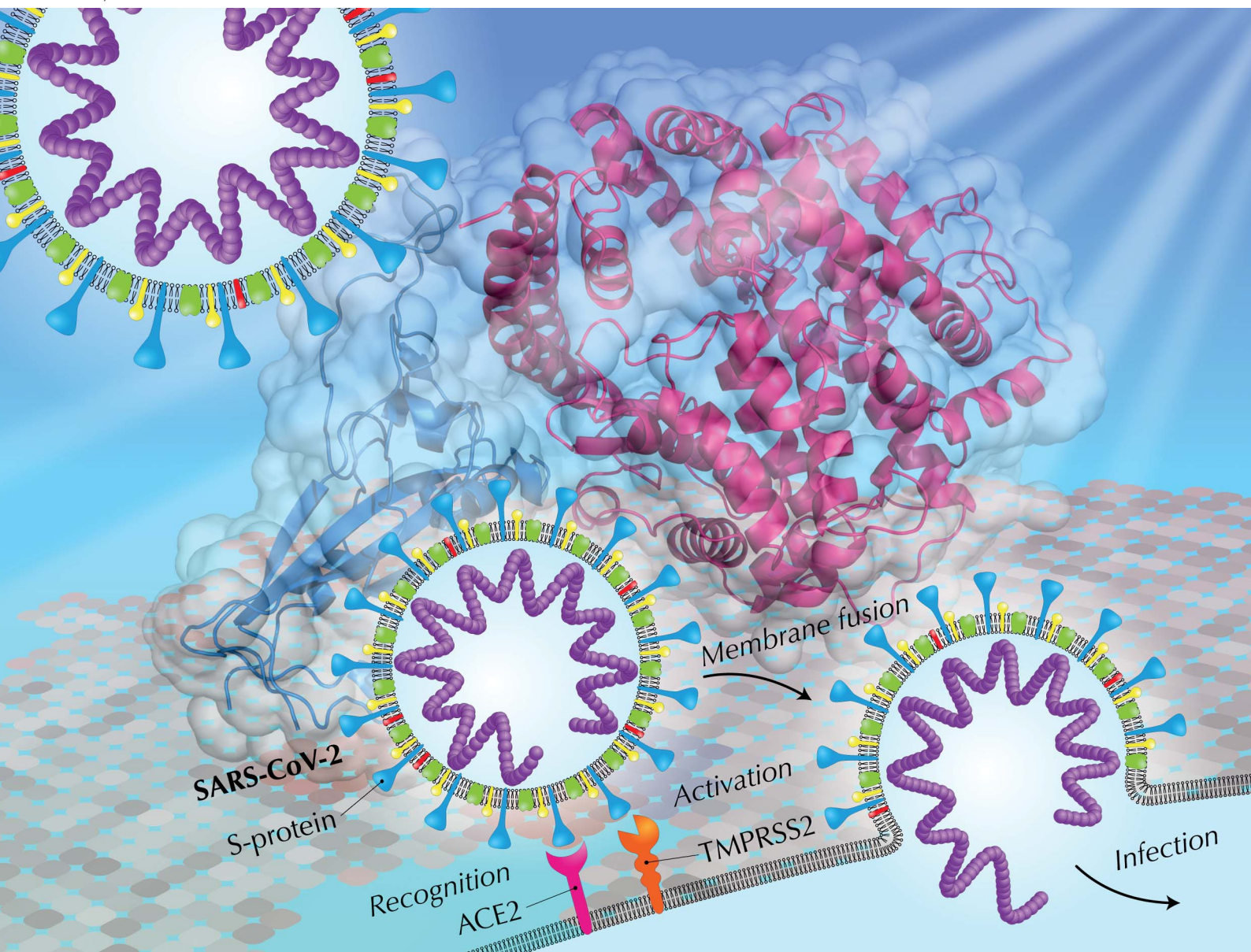


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Molecular recognition of SARS-CoV-2 spike glycoprotein: quantum chemical hot spot and epitope analyses†

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Due to the COVID-19 pandemic, researchers have attempted to identify complex structures of the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) spike glycoprotein (S-protein) with angiotensin-converting enzyme 2 (ACE2) or a blocking antibody. However, the molecular recognition mechanism—critical information for drug and antibody design—has not been fully clarified at the amino acid residue level. Elucidating such a microscopic mechanism in detail requires a more accurate molecular interpretation that includes quantum mechanics to quantitatively evaluate hydrogen bonds, XH/ π interactions (X = N, O, and C), and salt bridges. In this study, we applied the fragment molecular orbital (FMO) method to characterize the SARS-CoV-2 S-protein binding interactions with not only ACE2 but also the B38 Fab antibody involved in ACE2-inhibitory binding. By analyzing FMO-based interaction energies along a wide range of binding interfaces carefully, we identified amino acid residues critical for molecular recognition between S-protein and ACE2 or B38 Fab antibody. Importantly, hydrophobic residues that are involved in weak interactions such as CH–O hydrogen bond and XH/ π interactions, as well as polar residues that construct conspicuous hydrogen bonds, play important roles in molecular recognition and binding ability. Moreover, through these FMO-based analyses, we also clarified novel hot spots and epitopes that had been overlooked in previous studies by structural and molecular mechanical approaches. Altogether, these hot spots/epitopes identified between S-protein and ACE2/B38 Fab antibody may provide useful information for future antibody design, evaluation of the binding property of the SARS-CoV-2 variants including its N501Y, and small or medium drug design against the SARS-CoV-2.

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1. Introduction

Coronavirus disease 2019 (COVID-19), caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), has been declared a pandemic by the World Health Organization and has caused worldwide social and economic problems.¹ Despite its

significant infectious strength and the worldwide research efforts, to date, no specific treatment has been established against this new virus. The SARS-CoV-2 is composed of a variety of proteins, including the spike glycoprotein (S-protein),² which is believed to promote the invasion of host cells and the proliferation of the virus by binding to human angiotensin-converting enzyme 2 (ACE2) (Fig. 1A). The SARS-CoV-2 S-protein monomer

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† Electronic supplementary information (ESI) available: Table S1: IFIE and geometric IFP of XH–Y hydrogen bonds between SARS-CoV-2 S-protein and ACE2. Table S2: IFIE and geometric IFP of XH/ π interactions between SARS-CoV-2 S-protein and ACE2. Table S3: IFIE and geometric IFP of XH–Y

hydrogen bonds between SARS-CoV-2 S-protein and B38 Fab antibody. Table S4: IFIE and geometric IFP of XH/ π interactions between SARS-CoV-2 S-protein and B38 Fab antibody. Table S5: Predicted binding energies between the S-protein and ACE2/B38 Fab antibody without the IFIEs of sugar chain fragments. Table S6: Predicted binding energies (kcal mol^{−1}) between SARS-CoV-2 S-protein WT or N501Y variant and ACE2. Fig. S1: Structural differences on the binding interface of S-protein and ACE2. Fig. S2: Intermolecular interaction energies between SARS-CoV-2 chimeric S-protein and ACE2 in host cells. Fig. S3: Intermolecular interaction energies between SARS-CoV S-protein and ACE2 in host cells. Fig. S4: Intermolecular interaction energies between SARS-CoV-2 S-protein and B38 Fab antibody. Fig. S5: Hot spots of molecular recognition between ACE2 and SARS-CoV-2 S-protein. Fig. S6: Structure difference between the wild-type and N501Y mutant of SARS-CoV-2 S-protein. Fig. S7: Differences of IFIEs of ACE2 between the wild type (N501) and the variant (Y501) on SARS-CoV-2 S-protein. See DOI: 10.1039/d0sc06528e.

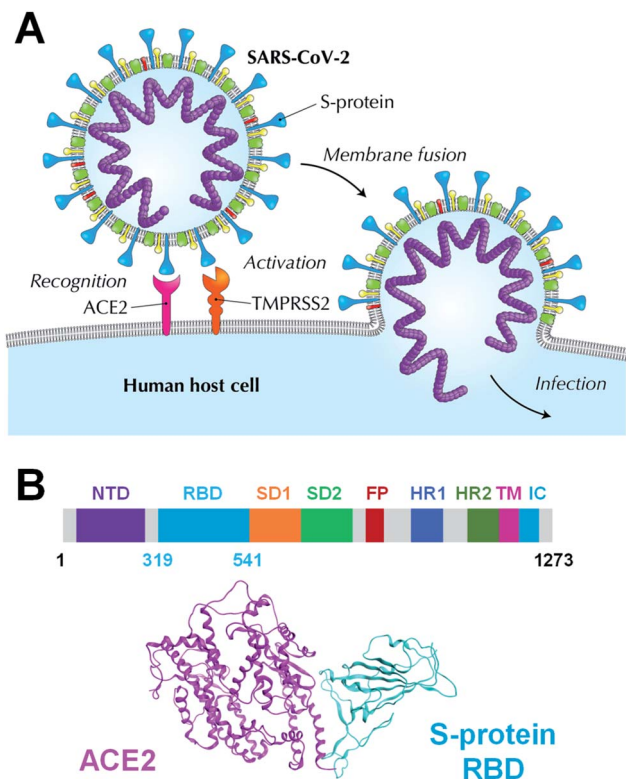


Fig. 1 Human host cell infection via ACE2 recognition by SARS-CoV-2 S-protein. (A) ACE2 is the host cell receptor responsible for mediating the SARS-CoV-2 infection, the new coronavirus responsible for coronavirus disease 2019 (COVID-19). (B) Overall topology of the SARS-CoV-2 S-protein monomer that comprises an N-terminal domain (NTD), receptor-binding domain (RBD), subdomain 1 (SD1), subdomain 2 (SD2), fusion peptide (FP), heptad repeat 1 (HR1), heptad repeat 2 (HR2), transmembrane region (TM), and an intracellular domain (IC). A complex between the SARS-CoV-2 S-protein and ACE2 is shown in cyan and magenta ribbon models.

comprises nine domains/regions: N-terminal domain (NTD), receptor-binding domain (RBD), subdomain 1 (SD1), subdomain 2 (SD2), fusion peptide (FP), heptad repeat 1 (HR1), heptad repeat 2 (HR2), transmembrane region (TM), and intracellular domain (IC) (Fig. 1B). The RBD domain plays a key role in cell infection *via* ACE2 recognition. Therefore, the SARS-CoV-2 S-protein is one of the target proteins used for antibody-drug design to prevent virus invasion, including S-protein neutralizing antibodies.^{2–5} To tackle the SARS-CoV-2 pandemic, researchers are extracting and designing neutralizing antibodies to block the binding of S-protein to ACE2.^{2–5} Understanding the binding mode of S-protein with ACE2 and existing neutralizing antibodies will help design potent neutralizing antibodies. Moreover, a better understanding of the molecular recognition differences of ACE2 among the SARS-CoV-2, its variant and other coronaviruses including SARS-CoV can improve the antibody-drug design. Several SARS-CoV-2 variants have already emerged and are rampant throughout the world.^{6–8} Thus, certain X-ray crystallographic and electron microscopic analyzes have already been performed, and geometry-based interactions have been discussed in previous studies.^{3,9–13}

Analyses of molecular recognition hot spots between the SARS-CoV-2 S-protein and ACE2 were conducted by several computational simulations, including electrostatic potential,^{14,15} molecular mechanics (MM)-based interaction energy analysis with classical force field,¹⁶ and quantum mechanics (QM)-based interaction energy analysis with or without evaluation of semiempirical dispersion energy.^{17–19} Molecular dynamics (MD) simulations for the complex formed by the S-protein and ACE2 were conducted to better understand their association.^{20–26} Approaches for drug/neutralizing antibody design targeting S-protein have been reported, such as a *de novo* design peptide inhibitors for SARS-CoV-2,^{27,28} a virtual screening of antiviral compounds for the SARS-CoV-2 (ref. 29) and sequence-based epitope analysis for the SARS-CoV-2, among others.³⁰ These previous studies have been based on the molecular recognition of S-protein and ACE2/antibody and were highly beneficial. These studies had focused on more on hydrogen bonds, which certainly play key roles in determining binding poses of a protein–protein interaction (PPI) system, than on hydrophobic interactions, which influence the binding affinity.³¹ Even when hydrophobic interactions were considered, they were limited by the non-quantitative or inaccurate evaluation using a structure-based, MM-based, or semi-empirical based analysis. Moreover, in the prediction of binding ability, the enthalpic interactions, such as electrostatic and dispersion interactions, as well as the effects of solvation and sugar chain have not been examined sufficiently. Therefore, we used *ab initio* electron-correlated QM theory^{32–35} that yields rigorous dispersion energy in order to quantitatively and accurately evaluate hydrophobic interactions such as CH/ π interactions.

