \frac{|\Delta R^{j}|_{i}}{\sum_{j=1}^{N} |\Delta R^{j}|_{i}/N}
$$

$$
536 \t\t (6)
$$

540 One of the most important aspects of DCI is that the entire network of interactions is explicitly included 541 in subsequent calculations without the need for dimensionality reduction techniques. If one considers 542 interactions such as communication directionality or dynamic coupling regulation between position pairs 543 as inherent properties of an anisotropic interaction network, it is critical to include the interactions of the 544 entire network to accurately model the effect one residue can have on another.

545 Here, we present two further extensions of DCI which allow us to uniquely model coupling directionality 546 and epistatic effects: DCI_{asym} and EpiScore, respectively. Interestingly, we can capture asymmetry between 547 different residues within a protein through DCI, as a coupling in and of itself is asymmetric within an ani-548 sotropic network. That is, each amino acid has a set of positions to which it is highly coupled, and this 549 anisotropy in connections gives rise to unique differences in coupling between a given *i j* pair of amino 550 acids which do not have direct interactions (Figure 2A). DCI_{asym}, then, is simply DCI_{ii} (the normalized dis-551 placement response of position *j* upon a perturbation to position *i*) – DCI_{ii} (Equation (7)). Using DCI_{asym} we 552 can determine a cause-effect relationship between the i j pair in terms of force/signal propagation be-553 tween these two positions.

554

 $DCI_{asym} = DCI_{ij} - DCI_{ji}$

- 555 (7)
-
- $\% \text{DCI}_{\text{asvm}} = \% \text{DCI}_{\text{ii}} \% \text{DCI}_{\text{ii}}$
- 557 (8)

559 where a positive DCI_{asym} value indicates communication from position *i* to position *j*.

560 EpiScore can identify or describe potential non-additivity in substitution behavior between residue pairs.

561 This metric can capture the differences in a normalized perturbation response to a position k when a force

562 is applied at two residues *i* and *j* simultaneously versus the average additive perturbation response when

563 each residue *i, j,* is perturbed individually (Figure 5A, Equation 9).

566
EpiScore =
$$
\frac{\%DCI_{[ij]k}}{\frac{1}{2}(\%DCI_{ik} + \%DCI_{jk})}
$$

564

- 565 (9)
- 567

568 EpiScore values < 1 (> 1) indicate that the additive perturbations of positions i and j generates a greater 569 (lesser) response at position k than the effect of a simultaneous perturbation. This means that, when 570 treated with a simultaneous perturbation at both sites i and j, the displacement response of k is lower 571 (higher) than the average effect of individual perturbations to *i* and *j*, one at a time. As EpiScore is a linear 572 scale, the further the value from 1, the greater the effect described above.

573 Molecular Dynamics (MD)

574 The production simulations of the variants Omicron, Omicron XBB, and Omicron XBB 1.5 were generated 575 using the AMBER software package. The mutations in the variants were modeled using PYMOL taking the 576 template as PDB 6M0J. The initial input proteins were parametrized utilizing the ff14SB force field (Maier 577 et al. 2015). To ensure adequate solvation of the protein, the solvation box was defined to encompass the 578 protein, maintaining a minimum distance of 16Å from the protein to the box edges, utilizing the explicit 579 TIP3P water model (Sun 1995). The neutralization of the solvated system was achieved through the addi-580 tion of sodium and chloride ions. The system was subjected to a steepest descent algorithm for 11000

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- 985 Supplemental Figures
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988 Supplementary figure S1. EpiScores with i = characteristic mutation sites, j = low EP sites, and k =

989 the binding interface of the open chain. EpiScores using variant sites are significantly different

990 (p<.001) from a set of EpiScores using random sites.

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993 Supplementary figure S2. EpiScores with i = characteristic mutation sites within the NTD, j = low

994 EP sites, and k = the binding interface of the open chain. NTD domain mutation sites result in

995 markedly lower EpiScores compared to elsewhere.

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1000 Supplementary figure S3. EpiScores with i = characteristic mutation sites, j = site 486, and k = the

1001 binding interface of the open chain. CAP and hACE2 and antibody binding site 486 displays

1002 epistasis with almost all XBB 1.5 variant sites at almost every hACE2 binding site. EpiScores

1003 using variant sites are significantly different (p<.001) from a set of EpiScores using random sites.

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