Conformational Variability Assessment of the Mutation Sites for D614G, B.1.1.7, and B.1.351 using SSSCPreds

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Analysis

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Abstract

Complementary techniques for the analysis of mutation sites at flexible regions, in which the position of atoms could not be determined by cryo-electron microscopy (Cryo-EM) such as the furin cleavage site of SARS-CoV-2, are necessary. The prediction data from SSSCPreds, a deep neural network-based prediction software of conformational flexibility or rigidity in proteins, can give insight into the conformational variability of mutation sites. We find that although the conformation of G614 is rigid, which is assigned as a left-handed (LH) α-helix-type one, that of D614 is flexible without the hydrogen bonding latch to T859. The rigidity of glycine which stabilizes the local conformation more effectively than that of aspartic acid with the latch, thereby contributes to the reduction of S1 shedding and increase in infectivity. Further it is predicted that no other amino acid allows the same conformation and stability as the glycine mutation in D614. The individual mutations in B.1.1.7 and B.1.351 have a lower effect and are not comparable to the overwhelming effectiveness of the D614G mutation. SSSCPreds provides important conformational flexibility insights into the deep neural network-based understanding of the current mutation sites and the potential for new ones in future.

Introduction

With the rapid expansion of the coronavirus disease 2019 (COVID-19) pandemic in 2020\textsuperscript{1}, a rate of 23.12 substitutions per year for SARS-CoV-2 is currently observed, and the evidence of possible reinfection with SARS-CoV-2 has been shown\textsuperscript{2}. The infection has spread through the process of natural selection so that predicting and tracking the impact of spontaneous mutations is necessary. Further, complementary techniques for the analysis of mutation sites at flexible regions, in which the position of atoms could not be determined by cryo-electron microscopy (Cryo-EM) such as the furin cleavage site of SARS-CoV-2\textsuperscript{3}, are needed.

Especially, the possibility of the next pandemic of Variant of Concern 202012/01 (VOC-202012/01, B.1.1.7) identified in UK\textsuperscript{4} and 501Y.v2 (B.1.351) having emerged in South Africa\textsuperscript{5} are deeply concerning. After March 2020, only two mutations, RNA-directed RNA polymerase P323L (ORF1ab P4715L, ORF1b P314L) and spike protein D614G have nearly overwhelmed the original mutation sites (https://nextstrain.org/ncov/global?c=gt-S_614). As for the receptor-binding motif (RBM) of spike protein, before B.1.1.7 and B.1.351, the mutation of S477N has expanded in Australia and in Europe but has not led to an extension of the pandemic (https://nextstrain.org/ncov/global?c=gt-S_477).

Recently, we reported a deep neural network-based prediction program of conformational flexibility or rigidity in proteins (SSSCPreds)\textsuperscript{6} using supersecondary structure code (SSSC)\textsuperscript{7–9}. The sequence flexibility/rigidity map of SARS-CoV-2 RBD (receptor binding domain), obtained from SSSCPreds, resembles the sequence-to-phenotype maps of ACE2-binding (angiotensin-converting enzyme 2-binding) affinity and expression, which was experimentally obtained by the deep mutational scanning\textsuperscript{10}. It suggests that the identical SSSC sequences among the ones predicted by three deep-neural-network-
based systems correlate well to the sequences with both lower ACE2-binding affinity and lower expression.

The frequency of mutations increases with the exponential increase in the number of infected people. The conformational flexibility of the protein sequences is deeply related to the ease of infection, and the accurate prediction is very important to make a countermeasure of COVID-19. In this paper, we report the conformational variability assessment of the mutation sites for D614G, and those of further strains B.1.1.7 and B.1.351.

**Results**

**D614G mutation**

As shown above, the D614G variant is now the dominant form worldwide. Recently, Gobeil and coworkers described that Cryo-EM structures reveal altered RBD disposition; antigenicity and proteolysis experiments reveal structural changes and enhanced furin cleavage efficiency of the G614 variant. However, the underlying factor of why glycine, and not other amino acids, can induce the effective strain replacement has not been explained.

The sequence flexibility/rigidity maps of all of the single amino acid mutations at the D614G mutation site using SSSCPreds indicate that only the mutation to glycine makes the other-type conformation ("T" conformation) rigid and reproduces the observed "T" conformations of Cryo-EM structures (Fig. 1a). On the other hand, although SSSCPred200 suggests the "T" conformation for D614, SSSCPred100 and SSSCPred predict the β-sheet-type conformations ("S" conformations). This means that the site of D614 is flexible without the hydrogen bonding latch between D614 and T859 (Fig. 1b,c).