The QM calculation of whole protein system consisting of hundreds to thousands of residues can be conducted using the fragment molecular orbital (FMO) approach.^{36–39} The hydrogen bonds, electrostatic interactions, salt bridges, and hydrophobic interactions were quantitatively analyzed using the inter-fragment interaction energy (IFIE) and its energy decomposition analysis (PIEDA) based on FMO calculations.^{40–42} In recent years, the FMO method has been widely used as a drug discovery tool.^{35,43–52} Recent studies have also provided detailed information on the underlying antigen–antibody or ligand-binding characteristics of several drug-targeted proteins, main protease,^{53–55} ribonucleic acid (RNA)-dependent RNA polymerase,⁵⁶ S-protein,^{17,19} in COVID-19. To date, the FMO method has been successfully used for identifying key amino acid residues and sugar chains for PPI, such as molecular recognition for an influenza hemagglutinin of a fragment antigen-binding (Fab) antibody^{57–67} and a measles hemagglutinin of a signaling lymphocytic activation molecule.^{68,69} In addition, it is essential to combine desolvation energy and enthalpic binding energy to predict binding ability.^{43,51} Therefore, the present study aimed to address additional calculations that could shed light on the molecular process underlying the SARS-CoV-2, the SARS-CoV-2 variant and SARS-CoV S-protein recognition by ACE2 or B38 Fab antibody, which remains to be understood in more detail, in particularly concerning hydrophobic interactions such as CH/ π interactions. In particular, the study explored the amino acid residues involved in, as well as types of interactions that make, hot spots for S-protein binding to ACE2.



This FMO-based hot spot and epitope analysis can be applied to other PPI systems in drug-antigen design as well.

2. Computational methods

To clarify the important interactions between the SARS-CoV-2 S-protein and ACE2, interaction energy analysis was performed according to the procedures described below.

2.1 Preparation of the SARS-CoV-2 S-protein and ACE2 complex

We retrieved the crystal structures of the complexes formed by the SARS-CoV-2 S-protein with ACE2 and B38 Fab antibody from the protein data bank (PDB) repository (with its codes of 6LZG and 7BZ5, respectively) for the FMO calculation. To compare the epitope candidate of S-proteins related to the SARS-CoV-2, FMO calculations were also performed for complexes between SARS-CoV-2 chimeric and SARS-CoV S-proteins with ACE2 (PDB IDs: 6VW1 and 2AJF, respectively). We would consider that the binding interface with the ACE2/B38 Fab antibody was robust because the amino acid residues on the binding surface in X-ray crystal structures almost overlap well (Fig. S1†). Thus, we performed the calculations and analysis employing prepared structures based on the X-ray crystal structures. Here, it will consider that the monomer structure of a complex between RBD and ACE2/antibody is suitable as a model to analyze molecular recognition. Although the S-protein existing on the virus surface basically forms a trimer structure,¹³ each RBD of S-protein trimer in the case of an open structure does not retain intramolecular interactions between another monomer of S-protein.⁶⁷ Since the trimer is almost observed as the open structure when it binds to ACE2 or an antibody, the monomer model will be appropriate to analyze molecular recognition between the S-protein and the ACE2/antibody.

The structures of the different viral protein/ACE2 or B38 Fab antibody complexes are shown in Fig. 2. All sugar chains and crystal water molecules were retained. Since Cl^- and Zn^{2+} ions existed far away from the binding surface between the SARS-CoV-2 S-protein and ACE2, they were deleted to simplify the interaction analysis. The missing atoms and missing residues were complemented by structure preparation with the Molecular Operating Environment (MOE) graphical software package (Chemical Computing Group, Montreal, QC, Canada).⁷⁰ Hydrogen atoms were added to each complex using Protonate 3D with MOE, and sequences termini were capped with amine ($-\text{NH}_2$) and carboxylic acid ($-\text{COOH}$) groups. These molecular changes were optimized concurrently using the Amber10:EHT force field with MOE.

Considering the common sequence of the SARS-CoV-2 (C361–E516) and the SARS-CoV (C348–E502), 50 and 36 amino acid mutations were identified in the common sequence and the receptor binding motif, respectively (Fig. 2E). Sequence homology between the SARS-CoV-2 and the SARS-CoV S-proteins was 84.4% for the whole Pro337–Phe515 and 47.2% for Tyr453–Tyr505. The sugar chain located on the SARS-CoV-2 chimeric/SARS-CoV S-protein and ACE2 binding surface was the only β -D-mannopyranose (BMA) extending from the side chain of N90 on ACE2 (Fig. 2B and C). The BMA sugar chain consists

of the terminal chain in the *N*-acetyl- β -D-glucosamine (NAG) and the BMA sugar chains (NAG–NAG–BMA) on N90_{ACE2}. These X-ray structures of ACE2 have a common sequence (S19–A614).

2.2 FMO method and intermolecular interaction energy analysis

The *ab initio* FMO method used was as follows. Briefly, a large molecule or molecular cluster was divided into small fragments and the molecular orbital (MO) calculations for each fragmented monomer and dimer were performed to obtain the properties of the entire system. The many-body effects were considered through the environmental electrostatic potentials. The total energies of the FMO calculations were given by:

$$E_{\text{total}} = \sum_I E'_I + \sum_{IJ} \Delta \tilde{E}_{IJ} \quad (1)$$

where E'_I represents the monomer energy without the environmental electrostatic potential ($\Delta \tilde{E}_{IJ}$) of the IFIE; and I and J are the fragment indices. In addition, PIEDA was used to decompose IFIE into its energy components $\Delta \tilde{E}_{IJ}$ including the electrostatic (ES), exchange–repulsion (EX), dispersion interaction (DI), and charge transfer with higher-order mixed terms (CT+mix), according to:

$$\Delta \tilde{E}_{IJ} = \Delta \tilde{E}_{IJ}^{\text{ES}} + \Delta \tilde{E}_{IJ}^{\text{EX}} + \Delta \tilde{E}_{IJ}^{\text{CT+mix}} + \Delta \tilde{E}_{IJ}^{\text{DI}} \quad (2)$$

Summation of IFIEs (IFIE-sum) over a set of multiple fragments (A) with a fragment J is expressed as following:

$$\Delta \tilde{E}_J^A = \sum_{I \in A} \Delta \tilde{E}_{IJ} \quad (3)$$

By analyzing IFIE-sum over a target protein, one can extract amino acid residues that specifically interact with the target protein, on its binding protein. Moreover, summing over a set of fragments (B) as

$$\Delta \tilde{E}^{AB} = \sum_{J \in B} \Delta \tilde{E}_J^A = \sum_{I \in A, J \in B} \Delta \tilde{E}_{IJ} \quad (4)$$

one can obtain the interaction energy between A and B sets which is used as the binding energy between two proteins in this study.

All FMO calculations were performed using the second-order Møller-Plesset perturbation theory (MP2) with 6-31G* basis set using the ABINIT-MP program.^{38,71,72} The MP2 electron-correlation energy corresponds to the DI term and can accurately and quantitatively evaluate CH/π and π – π interactions that cannot be sufficiently evaluated using the MM and semi-empirical methods. A Cholesky decomposition integral approximation⁷³ was applied to speed up the MP2 calculation while keeping the accuracy. The fragment unit for proteins was each amino acid residue and each sugar chain. During the standard fragmentation process it is desirable to cleave at a sp^3 hybridized single bond; therefore, single amino acid residues were cleaved between $\text{C}\alpha$ and $\text{C}=\text{O}$ in the main chain rather than at the sp^2 hybridized amide bond. Interaction energy between amino acid residues was evaluated using IFIE analysis with BioStation Viewer.⁷⁴ The FMO calculation results of the PDB ID entries



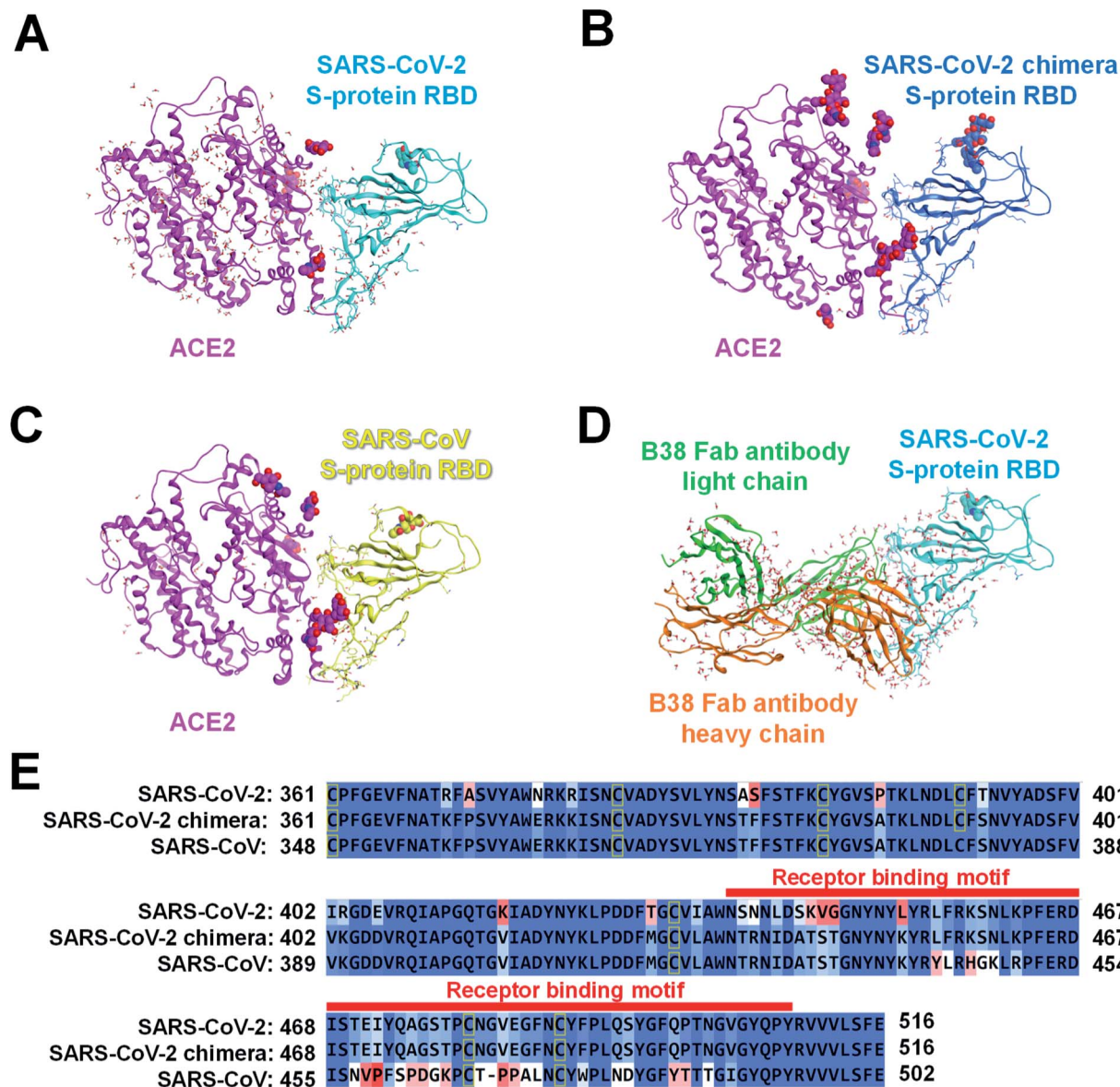


Fig. 2 Complexes of several S-proteins with ACE2 or B38 Fab antibody. The complexed structure of ACE2 (magenta ribbon; Chain A: seqs#19–615, 19–614, and 19–615 in (A), (B), and (C), respectively) with each S-protein is depicted using a ribbon model, and residues whose sequences differ between the three S-proteins are shown in a stick model. The crystal water molecules are shown by the ball and stick model, and the sugar chains are shown by the CPK space-filling model. The S-protein of (A) SARS-CoV-2 (Chain B: seq#333–527), (B) SARS-CoV-2 chimera (Chain E: seq#334–527), and (C) SARS-CoV (Chain E: seq#323–502) are illustrated in the cyan, blue, yellow ribbon model (PDB IDs: 6LZG, 6VW1, and 2AJF), respectively. (D) A complex of B38 Fab antibody (orange ribbon, Chain H, seq#0–217; green ribbon, Chain L, seq#0–215) with SARS-CoV-2 S-protein (cyan ribbon, Chain A, seq#334–528) is shown. (E) In sequence alignment of the SARS-CoV-2, SARS-CoV-2 chimeric, and SARS-CoV S-proteins, the well-aligned and poorly aligned residues are shown in color gradation as blue and red, respectively, evaluated using the BLOSUM62 scoring matrix.⁷⁰ A receptor binding motif on the binding surface with ACE2 is shown by a red belt.

6LZG, 6VW1, 2AJF, and 7BZ5 are registered in the FMO database (FMOB; <https://drugdesign.riken.jp/FMOB/>)^{75–77} with its codes (FMOB IDs) as 4NZVN, KR5L3, 596VZ, and Q86GY, respectively.

2.3 Geometric interaction fingerprint analysis

The hydrogen-bond, ion pair, and π -orbital interactions, such as XH/ π , cation/ π , and π - π interactions (Fig. 3), were detected by geometric interaction fingerprint (IFP) analysis based on the detected interatomic distance and angle using the “prolig_Calculate” function of the MOE software.⁷⁸ MM-based

energy restrictions defined by the “prolig_Calculate” function were employed to extract all possible atom pairs.

3. Results and discussion

3.1. How does the SARS-CoV-2 S-protein recognize ACE2 in human host cells?

In a first approach, a detailed structural analysis was performed to elucidate which key amino acid residues are important for the molecular recognition of ACE2 by the SARS-CoV-2 S-protein,



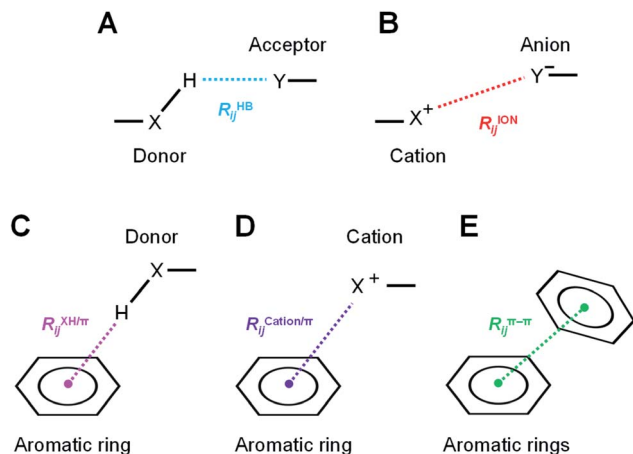


Fig. 3 Types of intermolecular and intramolecular interactions by geometric interaction fingerprint (IFP). Schematic diagrams of the (A) XH–Y hydrogen bond, (B) ion pair interaction, (C) XH/ π interaction, (D) cation/ π interaction, and (E) π – π interaction, respectively. Characteristic distances of each interaction were measured between detected heavy atoms or centroids of aromatic rings.

as well as which types of interactions were involved. Such information could help identify epitope candidates for novel antibodies to prevent the binding between the S-protein and ACE2. FMO-based interaction energy and geometric IFP analyses of the S-protein and ACE2 were performed.

To determine key residues for molecular recognition, it is necessary to identify amino acid residue pairs with attractive interactions between proteins. IFIE and IFIE-sum analyses and their energy decomposition analyses with PIEDA are useful for quantitatively evaluating interaction energies such as hydrogen bonds and CH/ π interactions. In 3.1, 3.2, and 3.3, IFIE-sums over S-protein and ACE2 were calculated using a common amino acid sequence aligned to compare among complexes because the X-ray crystal structures were composed of different lengths of amino acid sequences. Besides, the IFIE-sums did not include IFIEs of sugar chains and water molecules. Fig. 4 shows IFIE-sums of the SARS-CoV-2 S-protein (C361–E516) and ACE2 (S19–A614) using PIEDA. The data revealed that the ES components were dominant in their binding. Six acidic residues (D30, E35, E37, D38, E329, and D355) of ACE2 and five basic residues of the S-protein (R403, R408, K417, R457, and K458) showed a remarkable attractive interaction. These results were considered to be caused by total charges of molecular systems on ACE2 and S-protein of $-26e$ and $+2e$, respectively. In the CT+mix component, attractive interactions were shown by 10 fragment residues of ACE2 (Q24, D30, K31, H34, D38, Y41, Q42, Y83, K353, and G354) and nine residues of the S-protein (K417, Y449, F456, F486, N487, Q493, T500, N501, and G502). In the case of the DI components, characteristic interactions were identified on 15 residue fragments of ACE2 (Q24, T27, F28, D39, K31, H34, E35, D38, Y41, Q42, M82, Y83, K353, G354, and D355) and 14 residue fragments of the S-protein (K417, Y449, Y453, K455, F456, G476, F487, Y489, Q493, Q498, T500, N501, G502, and Y505). Since these amino acid residues are considered to play key roles in the molecular recognition between ACE2 and

the S-protein, the origin of the interaction energy of each fragment pair was further explored (Fig. 5, Section S2†). Nonetheless, it is important to note that in FMO analysis, in general, the ES component is effective over a long distance and tends to be overestimated as compared with the CT and DI ones. Thus, in IFIE-sums with a charged fragment, the weak CT and DI components that show hydrogen bonding and CH/ π interaction tend to be hidden by the ES one. Moreover, a direct comparison of the interaction strengths of the charged fragment pair and the neutral one can be challenging using IFIEs that include the ES components. Therefore, herein, the study focused on the amino acid residues contributing for attractive interactions in the CT and DI components caused by short-range interactions, such as hydrogen bonds and XH/ π interactions, to reveal the molecular recognition mechanism on the interface between the SARS-CoV-2 S-protein and ACE2.

Next, the relationship between the energy intensity and the observed interaction was described based on the above the interaction analysis between the SARS-CoV-2 spike and ACE2. Note, a more detailed description of the interactions between the SARS-CoV-2 S-protein and ACE2 based on Fig. 5 was provided in Section S2.†

3.1.1 XH–Y hydrogen bonds. The heavy atom pairs detected as hydrogen bonds in the geometric IFP and their FMO-based interaction energies are summarized in Table S1.† It was confirmed that the fragments showing attractive interaction energies of ES and CT+mix terms less than -30 and -3 kcal mol $^{-1}$ (Fig. 4B and D), respectively, were mainly associated with NH–O and OH–O hydrogen bonds with a distance between heavy atoms within 3 Å. The hydrogen bonds between the O oxygen atom of K353_{ACE2}, which was contained within the G354_{ACE2} fragment, and the hydrogen atom bonded to N nitrogen atom of G502_{Spike} (Fig. 5D), the OD1 oxygen atom of D38_{ACE2} and the hydrogen atom bonded to OH oxygen atom of Y449_{Spike} (Fig. 5G), the ND1 nitrogen atom of H34_{ACE2} and the hydrogen atom bonded to OH oxygen atom of Y453_{Spike} (Fig. 5H), and the OD1 oxygen atom of N487_{Spike} and the hydrogen atom bonded to NE2 nitrogen atom of Q24_{ACE2} (Fig. 5M) were typical examples such attractive interactions. The interaction energy between D30_{ACE2} and K417_{Spike} was the largest at -116.4 kcal mol $^{-1}$, where the distance between the heavy atoms of OD1 and OD2 oxygen atoms of D30_{ACE2} and NZ nitrogen atom of K417_{Spike} (Fig. 5J) were 2.9 and 3.8 Å, respectively. The atom pairs were also the only salt bridge pair between the S-protein and ACE2 that was detected from the geometric IFP. It was confirmed that these amino acid residues forming hydrogen bonds had attractive interaction energy lower than -5 kcal mol $^{-1}$, even in the ES component.

On the other hand, although weaker than the interaction energies of the NH–O and OH–O hydrogen bonds, CH–O hydrogen bonds also showed an attractive interaction energy that contributed to the molecular recognition of the SARS-CoV-2 S-protein and ACE2, such as the oxygen and carbon atom pairs for the O oxygen atom of T27_{ACE2}, which was contained within the F28_{ACE2} fragment, and the hydrogen atom bonded to CZ carbon atom of F456_{Spike} (Fig. 5K), as well as the OD1 oxygen



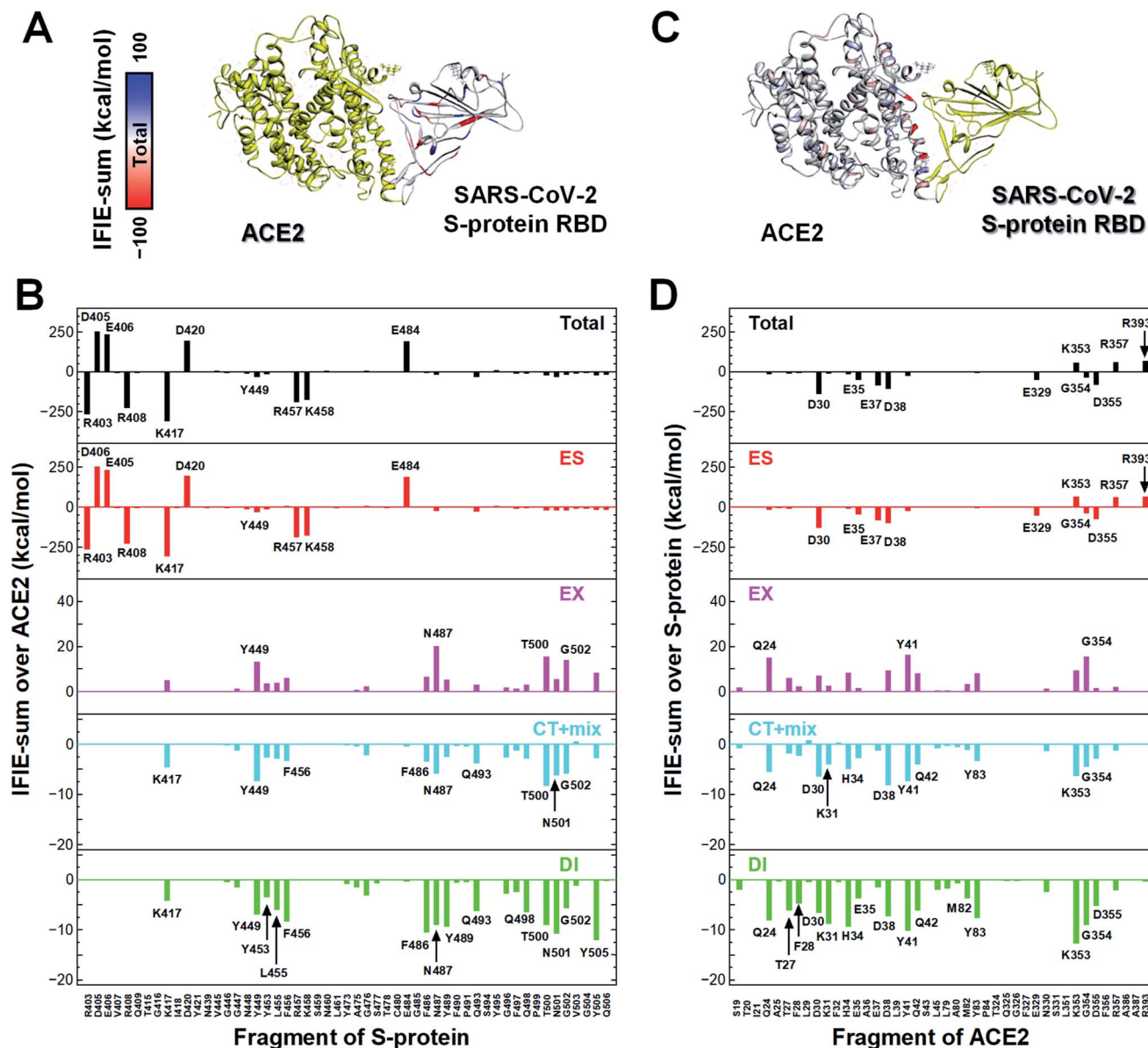


Fig. 4 Intermolecular interaction energies between the SARS-CoV-2 S-protein and ACE2 in host cells. The IFIE-sums over (A and B) ACE2 (S19–A614) and (C and D) S-protein (C361–E516) with each amino acid residue near the binding surface of the S-protein and ACE2 are shown. Visualization of IFIE-sums over the fragments of ACE2 (A) and SARS-CoV-2 S-protein (C) with attractive and repulsive interactions are represented by red and blue, respectively. Sugar chains are depicted by the ball and stick model. Figures (B) and (D) illustrate the IFIE-sums over ACE2 and SARS-CoV-2 S-protein, respectively, total, electrostatic (ES), exchange–repulsion (EX), charge transfer with higher-order mixed terms (CT+mix), and dispersion interaction (DI) components at the amino acid residue. The residue names of the corresponding fragments with interaction energies of ES, CT+mix, and DI components lower than -30 , -3 , and -3 kcal mol $^{-1}$, respectively, are shown.

atom of N487_{Spike} with the hydrogen atom bonded to CG carbon atom of Q24_{ACE2} (Fig. 5M).