Both observed "T" conformations of Cryo-EM structures for D614 and G614 (Protein Data Bank [PDB] ID code 6XR8 and 6XS6) are the same conformation11, which is assigned as a left-handed (LH) α-helix-type one (Fig. 1b,c)12. In general, the LH α-helix is stabilized by only glycine because glycine does not have chirality. The rigidity of glycine which stabilizes the local conformation more effectively than that of aspartic acid with the latch, thereby contributes to the reduction of S1 shedding13 and increase in infectivity without the latch between D614 and T859.

**B.1.1.7 and B.1.351**

As shown in Fig. 2 and Extended Data Fig. 1, the SSSCPreds data of the expanded S477N variant before B.1.1.7 and B.1.351 indicate that the S477N mutation increases the rigidity of the protein foundation GFNCYFPLQ. The foundation is located on the edge of flexible regions, in which the position of atoms could not be determined. The ratio of frequencies of the S477N mutation have gradually increased, but the mutation has not contributed to the pandemic. The SSSCPreds data of the N501Y mutation for B.1.1.7 show a similar increased stability of the foundation, but those of the E484K and N501Y mutations for B.1.351 suggest the equivalent rigidity of the original sequence (Fig. 2, Extended Data Fig. 1, and
Extended Data Fig. 5). The high ACE2-binding affinity of the single N501Y mutation has been reported\textsuperscript{10}. Although the K417N mutation site can also contact with ACE2, the SSSCPred data suggest flexibility of the nearby site (Extended Dataset 1 and Extended Data Fig. 4).

The mutations of HV69-70 deletion, Y144 deletion, and P681H for B.1.1.7 and D80A, D215G, and R246I for B.1.351 also correspond to the edge of flexible regions, in which the position of atoms could not be determined (Fig. 2, Extended Dataset 1, and Extended Data Fig. 2). The SSSCPred data of P681H mutation indicate the stabilization of the sequence CASYQT with identical β-sheet-type conformations before the furin cleavage site (Extended Data Fig. 3).

From the SSSCPred data, the mutations T716I and D1118H for B.1.1.7 (Fig. 2, Extended Dataset 1, and Extended Data Fig. 3) and the mutation A701V for B.1.351 (Extended Dataset 1 and Extended Data Fig. 5) increase the rigidity of the near sites. These mutation sites are located on the loop regions. The mutations T716I, S982A, and D1118H for B.1.1.7 and the mutation A701V for B.1.351 seem to contribute to the process that the transformation to post-fusion hairpin state in the membrane fusion proceeds smoothly.

**Discussion**

In this study, the conformational variability of the mutation sites for D614G, B.1.1.7, and B.1.351 has been evaluated by using SSSCPred. The overwhelming D614G mutation is rationalized by the more rigid conformation of glycine than that of aspartic acid, which is assigned as a LH α-helix-type one. In view of the conformational variability except the strength of binding affinity, individual mutations comparable to the overwhelming D614G mutation are not found in B.1.1.7 and B.1.351. As of September, the frequencies of N477 against S477 (https://nextstrain.org/ncov/global?c=gt-S_477) was similar to the present frequencies of Y501 against N501 (https://nextstrain.org/ncov/global?c=gt-S_501). B.1.1.7 has Q27stop of ORF8 (https://nextstrain.org/ncov/global?c=gt-ORF8_27). The mutation Q27stop of ORF8 seems to be correlated with milder disease\textsuperscript{14}. Another factor that leads to a new more virulent form is the P681H mutation at the furin cleavage site. The P681H mutation has appeared as of August (https://nextstrain.org/ncov/global?c=gt-S_681), but no report of a highly virulent form exists. The E484K mutation of B.1.351 (https://nextstrain.org/ncov/global?c=gt-S_484) has already appeared as of September, but the toxicity of B.1.351 seems to be equivalent to that of the original sequence. The S24L mutation of ORF8 is now going around in US (https://nextstrain.org/ncov/global?c=gt-ORF8_24) but are not overwhelming. It is suggested with the nextstrain data that the integrated effect of mutations contributes to the expansion of B.1.1.7 and B.1.351 but are not overwhelming.

**Declarations**

**ONLINE METHODS**
The FASTA-format files containing the amino acid sequences and SSSCs of protein subunits were obtained from the observed Protein Data Bank (PDB) files\textsuperscript{15} by using the SSSCview program (available online at https://staff.aist.go.jp/izumi.h/SSSCPreds/index-e.html)\textsuperscript{9}.

The original and mutation sequences of protein subunits were converted to the predicted SSSCs by using the SSSCPreds program (available online at https://staff.aist.go.jp/izumi.h/SSSCPreds/index-e.html)\textsuperscript{6}.

**Data availability**

The reference models and the original amino acid sequences were downloaded from the PDB. The mutation sequences were obtained based on References 4 and 5.

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**Author contributions**

H.I. designed methodology and software and performed research; L.A.N. and R.K.D. organized research; and H.I., L.A.N., and R.K.D. wrote the paper.

**Competing interests**

The authors declare no competing financial interest.

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