3.1.2 XH/ π interactions. Besides the π -orbital of the aromatic ring, the heavy atom pairs detected as XH/ π in the geometric IFP and their FMO-based interaction energies are summarized in Table S2.† It was confirmed that the fragments showing attractive interaction energy of DI component lower than -3 kcal mol $^{-1}$ (Fig. 4B and D) were mainly associated with XH/ π interactions with a distance between the centroid of the aromatic ring and heavy atom of X within 4 Å. For example, there were CH/ π interactions *via* π and σ_{CH} orbitals on the

imidazole ring of H34_{ACE2} and the hydrogen atoms bonded to CD1 and CD2 carbon atoms of L455_{Spike} (Fig. 5H), the phenol ring of Y83_{ACE2} and the hydrogen atoms bonded to CE1 and CZ carbon atoms of F486_{Spike} (Fig. 5N), and the benzene ring of F486_{Spike} and the hydrogen atoms bonded to CB carbon atom of M82_{ACE2} (Fig. 5N). Moreover, it was clarified that not only the CH of the hydrophobic amino acid residues but also CH/ π interaction between the phenol ring of Y505_{Spike} and the hydrogen atom bonded to CA carbon atom of K353_{ACE2} (Fig. 5F) contributed to the SARS-CoV-2 and ACE2 binding.

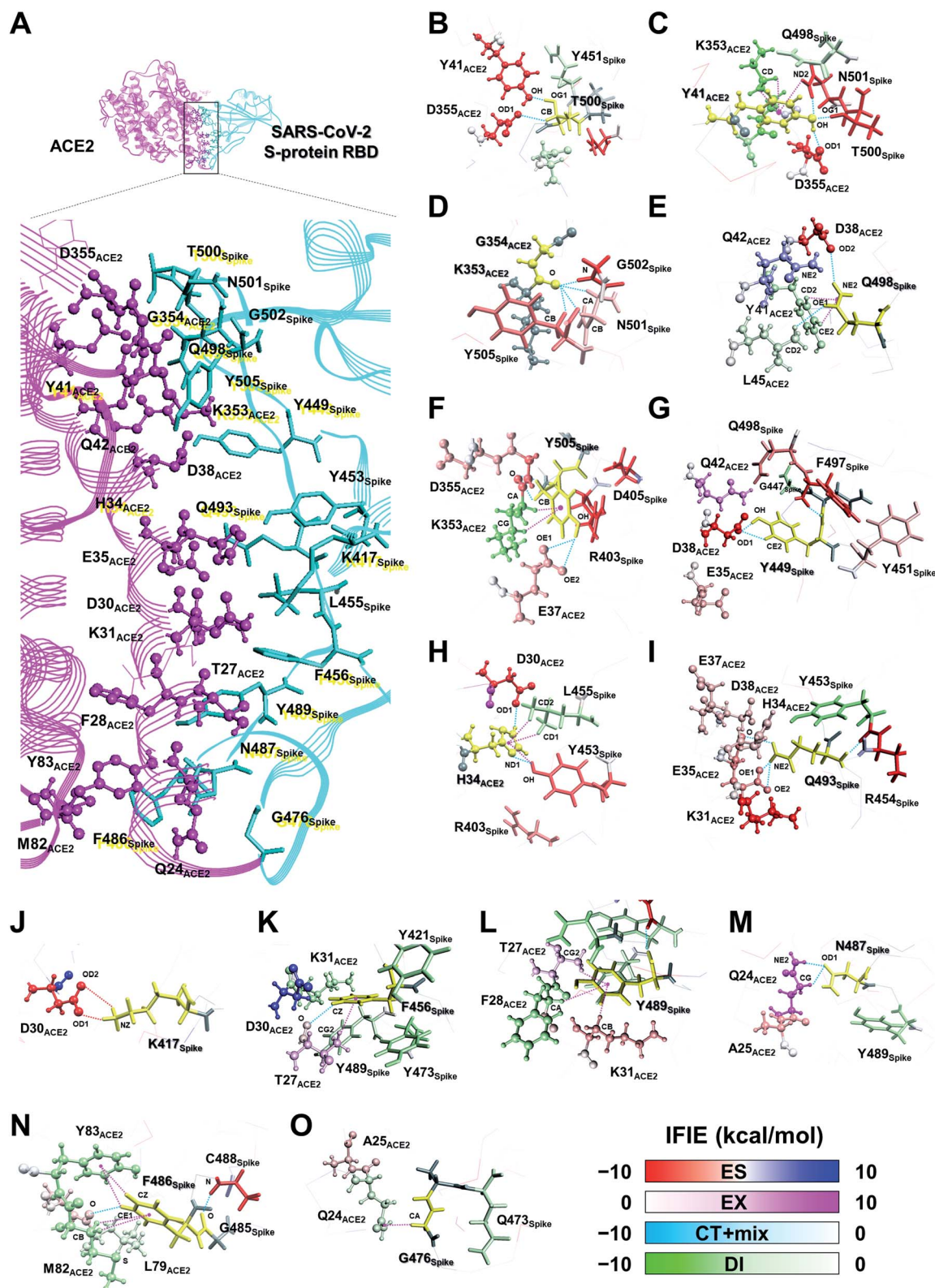


Fig. 5 3D visualization of the interaction energies between the SARS-CoV-2 S-protein amino acid residues near the contact surface with ACE2 (A). The fragments of T500_{Spike} (B), Y41_{ACE2} (C), G354_{ACE2} (D), Q498_{Spike} (E), Y505_{Spike} (F), Y449_{Spike} (G), H34_{ACE2} (H), Q493_{Spike} (I), K417_{Spike} (J), F456_{Spike} (K), Y489_{Spike} (L), N487_{Spike} (M), F486_{Spike} (N), and G476_{Spike} (O) are shown in yellow. The main components of the attractive and repulsive interaction energies are represented by the following color scheme: ES component, red and blue; DI component, green and white. The SARS-CoV-2 S-protein and ACE2 are shown using stick and ball model and stick model, respectively. Hydrogen bonds, ion pair interactions, and π/π interactions are shown by the cyan, red, and magenta dotted lines.



In addition to the CH/ π interactions described, OH/ π interaction between the imidazole ring of H34_{ACE2} and the hydrogen atom bonded to OH oxygen atom of Y453_{Spike} (Fig. 5H), and the NH/ π interaction between the phenol ring of Y41_{ACE2} and the hydrogen atom bonded to ND2 nitrogen atom of N501_{Spike} (Fig. 5C), were also found. It was confirmed that interaction energies of DI component for H34_{ACE2} with Y453_{Spike}, and Y41_{ACE2} with N501_{Spike} had stable interaction energies of -3.4 and -4.6 kcal mol⁻¹, respectively.

3.1.3 Ion pair, cation/ π , and π - π interactions. Based on PIEDA and geometric IFP analyses, ion pair interaction was detected only on the NZ nitrogen atom of the side chain on K417_{Spike} and between OD1 and OD2 oxygen atoms of the side chain on E30_{ACE2}. In addition, there were no cation/ π and π - π interactions between the S-protein and ACE2.

Taken together, the FMO data revealed that interaction networks were formed between the SARS-CoV-2 S-protein and ACE2 *via* hydrogen bond, XH/ π , and salt bridge interactions spanning multiple residues. The 15 residues of ACE2 selected by CT and DI energy analyses play key roles in the recognition of the SARS-CoV-2 S-protein, representing hot spot residues for inhibiting the binding to ACE2 in the context of drug and antibody design. Although the XH-Y hydrogen bond and XH/ π interactions were detected based on the geometric IFP analysis, this approach may fail to detect some molecular interactions. In particular, PIEDA can detect various types of XH/ π interactions and also evaluate them quantitatively, whereas structure-based analysis detects only typical interactions. In fact, the CH/ π interaction between the phenol ring of Y489_{Spike} and the hydrogen atom bonded to CA carbon atom of F28_{ACE2} (Fig. 5L), which was not detected by geometric IFP analysis, was found by interaction energy of DI component with PIEDA. By the geometric IFP, the XH/ π interactions with the π -orbital of the amide were not detected because this analysis method targeted the π -orbital of the aromatic ring. By PIEDA; however, it was confirmed that the XH/ π interactions with the π -orbital of the amide in the side chain on Q498_{Spike} and Q24_{ACE2} were also crucial for molecular recognition between the SARS-CoV-2 S-protein and ACE2 (Fig. 5E and O).

The amino acid residue pairs that formed the NH-O and OH-O hydrogen bonds were in good agreement with similar interactions reported in previous studies.^{10-12,79} Herein, it was the first report describing that the formation of CH-O hydrogen bonds with weak interaction energy are important for molecular recognition of ACE2 by the SARS-CoV-2. In addition, although it was described in reports of X-ray crystallography and electron microscopic analyses that hydrophobic residues are involved in molecular recognition as an effect of van der Waals interaction with ambiguous contributions, the findings here reported provide further clarification that such hydrophobic residues form XH/ π interactions.

3.2. What are the differences and similarities in the ACE2 recognition mechanism among three SARS S-proteins?

To clarify the differences in the molecular recognition of ACE2 between the SARS-CoV-2 (Fig. 4), the SARS-CoV-2 chimeric

(Fig. S2†), and the SARS-CoV S-proteins (Fig. S3†), the FMO-based interaction energies of all three structures were analyzed and compared. Furthermore, the difference in the binding ability of ACE2 (S19-A614) between the SARS-CoV-2 (C348-E502), the SARS-CoV-2 chimeric (C348-E502), and the SARS-CoV S-proteins was also investigated by QM-based interaction energy analysis (Fig. 6 and 7). To easy understanding, Fig. S5† shows the hot spots on molecular surface of the ACE2 and the SARS-CoV-2 S-protein complex (PDB ID: 6LZG) based on our research. This approach revealed hot spot residues of ACE2 for designing low, medium, and peptide inhibitors, as well as epitope candidates of S-protein, to prevent S-protein and ACE2 binding.

3.2.1 Hot spot analysis of ACE2. The three types of S-proteins were compared with each other for their ability to interact with ACE2, and key amino acid residues that were candidates for hot spot for inhibiting S-protein binding were examined. Fig. 6 shows a heat map of ES, EX, CT, and DI interaction energy components for each amino acid residue in ACE2 on the binding surface between ACE2 and the S-proteins. As a result, 22 amino acid residues of ACE2 were found to be important for binding to the three S-proteins, among which 15 were common amino acid residues (Q24, T27, D30, K31, H34, E35, E37, D38, Y41, Q42, Y83, E329, K353, G354, and D35) on ACE2 that could be considered essential amino acid residues for recognition of the S-protein. It was also confirmed that the SARS-CoV-2 and the SARS-CoV-2 chimeric S-proteins formed several short-range interactions with the amino acid residues of ACE2 compared with the SARS-CoV S-protein. The amino acid residues in which the CT and DI interactions were more strongly bound than -3 kcal mol⁻¹ could form short-range interactions, such as hydrogen bonds and XH/ π interactions. It is expected that the design of small/medium molecules targeting these 22 amino acid residues may lead to effective inhibition of S-protein binding.

3.2.2 Hot spot analysis of S-protein. Similar analyses of potential hot spots for ACE2 on the S-protein (Fig. 7) revealed 34, 37, and 34 amino acid residues on the SARS-CoV-2, the SARS-CoV-2 chimeric, and the SARS-CoV S-proteins, respectively, that attractively interacted with ACE2. The SARS-CoV-2 and the SARS-CoV-2 chimeric S-proteins had similar interaction results as the amino acid sequences of the receptor binding motif were almost the same (Fig. 2). In addition, 28 amino acid residues showed a common attractive interaction among the three S-proteins, being considered to be essential hot spots for the S-protein and ACE2 recognition regardless of the protein type. It was also found that, similar to the hot spot analysis of ACE2, the complex of the SARS-CoV-2 and the SARS-CoV-2 chimera had more amino acid residues that formed short-range interactions than the complex with the SARS-CoV. The 28 common hot spots including 17 conserved residues were highly homologous among the three SARS S-proteins and can be designated as essential hot spots for recognizing ACE2 in S-proteins (SARS-CoV-2, SARS-CoV, and MARS virus).

As shown in Fig. 6 and 7, the SARS-CoV-2 S-protein had more hot spots that interacted with ACE2 by CT and DI interaction than the SARS-CoV S-protein; in addition, the SARS-CoV-2 S-



IFIE-sum over S-protein with each amino acid residue of ACE2 (kcal/mol)																	^b Importance for hot spots of ACE2
ACE2	SARS-CoV-2					SARS-CoV-2 chimera					SARS-CoV						
Residue	ES	EX	CT	DI	^a Hot spot	ES	EX	CT	DI	^a Hot spot	ES	EX	CT	DI	^a Hot spot		
SER19	4.2	2.0	-0.7	-2.0		5.1	3.8	-1.0	-2.5		-0.3	0.0	-0.4	-0.7			
THR20	-2.1	0.0	0.0	-0.1		-2.5	0.0	0.0	0.0		-0.9	0.0	0.0	-0.1			
ILE21	-0.5	0.0	0.0	-0.1		-0.6	0.0	0.0	0.0		-2.0	0.0	0.0	0.0			
GLN24	-16.5	15.0	-5.5	-8.2	+	-11.0	4.6	-3.1	-6.0	+	-5.9	3.1	-1.2	-3.1	+	+++	
ALA25	-5.5	0.0	-0.1	-0.4	+	-1.6	0.0	-0.2	-0.6		-1.9	0.0	0.0	-0.3		+	
THR27	-10.2	6.2	-1.8	-6.2	+	-4.0	0.7	-1.9	-3.6	+	-2.9	2.7	-1.0	-4.2	+	+++	
PHE28	-3.3	2.4	-2.3	-4.8	+	-1.9	2.4	-2.2	-5.1	+	0.9	0.9	-1.1	-3.0		++	
LEU29	-1.3	0.1	0.8	-0.5		0.2	0.1	0.4	-0.5		-0.8	0.1	0.3	-0.5			
ASP30	-132.2	7.0	-6.5	-6.7	+	-30.2	0.7	-0.9	-1.9	+	-54.9	0.0	-0.1	-0.5	+	+++	
LYS31	-2.2	2.8	-4.0	-8.8	+	-34.6	9.9	-6.6	-11.3	+	30.0	3.4	-2.2	-5.9	+	+++	
PHE32	1.5	0.1	0.3	-0.5		0.9	0.1	0.6	-0.4		0.7	0.0	0.3	-0.4			
HIS34	-9.4	8.4	-5.0	-9.3	+	-15.8	9.0	-3.8	-8.0	+	-9.1	1.6	-2.1	-4.5	+	+++	
GLU35	-48.0	1.7	-2.7	-3.7	+	-33.3	6.5	-4.9	-4.7	+	-32.1	0.0	0.0	-0.2	+	+++	
ALA36	1.1	0.0	0.1	-0.1		-0.5	0.0	0.0	-0.1		0.4	0.0	0.0	0.0			
GLU37	-82.3	0.2	-1.2	-1.5	+	-62.9	1.4	-2.7	-2.9	+	-62.5	0.4	-1.7	-1.6	+	+++	
ASP38	-100.9	9.5	-8.2	-7.4	+	-55.8	5.0	-5.1	-5.2	+	-68.6	4.9	-7.2	-6.7	+	+++	
LEU39	2.1	0.0	0.0	-0.1		2.1	0.0	0.0	-0.1		0.0	0.0	0.0	-0.1			
TYR41	-24.8	16.2	-7.3	-10.2	+	-19.1	17.9	-6.8	-10.5	+	-14.2	12.8	-5.5	-10.5	+	+++	
GLN42	-2.3	8.0	-4.0	-6.2	+	-5.2	3.4	-2.5	-3.3	+	-10.1	4.8	-2.9	-4.9	+	+++	
SER43	1.8	0.0	0.0	0.0		0.7	0.0	0.0	0.0		0.8	0.0	0.0	-0.1			
LEU45	0.6	0.6	-0.8	-2.0		-1.3	0.6	-0.9	-1.9		0.3	1.1	-0.8	-2.5			
LEU79	-1.0	0.8	-0.3	-1.7		-2.3	3.0	-0.6	-2.4		-0.3	0.6	-0.2	-1.5			
ALA80	-3.5	0.1	-0.6	-0.7	+	-2.5	0.0	-0.1	-0.3		-0.6	0.0	0.0	-0.1		+	
MET82	1.4	3.5	-1.1	-3.7	+	1.0	3.1	-0.9	-3.7	+	-1.2	4.8	-0.8	-2.9		++	
TYR83	-5.0	8.1	-3.4	-7.7	+	-12.7	8.3	-4.7	-8.4	+	-4.9	5.0	-1.7	-3.9	+	+++	
PRO84	0.2	0.0	0.0	0.0		1.3	0.0	0.0	0.0		0.4	0.0	0.0	0.0			
THR324	0.7	0.0	0.0	-0.1		0.0	0.0	0.0	-0.1		0.2	0.0	0.0	-0.2			
GLN325	-0.5	0.0	0.0	-0.3		-2.8	0.0	0.0	-0.2		-4.5	0.4	-1.1	-2.0	+	+	
GLY326	2.9	0.0	0.0	-0.2		-0.7	0.0	0.0	-0.2		0.8	0.0	-0.1	-0.6			
PHE327	2.2	0.0	0.0	-0.1		2.3	0.0	0.0	-0.1		3.4	0.0	0.0	-0.1			
GLU329	-53.0	0.0	0.0	0.0	+	-80.4	0.0	-2.4	-1.8	+	-112.2	5.7	-6.3	-5.0	+	+++	
ASN330	-2.1	1.3	-1.4	-2.4		-1.6	1.5	-1.4	-2.5		-7.2	6.1	-2.4	-4.8	+	+	
SER331	0.7	0.0	0.0	0.0		2.8	0.0	0.0	0.0		2.5	0.0	0.0	0.0			
LEU351	1.1	0.0	0.0	0.0		1.5	0.0	0.0	-0.1		1.3	0.0	0.0	0.0			
LYS353	65.0	9.4	-6.4	-12.8	+	20.6	17.5	-6.0	-13.4	+	28.2	14.1	-6.8	-14.4	+	+++	
GLY354	-39.3	15.6	-4.5	-9.1	+	-37.0	13.2	-3.6	-8.8	+	-46.8	31.2	-7.7	-10.2	+	+++	
ASP355	-75.1	1.7	-2.9	-5.2	+	-57.2	1.5	-2.5	-4.9	+	-68.0	4.4	-2.5	-6.3	+	+++	
PHE356	2.6	0.0	-0.1	-0.2		3.1	0.0	-0.1	-0.2		3.7	0.0	-0.1	-0.4			
ARG357	62.8	2.2	-1.2	-2.1		48.2	1.4	-0.8	-1.8		52.3	1.5	-1.0	-1.6			
ALA386	-1.4	0.0	0.0	0.0		-0.8	0.0	0.0	0.0		-0.7	0.0	0.0	0.0			
ALA387	-3.3	0.0	0.0	-0.1	+	-1.3	0.0	0.0	-0.1		-1.2	0.0	0.0	0.0		+	
ARG393	67.2	0.0	-0.1	-0.4		42.6	0.1	-0.3	-0.6		50.8	0.0	-0.1	-0.2			
^c Count	16	0	10	15	20	14	0	9	14	17	14	0	5	13	17	22	

Fig. 6 The IFIE-sums over the S-protein of the SARS-CoV-2 (C348–E502), the SARS-CoV-2 chimera (C348–E502), and the SARS-CoV (C348–E502) with each amino acid residue of ACE2 by PIEDA. Attractive interaction energies of the electrostatic (ES), charge transfer with higher-order mixed terms (CT), and dispersion interaction (DI) are indicated using red, light blue, and green gradations, respectively, and repulsive interaction energy of the exchange–repulsion (EX) is indicated using magenta gradation. ^aHot spots with an interaction energy of -3 kcal mol^{-1} or less in any of the ES, EX, CT, and DI components, were labeled as "+". ^bThe number of "+" shows importance of hot spots between three complexes. ^cCount is number of hot spot residues.

protein is considered to bind more firmly to ACE2 than the SARS-CoV S-protein.

These results are consistent with previous findings^{2,17,28,80} in Section S5.† We also discovered 17 and 4 new hot spots of the SARS-CoV-2 S-protein and the ACE2, respectively (Fig. S5†), that were overlooked in other QM researches for the following reasons: Lim *et al.* reported the interaction analysis based on the FMO calculations using the self-consistent charge density-functional tight-binding method with the third-order expansion using semi-empirical dispersion (DFTB3/D) method.¹⁷ The DFTB3/D method with a partially semi-empirical approach tended to underestimate CT and DI components compared to the MP2 method with *ab initio* electron-correlated QM theory.

In addition, there were several amino acid residues for which the tendency of attractive and repulsive interactions of CT and DI components was different between DFTB3/D and MP2 methods.

Focusing on amino-acid-residue pairs forming NH–O and OH–O hydrogen bonds, Gómez *et al.* performed higher precision QM calculation with computed accurate interaction energies using highly correlated domain based local pair-natural orbital-coupled cluster (DLPNO-CCSD(T)/aug-cc-pVDZ) level to investigate accurate interaction energies for its excision model not considering the entire complex.¹⁸ On the other hand, we analyzed interactions for all the amino-acid-residue pairs of the entire system taking into account their



IFIE-sums over ACE2 with each amino acid of S-protein (kcal/mol)																	^b Importance for hot spots of three S-proteins	
SARS-CoV-2 S-protein						SARS-CoV-2 chimera S-protein						SARS-CoV S-protein						
Residue	ES	EX	CT	DI	^a Hot spot	Residue	ES	EX	CT	DI	^a Hot spot	Residue	ES	EX	CT	DI		^a Hot spot
ARG403	-265.3	0.0	0.0	0.0	+	LYS403	-263.9	0.0	0.0	-0.1	+	LYS390	-271.1	0.0	0.0	-0.1	+	+++
ASP405	254.8	0.0	0.0	0.0		ASP405	254.8	0.0	0.0	0.0		ASP392	261.7	0.0	0.0	0.0		
GLU406	234.6	0.0	0.0	0.0		ASP406	237.1	0.0	0.0	0.0		ASP393	248.1	0.0	0.0	0.0		
VAL407	-6.8	0.0	0.0	0.0	+	VAL407	-7.2	0.0	0.0	0.0	+	VAL394	-6.7	0.0	0.0	0.0	+	+++
ARG408	-228.1	0.0	0.0	0.0	+	ARG408	-231.2	0.0	0.0	0.0	+	ARG395	-242.5	0.0	0.0	0.0	+	+++
GLN409	-6.9	0.0	0.0	0.0	+	GLN409	-7.0	0.0	0.0	0.0	+	GLN396	-7.4	0.0	0.0	0.0	+	+++
THR415	-1.1	0.0	0.0	0.0		THR415	-3.5	0.0	0.0	0.0	+	THR402	-0.7	0.0	0.0	0.0		+
GLY416	3.6	0.0	0.0	0.0		GLY416	4.6	0.0	0.0	0.0		GLY403	-1.8	0.0	0.0	0.0		
LYS417	-308.6	5.1	-4.5	-4.2	+	VAL417	-4.5	0.0	0.0	-0.2	+	VAL404	-4.9	0.0	0.0	-0.1	+	+++
ILE418	-6.1	0.0	0.0	0.0	+	ILE418	-7.2	0.0	0.0	0.0	+	ILE405	-7.5	0.0	0.0	0.0	+	+++
ASP420	197.5	0.0	0.0	0.0		ASP420	196.5	0.0	0.0	0.0		ASP407	210.0	0.0	0.0	0.0		
TYR421	-1.5	0.0	0.0	0.0		TYR421	-3.9	0.0	0.0	0.0	+	TYR408	-9.9	0.0	0.0	0.0	+	++
ASN439	-5.0	0.0	0.0	0.0	+	ARG439	-296.8	0.0	-2.4	-1.7	+	ARG426	-333.2	5.7	-6.6	-5.5	+	+++
VAL445	6.2	0.0	0.0	-0.1		SER445	5.9	0.0	0.0	0.0		SER432	3.2	0.0	0.0	0.0		
GLY446	-7.8	0.0	-0.2	-0.5	+	THR446	2.8	0.0	0.0	-0.1		THR433	7.1	0.0	0.0	-0.2		+
GLY447	-2.4	1.4	-1.3	-1.5		GLY447	5.6	0.0	0.0	0.0		GLY434	1.4	0.0	0.0	-0.1		
ASN448	-11.8	0.0	0.0	-0.1	+	ASN448	-7.3	0.0	0.0	0.0	+	ASN435	-10.6	0.0	0.0	0.0	+	+++
TYR449	-31.8	13.1	-7.4	-6.9	+	TYR449	-23.3	5.0	-4.8	-5.4	+	TYR436	-33.3	8.3	-7.5	-7.6	+	+++
TYR453	-15.0	3.7	-2.6	-3.5	+	TYR453	-9.9	4.8	-2.4	-3.5	+	TYR440	-7.7	1.0	-1.3	-2.0	+	+++
LEU455	-0.5	3.9	-2.9	-6.0	+	LEU455	1.7	3.3	-1.5	-5.1	+	TYR442	1.2	0.9	-1.4	-4.1	+	+++
PHE456	7.1	6.1	-3.4	-8.4	+	PHE456	2.0	4.5	-2.8	-7.0	+	LEU443	5.5	1.1	-0.3	-1.6		++
ARG457	-188.6	0.0	0.0	0.0	+	ARG457	-185.4	0.0	0.0	0.0	+	ARG444	-194.7	0.0	0.0	0.0	+	+++
LYS458	-177.2	0.0	0.0	0.0	+	LYS458	-174.8	0.0	0.0	0.0	+	HIS445	4.0	0.0	0.0	0.0		++
SER459	-0.3	0.0	0.0	0.0		SER459	-7.7	0.0	0.0	0.0	+	GLY446	-7.0	0.0	0.0	0.0	+	+
ASN460	5.1	0.0	0.0	0.0		ASN460	4.9	0.0	0.0	0.0		LYS447	-205.0	0.0	0.0	0.0	+	+
LEU461	-5.2	0.0	0.0	0.0	+	LEU461	-4.6	0.0	0.0	0.0	+	LEU448	-5.2	0.0	0.0	0.0	+	+++
TYR473	-4.8	0.4	-0.2	-0.8	+	TYR473	-4.8	0.0	-0.2	-0.5	+	PHE460	-5.3	0.1	-0.1	-0.4	+	+++
ALA475	0.2	1.0	-0.4	-1.5		ALA475	-0.4	0.2	-0.7	-1.3		PRO462	-5.5	0.1	-0.3	-1.3	+	++
GLY476	7.4	2.5	-2.1	-3.2	+	GLY476	2.9	4.8	-2.2	-3.6	+	ASP463	203.9	0.0	-0.4	-0.6		++
SER477	-3.8	0.0	0.1	-0.8	+	SER477	-4.4	0.0	0.5	-0.8	+	GLY464	-5.2	0.0	0.0	0.0	+	+++
THR478	-5.6	0.0	0.0	0.0	+	THR478	-4.2	0.0	0.0	0.0	+	LYS465	-203.4	0.0	0.0	0.0	+	+++
CYS480	-2.7	0.0	0.0	-0.2		CYS480	-5.9	0.0	0.0	-0.1	+	CYS467	-2.5	0.0	0.0	-0.1		+
GLU484	191.0	0.0	-0.4	-0.4		GLU484	215.8	0.0	0.0	-0.1		PRO470	-7.0	0.0	0.0	0.0	+	+
GLY485	-0.4	0.0	0.0	0.0		GLY485	1.1	0.0	0.0	0.0		ALA471	5.7	0.0	0.0	-0.1		
PHE486	-0.4	6.6	-3.5	-10.6	+	PHE486	-3.9	9.3	-3.1	-10.5	+	LEU472	0.1	5.4	-1.0	-4.3	+	+++
ASN487	-26.4	20.1	-5.9	-9.1	+	ASN487	-20.3	8.1	-4.9	-7.2	+	ASN473	-7.7	7.8	-2.1	-5.5	+	+++
TYR489	0.9	5.3	-2.5	-9.4	+	TYR489	-1.7	3.8	-3.3	-9.5	+	TYR475	2.2	5.3	-2.7	-9.5	+	+++
PHE490	1.7	0.0	-0.3	-0.6		PHE490	-1.9	0.0	-0.1	-0.3		TRP476	-10.8	0.0	0.0	0.0	+	+
PRO491	-1.0	0.0	-0.4	-0.5		PRO491	-2.9	0.0	-0.6	-0.6		PRO477	7.0	0.0	0.0	0.0		
GLN493	-27.5	3.1	-3.7	-6.3	+	GLN493	-54.2	13.2	-8.6	-9.1	+	ASN479	-0.2	0.6	-0.6	-2.1		++
SER494	-4.0	0.0	0.0	0.0	+	SER494	-3.8	0.0	0.0	0.0	+	ASP480	229.8	0.0	0.0	0.0		++
TYR495	8.4	0.0	0.0	-0.1		TYR495	5.1	0.0	0.0	0.0		TYR481	5.2	0.0	0.0	0.0		
GLY496	0.9	2.0	-2.6	-2.8		GLY496	-0.8	5.4	-2.5	-1.7		GLY482	3.1	0.1	-0.7	-1.2		
PHE497	-9.5	1.3	-1.3	-2.4	+	PHE497	-14.2	5.1	-0.3	-3.1	+	PHE483	-7.4	1.2	-1.1	-1.9	+	+++
GLN498	-4.9	3.2	-2.8	-6.5	+	GLN498	-13.0	4.9	-3.5	-6.4	+	TYR484	-14.2	5.7	-4.4	-9.7	+	+++
PRO499	1.4	0.0	0.0	-0.1		PRO499	3.8	0.0	0.0	-0.1		THR485	3.4	0.0	-0.1	-0.2		
THR500	-19.9	15.6	-8.3	-9.1	+	THR500	-12.1	16.6	-7.8	-9.4	+	THR486	-13.5	11.4	-7.2	-9.0	+	+++
ASN501	-20.4	5.6	-6.2	-10.8	+	ASN501	-8.5	5.3	-4.8	-10.1	+	THR487	-8.3	17.0	-7.6	-13.6	+	+++
GLY502	-21.0	13.9	-5.9	-5.7	+	GLY502	-19.4	11.8	-5.7	-5.7	+	GLY488	-28.7	29.4	-9.2	-7.9	+	+++
VAL503	-10.5	0.1	0.6	-1.2	+	VAL503	-10.1	0.1	0.7	-1.3	+	ILE489	-10.3	0.6	1.3	-3.0	+	+++
GLY504	-9.4	0.0	0.0	0.0	+	GLY504	-8.3	0.0	0.0	0.0	+	GLY490	-8.3	0.0	0.0	0.0	+	+++
TYR505	-15.4	8.4	-2.7	-12.1	+	TYR505	-20.9	9.3	-3.9	-13.5	+	TYR491	-15.9	8.0	-3.1	-11.8	+	+++
GLN506	-18.4	0.0	0.0	-0.2	+	GLN506	-15.1	0.0	0.0	-0.2	+	GLN492	-13.1	0.0	-0.1	-0.5	+	+++
^c Count	29	0	9	15	34	^c Count	33	0	10	15	37	^c Count	31	0	7	12	34	42

Fig. 7 IFIE-sums over ACE2 (S19–A614) with each amino acid residue of the SARS-CoV-2, the SARS-CoV-2 chimera, and the SARS-CoV S-proteins by PIEDA. Attractive interaction energies of the electrostatic (ES), charge transfer with higher-order mixed terms (CT), and dispersion interaction (DI) are indicated using red, light blue, and green gradations, respectively, and repulsive interaction energy of the exchange–repulsion (EX) is indicated using magenta gradation. ^aHot spots with an interaction energy of -3 kcal mol⁻¹ or less in any of the ES, EX, CT, and DI components, were labeled as “+”. ^bThe number of “+” shows importance of hot spots between three complexes. ^cCount is number of hot spot residues.

polarization caused by the surrounding environment. We focused not only on typical hydrogen bonds above but also on the CH–O hydrogen bonds and the XH/ π interactions to address a wide range of possibilities of the key residues of molecular recognition. Therefore, we succeeded in finding known and new hot spots that were structurally and quantum chemically valid.

3.3. Is FMO-based epitope analysis of the SARS-CoV-2 S-protein useful?

Based on the complex between the SARS-CoV-2 S-protein and ACE2, the expected epitope candidates analyzed by IFIE-sum

over B38 Fab antibody using PIEDA were further explored as potential neutralizing epitopes of S-protein. Using a complex of the SARS-CoV-2 S-protein (C361–E516) and B38 Fab antibody (D0–S217 on heavy chain and D0–C215 on light chain), PIEDA and geometric IFP analyses were performed similarly as described in the Sections 3.1 and 3.2. The actual epitope residues of the B38 Fab antibody on the SARS-CoV-2 S-protein were revealed and compared to the candidate epitope residues upon ACE2 binding.

First, FMO-based interaction energy and geometric IFP analyses of the S-protein and B38 Fab antibody were performed to clarify the amino acid residues and their interaction type that were critical for molecular recognition. FMO-based



other hand, the amino acid residues of S-protein in which attractive interaction was observed with ACE2 but not with the B38 Fab antibody were V407, R408, I418, N439, G446, N448, Y449, Y453, R457, T478, Q493, F497, and T500. It was also revealed new amino acid residues through which the S-protein interacted with the B38 Fab antibody (D405, E406, T415, G416, D420, Y421, N460, Y473, A475, E484, and G496). Several amino acid residues of S-protein that had an attractive interaction with ACE2 did not interact with the B38 Fab antibody. Nevertheless, the B38 Fab antibody showed more CT and DI interactions with the S-protein amino acid residues that did not interact with ACE2.

The numbers of amino acid residues in the SARS-CoV-2 S-protein showing CT and DI interaction were 9 and 15, respectively, when complexed with ACE2; whereas it was 14 and 20, respectively, when the S-protein was complexed with the B38 Fab antibody. This indicated that B38 Fab acquired more hydrogen bonds and XH/ π interactions than ACE2. From these results, the 21 amino acid residues of the SARS-CoV-2 S-protein shared by the ACE2 and B38 Fab antibody interactions would be essential epitopes that directly inhibit ACE2 recognition and binding. This data (Fig. 8) is expected to provide useful information for the development of potentially therapeutic antibodies. For example, for residues without attractive interaction with the antibody, the substitution of amino acid residues that mimic the interaction with ACE2 will lead to the design of antibodies with higher binding capacity. These amino acid residues were also consistent with the description of NH-O and OH-O hydrogen bonds reported by a previous study.³ New NH-O and OH-O hydrogen bonds, not shown by the previous study,³ were identified and their interaction was clarified by FMO calculations. This study demonstrated that CH-O hydrogen bonds and XH/ π interactions are crucial for the molecular recognition between the SARS-CoV-2 S-protein and B38 Fab antibody, with 21 epitope residues of the SARS-CoV-2 S-protein being critical for antibody design.

The difference in the interaction energy of each amino acid residue of the S-protein between ACE2 and B38 Fab antibody will provide worthwhile information for improving B38 Fab and other antibodies. Lim *et al.* has shown two large hot spot regions from FMO-based interaction analysis of ACE2 and S-proteins of the SARS-CoV-2.¹⁷ Since the hot spots of S-protein with the several neutralizing antibodies of the SARS-CoV coincided with the 2nd hot spot region, the authors speculated that the 2nd hot spot region was crucial for drug design against the SARS-CoV-2. However, our results suggest that all key hot spots for molecular recognition between the SARS-CoV-2 S-protein and ACE2 may play a significant role in the design of neutralizing antibodies. This is because the first and the second sizable hot spot regions of S-protein, defined by Lim *et al.*, were both found as epitopes of the complex between S-protein and B38 Fab antibody.

As Fig. 8 shows, in ACE2 binding, we confirmed that the primary contribution of local hot spots was long-distance electrostatic interaction, such as that of R403 and K417; however, in B38 Fab antibody binding, the local hot spots (*e.g.*, R403 and K417) and their surrounding amino acid residues acquired

IFIE-sums over ACE2 with each amino acid of S-protein (kcal/mol)						IFIE-sums over B38 Fab antibody with each amino acid of S-protein (kcal/mol)						^b Importance for hot spots and epitopes
SARS-CoV-2 S-protein						SARS-CoV-2 S-protein						
Residue	ES	EX	CT	DI	^a Hot spot	Residue	ES	EX	CT	DI	^a Epitope	
ARG403	-265.3	0.0	0.0	0.0	+	ARG403	-24.0	12.7	-7.2	-9.0	+	++
ASP405	254.8	0.0	0.0	0.0		ASP405	11.8	2.0	-1.8	-3.6	+	+
GLU406	234.6	0.0	0.0	0.0		GLU406	-13.8	0.1	-0.9	-1.8	+	+
VAL407	-6.8	0.0	0.0	0.0	+	VAL407	-3.8	0.0	0.0	0.0	+	++
ARG408	-228.1	0.0	0.0	0.0	+	ARG408	14.6	0.0	0.0	-0.2		+
GLN409	-6.9	0.0	0.0	0.0	+	GLN409	-8.8	0.2	-0.7	-0.9	+	++
THR415	-1.1	0.0	0.0	0.0		THR415	-16.0	13.0	-6.0	-6.2	+	+
GLY416	3.6	0.0	0.0	0.0		GLY416	6.1	4.1	-2.1	-3.8	+	+
LYS417	-308.6	5.1	-4.5	-4.2	+	LYS417	-26.7	7.4	-2.8	-11.1	+	++
ILE418	-6.1	0.0	0.0	0.0	+	ILE418	-3.8	0.0	0.1	-0.2	+	++
ASP420	197.5	0.0	0.0	0.0		ASP420	-100.1	34.7	-14.3	-5.9	+	+
TYR421	-1.5	0.0	0.0	0.0		TYR421	-16.7	14.2	-6.3	-9.9	+	+
ASN439	-5.0	0.0	0.0	0.0	+	ASN439	-2.1	0.0	0.0	0.0		+
VAL445	6.2	0.0	0.0	-0.1		VAL445	-0.2	0.0	0.0	0.0		
GLY446	-7.8	0.0	-0.2	-0.5	+	GLY446	-0.2	0.0	0.0	0.0		+
GLY447	-2.4	1.4	-1.3	-1.5		GLY447	-0.4	0.0	0.0	0.0		
ASN448	-11.8	0.0	0.0	-0.1	+	ASN448	-1.0	0.0	0.0	0.0		+
TYR449	-31.8	13.1	-7.4	-6.9	+	TYR449	-0.9	0.0	0.0	-0.1		+
TYR453	-15.0	3.7	-2.6	-3.5	+	TYR453	-0.1	0.2	-0.7	-1.4		+
LEU455	-0.5	3.9	-2.9	-6.0	+	LEU455	0.8	1.2	-1.8	-3.5	+	++
PHE456	7.1	6.1	-3.4	-8.4	+	PHE456	-27.7	18.3	-8.4	-9.1	+	++
ARG457	-188.6	0.0	0.0	0.0	+	ARG457	44.1	0.0	-0.7	-0.6		+
LYS458	-177.2	0.0	0.0	0.0	+	LYS458	-10.9	39.6	-14.6	-12.9	+	++
SER459	-0.3	0.0	0.0	0.0		SER459	-0.7	1.3	-1.5	-2.4		
ASN460	5.1	0.0	0.0	0.0		ASN460	-11.6	1.7	-1.4	-2.7	+	+
LEU461	-5.2	0.0	0.0	0.0	+	LEU461	1.4	0.0	0.0	0.0		+
TYR473	-4.8	0.4	-0.2	-0.8	+	TYR473	-26.3	20.3	-7.7	-6.3	+	++
ALA475	0.2	1.0	-0.4	-1.5		ALA475	-5.3	2.8	-2.3	-3.8	+	+
GLY476	7.4	2.5	-2.1	-3.2	+	GLY476	-36.8	11.9	-8.9	-8.9	+	++
SER477	-3.8	0.0	0.1	-0.8	+	SER477	5.7	0.5	2.3	-1.7		+
THR478	-5.6	0.0	0.0	0.0	+	THR478	4.7	0.0	0.0	-0.1		+
CYS480	-2.7	0.0	0.0	-0.2		CYS480	-0.8	0.0	-0.1	-0.1		
GLU484	191.0	0.0	-0.4	-0.4		GLU484	-28.6	0.0	0.0	0.0	+	+
GLY485	-0.4	0.0	0.0	0.0		GLY485	0.8	0.0	0.0	0.0		
PHE486	-0.4	6.6	-3.5	-10.6	+	PHE486	-7.5	5.7	-3.1	-8.0	+	++
ASN487	-26.4	20.1	-5.9	-9.1	+	ASN487	-39.4	15.7	-6.7	-8.2	+	++
TYR489	0.9	5.3	-2.5	-9.4	+	TYR489	-5.4	3.7	-3.6	-7.9	+	++
PHE490	1.7	0.0	-0.3	-0.6		PHE490	-3.0	0.0	-0.3	-0.5		
PRO491	-1.0	0.0	-0.4	-0.5		PRO491	-2.7	0.1	-0.9	-0.7		
GLN493	-27.5	3.1	-3.7	-6.3	+	GLN493	-3.7	0.4	-0.7	-1.4	+	++
SER494	-4.0	0.0	0.0	0.0	+	SER494	1.5	0.0	0.0	0.0		+
TYR495	8.4	0.0	0.0	-0.1		TYR495	-0.2	1.0	-1.2	-2.1		
GLY496	0.9	2.0	-2.6	-2.8		GLY496	-10.1	1.2	-1.7	-2.4	+	+
PHE497	-9.5	1.3	-1.3	-2.4	+	PHE497	3.2	0.2	-0.4	-0.9		+
GLN498	-4.9	3.2	-2.8	-6.5	+	GLN498	-15.2	15.5	-2.7	-7.8	+	++
PRO499	1.4	0.0	0.0	-0.1		PRO499	-0.3	0.0	0.0	-0.1		
THR500	-19.9	15.6	-8.3	-9.1	+	THR500	0.9	0.0	-0.2	-0.4		+
ASN501	-20.4	5.6	-6.2	-10.8	+	ASN501	-34.1	12.8	-8.5	-8.6	+	++
GLY502	-21.0	13.9	-5.9	-5.7	+	GLY502	-14.0	13.9	-6.1	-5.5	+	++
VAL503	-10.5	0.1	0.6	-1.2	+	VAL503	-6.4	0.1	0.4	-1.0	+	++
GLY504	-9.4	0.0	0.0	0.0	+	GLY504	-7.0	0.0	0.0	-0.1	+	++
TYR505	-15.4	8.4	-2.7	-12.1	+	TYR505	-23.8	13.0	-7.2	-17.8	+	++
GLN506	-18.4	0.0	0.0	-0.2	+	GLN506	-6.7	0.0	0.0	0.0	+	++
^c Count	29	0	9	15	34	^c Count	28	0	14	20	31	44

Fig. 8 Difference of the IFIE-sums over the SARS-CoV-2 S-protein (C361–E516) between ACE2 and B38 Fab antibody. Interaction of each amino acid residue on S-protein with ACE2 and B38 Fab antibody are listed. Attractive interaction energies of the electrostatic (ES), charge transfer with higher-order mixed terms (CT), and dispersion interaction (DI) are identified using red, light blue, and green gradations, respectively, and repulsive interaction energy of the exchange–repulsion (EX) is identified using magenta gradation. ^aHot spots and epitopes with an interaction energy of -3 kcal mol^{-1} or less in any of the ES, EX, CT, and DI components, were labeled as “+”. ^bThe number of “+” shows importance for hot spots and epitopes. ^cCount is number of hot spot and epitope residues.



3.4.2 Solvation effect. To account for the solvation effect, the statistically corrected IFIE (SCIFIE)⁸¹ and FMO method combined with molecular mechanics Poisson–Boltzmann surface area (FMO+MM-PBSA)⁴³ approaches were used for predicting the binding energy (Table 1). The SCIFIE approach is a method to consider electrostatic and solvation shielding effects. In this research, a statistically corrected ES interaction energy ($\Delta\tilde{E}_{SC-ES}^{II}$), which felt the shielding effect by both the

$$\Delta E_{\text{bind}} = \sum_{IJ} \Delta \tilde{E}_{IJ}^{\text{ES}} + \Delta \tilde{E}_{IJ}^{\text{EX}} + \Delta \tilde{E}_{IJ}^{\text{CT+mix}} + \Delta \tilde{E}_{IJ}^{\text{DI}} \quad (5)$$

S-protein	SARS-CoV-2	SARS-CoV-2 chimera	SARS-CoV	SARS-CoV-2
Binding protein	ACE2	ACE2	ACE2	B38 Fab antibody
K_d (nM)	44.2	23.2	185	—
pK_d	7.35	7.63	6.73	—
ES	−887.15	−645.09	−818.50	−432.38
EX	122.71	121.73	110.62	269.35
CT+mix	−70.93	−67.92	−56.88	−130.86
DI	−125.36	−123.48	−105.42	−179.50
Total	−960.73	−714.75	−870.19	−473.38
SC-ES	−612.88	−491.07	−549.43	−498.64
Total (SC)	−686.46	−560.74	−601.11	−539.65
Desolv	592.53	395.91	531.82	98.78
Total (Desolv)	−368.20	−318.84	−338.36	−374.61
Total (SC+Desolv)	−93.92	−164.83	−69.29	−440.87

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solvent and the inside of the molecule, was used to analyze the ES component. Then the modified IFIE with the SC-ES component is given by

$$\Delta\tilde{E}_{IJ}^{\text{SC}} = \Delta\tilde{E}_{IJ}^{\text{SC-ES}} + \Delta\tilde{E}_{IJ}^{\text{EX}} + \Delta\tilde{E}_{IJ}^{\text{CT+mix}} + \Delta\tilde{E}_{IJ}^{\text{DI}} \quad (6)$$

and the total binding energy using SCIFIE approach was evaluated by:

$$\Delta E_{\text{bind}} = \sum_{I,J} \Delta\tilde{E}_{IJ}^{\text{SC}} \quad (7)$$

where I is the fragment of the S-protein and J is the fragment of the ACE2/B38 Fab antibody. On the other hand, the FMO+MM-PBSA approach⁴³ is a method used to evaluate binding energy incorporating the solvation effect by combining the desolvation energy with the sum of IFIEs as follows:

$$\Delta G^{\text{Desolv}} = G_{AB}^{\text{Solv}} - G_A^{\text{Solv}} - G_B^{\text{Solv}} \quad (8)$$

where ΔG^{Desolv} is desolvation energy, and G^{Solv} is solvation energy with A representing the S-protein and B the ACE2/B38 Fab antibody. The total binding energy was predicted using FMO+MM-PBSA approach as follows:

$$\Delta E_{\text{bind}} = \sum_{I,J} \Delta\tilde{E}_{IJ} + \Delta G^{\text{Desolv}} \quad (9)$$

Lastly, the total binding energy was predicted using SCIFIE combined with the desolvation evaluated using MM-PBSA method as follows:

$$\Delta E_{\text{bind}} = \sum_{I,J} \Delta\tilde{E}_{IJ}^{\text{SC}} + \Delta G^{\text{Desolv}} \quad (10)$$

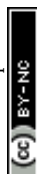
These total binding energies are shown in Table 1. While $\Delta\tilde{E}_{IJ}^{\text{SC-ES}}$ gives a weak contribution by shielding effect, the binding energy alone using the SCIFIE approach for the four complexes was not in agreement with the experimental binding affinity. By adding the desolvation energies from the MM-PBSA approach, the excessive electrostatic interaction energies were suppressed, and the predicted binding energies approximate to reproduce the experimental binding ability under the physiological condition. Moreover, the binding energy including both statistical correction and desolvation (SC+Desolv) for the three complexes, excluding the SARS-CoV-2 chimeric S-protein and ACE2 complex, seemed to be in agreement with the order of the binding affinity.^{2,80} The large difference seen in the interaction energy due to the electric charge of the molecules was eliminated by adding shielding and the desolvation effects, and the binding energies became comparable to that of the experimental value. Thus, when the PPI binding ability is estimated by FMO calculation among differently charged molecular systems, it may be useful to incorporate both the shielding and desolvation effects.

Taken together, it remains challenging to predict the strength of the binding ability of the differently charged proteins, such as ACE2, having highly negative charges, and of the B38 Fab antibody, having highly positive charges with a receptor such as S-protein using bare IFIE. However, incorporating desolvation energy and SCIFIE demonstrates that the predicted binding energies between differently charged

proteins could be improved for suppressing the overestimated ES interactions. Meanwhile, ACE2 comprises large amount of aspartic acids, glutamic acids, and histidines that can have multiple protonated states. Therefore, while the calculation was performed using only one protonation state in this study, it will be necessary to examine multiple plausible protonation states that may exist *in vivo* and investigate the predicted binding energy of PPI.

3.4.3 Sugar chain effect. The role of sugar chains in the molecular recognition of S-protein and ACE2/B38 Fab antibody was investigated. The sugar chain located on the SARS-CoV-2/SARS-CoV S-protein and ACE2 binding surface was the only BMA sugar chain, and was the terminal sugar chain of the NAG-NAG-BMA sugar chain extending from the side chain of N90 on ACE2 (Fig. 2B and C). In all complexes between the SARS-CoV-2 S-protein and ACE2 (PDB IDs: 6LZG, 6M0J, and 6M17) a sugar chain was not observed, in which a sugar chain consisting of three sugars from N90 of ACE2 was likely to reach and interact with the S-protein. There was one sugar chain (NAG) from N90 of ACE2, which did not directly interact with the SARS-CoV-2 S-protein (PDB ID: 6LZG). Moreover, no sugar chains were identified on the binding surface between the SARS-CoV-2 S-protein and B38 Fab antibody.

Table S5† lists the predicted binding energies using the IFIEs between the S-protein and ACE2/B38 Fab antibody without interaction energy of the sugar chain fragment. By comparing Tables 1 and S5,† the predicted binding energies of IFIEs between the SARS-CoV-2 chimeric S-protein and ACE2 was stabilized at $-15.5 \text{ kcal mol}^{-1}$ by the sugar chains, whereas that of the SARS-CoV S-protein and ACE2 was stabilized at $-2.8 \text{ kcal mol}^{-1}$. In the SARS-CoV-2 chimeric S-protein/ACE2 complex, the sugar chain on ACE2 attractively interacted with R408 *via* hydrogen bond and was critical for the molecular recognition of the S-protein. Moreover, the IFIEs between the BMA703 sugar chain on ACE2 and R408 of the SARS-CoV-2 S-protein showed an attractive interaction of $-16.2 \text{ kcal mol}^{-1}$ (ES: $-14.4 \text{ kcal mol}^{-1}$, EX: $1.5 \text{ kcal mol}^{-1}$, CT+mix: $-1.3 \text{ kcal mol}^{-1}$, DI: $-2.0 \text{ kcal mol}^{-1}$), which accounts for most of the summation of IFIEs in sugar chain effects. In the complex between the SARS-CoV S-protein and ACE2, weak attractive interaction between the BMA1092 sugar chain on ACE2 with the S-protein was confirmed, representing a repulsive interaction energy of $+3.0 \text{ kcal mol}^{-1}$; however, several residues (*e.g.*, D392, D393, and T402) of the S-protein interacted with the BMA1092 sugar chain with IFIEs of $-1.0 \text{ kcal mol}^{-1}$. In this study, only three sugar chains from N90 on ACE2 played key roles in the molecular recognition between the SARS-CoV-2 chimeric S-protein and ACE2. Since sugar chains are generally present on the molecular surface, the electron density tends to be difficult to see in the experimental setting. Thus, it will be necessary to discuss not only one X-ray crystal structure but also other X-ray crystal structures and structural fluctuations by MD simulations. The flexibility of sugar chains on the SARS-CoV-2 S-protein and the ACE2 was already investigated by several MD simulations.^{22–26} One of the results that the sugar chain on N90_{ACE2} works favorably for molecular recognition of the S-protein was reported as follows: the sugar chain on N90_{ACE2} was close enough



to the S-protein to repeatedly form interactions.²⁴ Although the sugar chains (NAG–NAG–BMA) of N90_{ACE2} were present near the interface with the S-protein, the sugar chain (BMA) of N90_{ACE2} and the S-protein did not always form hydrogen bonds judging from our FMO calculation results. In other words, it was suggested that the sugar chains of N90_{ACE2} would be located near the S-protein; however, they may not form a robust interaction with the S-protein. In fact, only one sugar chain (NAG) of N90_{ACE2}, which could not reach the interface with the S-protein, was observed in the complex of the SARS-CoV-2 S-protein and the ACE2, and it will be considered that the remaining sugar chains (NAG–BMA) of N90_{ACE2} would be fluctuating and may not be observed by X-ray crystal structures.

3.4.4 Mutation effect of the SARS-CoV-2 S-protein (N501Y). Currently, the highly transmissible variants of the United Kingdom (N501Y) and South Africa (N501Y, E484K, and K417N) are prevalent in various regions.^{6,7} Here, we investigated a mutation N501Y of RBD in common between S-protein variants of the United Kingdom and South Africa to understand the mutation effect by using modeling structure (FMODB ID: 7J11K), as seen in Section S7.† From interaction energy analysis (Fig. S7†), the N501Y mutation of the S-protein enhanced the attractive interaction because of the hydrogen bond and the XH/ π interactions with Y41_{ACE2} and K353_{ACE2} more than a wild type of the S-protein (Fig. S6†). It was confirmed that the N501Y mutation of the S-protein attractively enhances the DI energy of Y41_{ACE2} on the peptide motif (E37–Q42), which was essential for recognizing the S-protein, as proposed by Larue *et al.*²⁸ Along with these attractive interactions, it was confirmed that the binding energy including both statistical correction and desolvation (SC+Desolv) of the ACE2 with the mutant S-protein (N501Y) was further strengthened by *ca.* -10 kcal mol⁻¹ compared to that with the wild type S-protein as seen in Tables 1 and S6.† These FMO results may explain one of the reasons for the high infectivity of the mutant (N501Y). Recently, Zhu *et al.* analyzed the complex between the SARS-CoV-2 S-protein of the N501Y variant and a neutralizing antibody by cryo-electron microscopy (not yet published at February 7, 2021).⁸ They reported that the overall conformation and the interactions between the S-protein and the antibody of the variant almost were unchanged compared with those of the wild type; however, there was replacement of the asparagine residue by the bulkier tyrosine side chain results in subtle local rearrangements. Also, the binding ability of ACE2 with the variant was enhanced considerably compared with the wild type. Their results are in good agreement with our FMO results using the modeling structure of the complex between the S-protein of the N501Y variant and the ACE2. We hope that the experimental structure of the complex between the S-protein of the novel variant and ACE2/antibody is published for more detailed analysis.

4. Conclusion

The amino acid residues that are the key to the molecular recognition of three SARS S-proteins and ACE2/B38 Fab antibody were revealed by FMO-based interaction energy analysis with *ab initio* electron-correlated theory. The collected data

provided new insights in the importance of forming a complex interaction network for the molecular recognition between the S-protein and ACE2/B38 Fab antibody, not only *via* NH–O and OH–O hydrogen bonds and salt bridges, but also *via* the CH–O hydrogen bonds and XH/ π interactions. Since the XH/ π interaction is specifically found by the interaction energy of DI components, it is difficult to accurately understand XH/ π interaction solely through MM-based electrostatic interaction analysis and structure-based geometry analysis. Moreover, QM-based hot spot and epitope analyses by FMO calculations were useful in clarifying the type and strength of molecular interactions, such as hydrogen bond and XH/ π interactions. Prediction of the binding ability between the three types of SARS S-proteins and ACE2/antibody was performed by FMO-based interaction energy where incorporation of the shielding and desolvation effects was an essential factor. Since sugar chains are generally present on the molecular surface and have a disordered structure, the role of sugar chains in molecular recognition between the S-protein and ACE2 should be examined using structural fluctuation sampling, such as MD simulation, in future investigations. In addition, we investigated IFIE and binding energy between the ACE2 and the mutant N501Y on the SARS-CoV-2 S-protein regarding the mutation in common of the variants between the United Kingdom and South Africa, and our results can explain the high infectivity of the mutant.

We plan to release all the FMO data in this study on the public database, FMODB,^{75–77} so that all researchers can access it and utilize it for designing effective antibody-drugs. Also, our group has recently performed over 336 FMO calculations for the COVID-19-related proteins such as S-protein, main protease, RNA-dependent RNA polymerase,⁵⁶ based on the representative PDB structures selected in the PDB Japan database in line with this global fight effort against the coronavirus epidemic. These data have already been published on FMODB^{75–77} and can easily be analyzed IFIEs for inter- and intramolecular interactions on the Web interface. Finally, we expect that these findings of novel hot spots/epitopes between the SARS-CoV-2 S-protein and ACE2/B38 Fab antibody will provide useful information for future antibody design, evaluation of the binding property of variant, and small or medium drug design that overcome COVID-19.

Author contributions

The manuscript was written by CW and YO. The FMO calculations were performed and analyzed by CW. All authors discussed the results and have given approval to the final version of the manuscript.

Abbreviations

ACE2	Angiotensin-converting enzyme 2
BMA	β -D-mannopyranose
COVID-19	Coronavirus disease 2019
CT+mix	Charge transfer with higher-order mixed terms
Desolv	Desolvation



DFTB3/D	Self-consistent charge density-functional tight-binding method with the third-order expansion using semi-empirical dispersion
DI	Dispersion interaction
DLPNO-CCSD(T)	domain based local pair-natural orbital-coupled cluster
ES	Electrostatic
EX	Exchange–repulsion
Fab	Fragment antigen-binding
FMO	Fragment molecular orbital
FMODB	FMO database
FP	Fusion peptide
HR1	Heptad repeat 1
HR2	Heptad repeat 2
IC	Intracellular domain
IFIE	Inter-fragment interaction energy
IFIE-sum	Summation of IFIEs
IFP	Interaction fingerprint
Kd	Binding dissociation constant
MD	Molecular dynamics
MM	Molecular mechanics
MM-PBSA	Molecular mechanics Poisson–Boltzmann surface area
MO	Molecular orbital
MOE	Molecular Operating Environment
MP2	Second-order Møller–Plesset perturbation theory
NAG	N-acetyl- β -D-glucosamine
NTD	N-terminal domain
PDB	Protein data bank
PIEDA	Pair interaction energy decomposition analysis
PPI	Protein–protein interaction
QM	Quantum mechanics
RBD	Receptor-binding domain
RNA	Ribonucleic acid
SARS-CoV-2	Severe acute respiratory syndrome coronavirus 2
SC	Statistically correction
SC-ES	Statistically corrected electrostatic
SCIFIE	Statistically corrected IFIE
SD1	Subdomain 1
SD2	Subdomain 2
Solv	Solvation
S-protein	Spike glycoprotein
TM	Transmembrane region

Conflicts of interest

There are no conflicts to declare.

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