Immunoinformatics-aided identification of T cell and B cell epitopes in the surface glycoprotein of 2019-nCoV

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Abstract
The 2019 novel coronavirus (2019-nCoV) outbreak has caused a large number of deaths with thousands of confirmed cases worldwide, especially in East Asia. This study took an immunoinformatics approach to identify significant cytotoxic T lymphocyte (CTL) and B cell epitopes in the 2019-nCoV surface glycoprotein. Also, interactions between identified CTL epitopes and their corresponding major histocompatibility complex (MHC) class I supertype representatives prevalent in China were studied by molecular dynamics simulations. We identified five CTL epitopes, three sequential B cell epitopes and five discontinuous B cell epitopes in the viral surface glycoprotein. Also, during simulations, the CTL epitopes were observed to be binding MHC class I peptide-binding grooves via multiple contacts, with continuous hydrogen bonds and salt bridge anchors, indicating their potential in generating immune responses. Some of these identified epitopes can be potential candidates for the development of 2019-nCoV vaccines.

KEYWORDS
2019-nCoV, coronavirus, COVID-19, epitope prediction, immunoinformatics, SARS-CoV-2

1 | INTRODUCTION

The 2019 novel coronavirus (2019-nCoV; Family Coronaviridae), also known as the Wuhan coronavirus, has created a global emergency the likes of which had not been seen since the 2003 severe acute respiratory syndrome (SARS) outbreak. As of 27th January 2020, the death toll from the coronavirus stood at 80, with more than 2500 confirmed cases in China, as well as in 11 other countries.1

The virus, first identified in Wuhan, Hubei, China, causes a contagious, respiratory infection. The incubation period is reportedly 2 to 10 days, and studies are underway to determine whether the transmission is possible from asymptomatic individuals.1 Similar to other coronaviruses, 2019-nCoV reportedly possesses a surface glycoprotein. In more established coronavirus pathogenesis, this protein has been shown to bind host cellular receptors and to mediate membrane fusion.2 More importantly, it has been described as a potential vaccine target in both SARS-coronavirus (SARS-CoV) and Middle East respiratory syndrome coronavirus (MERS-CoV).3,4

This study aimed to identify significant cytotoxic T cell (CTL) and B cell epitopes within the 2019-nCoV surface glycoprotein using bioinformatic analysis. We also studied the potential interaction of the putative CTL epitopes with their corresponding representative major histocompatibility complex (MHC) class I supertypes most frequently observed in the Chinese population. We investigated hydrogen bonding, salt-bridges, and residue-residue contacts between CTL binding epitopes and the MHC class I molecules by molecular dynamics (MD) simulations.

2 | METHODS

Amino acid sequence of the 1273 AA, viral surface glycoprotein was retrieved in FASTA format from NCBI Protein Database (YP_009724390.1). Because of the novel nature of 2019-nCoV, we compared the glycoprotein's alignment with the spike glycoprotein of SARS-CoV (UniProt: P59594) to ensure sequence similarity in EMBOSS Water pairwise sequence aligner.5
NetCTL 1.2 server ([http://www.cbs.dtu.dk/services/NetCTL](http://www.cbs.dtu.dk/services/NetCTL)) was used to identify CTL epitopes within the glycoprotein sequence. NetCTL uses NetMHC server’s artificial neural networks (ANNs) to predict MHC binding, NetChop-3.0 to predict C-terminal cleavages and a weight matrix to calculate TAP transport efficiency to generate predictions. The scores of the three prediction methods are merged into a single score, which can be translated as sensitivity and specificity for the epitope-MHC-I binding event. We predicted MHC-peptide-binding for the following MHC class I supertypes—A2, A3, B7, B44, and B62. Together, these supertypes account for the majority of human leukocyte antigen A (HLA-A) (76.5%) and HLA-B (67.1%) distribution in the Chinese population.

Epitopes with a combined score of >0.75 (sensitivity = 0.80, specificity = 0.97) were further confirmed in CTLPred ([http://crdd.osdd.net/raghava/ctlpred](http://crdd.osdd.net/raghava/ctlpred)), where a consensus scoring approach was taken with thresholds of 0.51 (ANN) and 0.36 (support vector machines).

The three highest-scoring epitopes for each supertype were selected for analyses of binding affinity to their corresponding representative supertypes in NetMHCpan 4.0 server which predicted epitope-MHC-I binding using ANN trained on quantitative binding data and mass spectroscopy-derived MHC eluted ligands ([http://www.cbs.dtu.dk/services/NetMHCpan](http://www.cbs.dtu.dk/services/NetMHCpan)). Default threshold % ranks of 0.5 for strong binders, and 2 for weak binders were used.

Binding CTL epitopes were selected for homology modeling in PEP-FOLD3, and structures with the lowest coarse-grained energy were selected for protein-protein docking with corresponding HLA

### TABLE 1 CTL epitopes identified in 2019-nCoV surface glycoprotein and their corresponding MHC class I supertype representative

| Epitope     | Epitope score—ANN/SVM | Antigenicity (score) | HLA (% rank) |
|-------------|------------------------|----------------------|--------------|
| YLQPRTFLL   | 0.83/0.64              | Y (0.45)             | HLA-A*02:01 (0.01) |
| GVYFASTEK   | 0.58/0.98              | Y (0.71)             | HLA-A*03:01 (0.00) |
| EPVLKGVKL   | 0.73/0.61              | Y (1.23)             | HLA-B*07:02 (0.28) |
| VVNQNAQAL   | 0.77/0.78              | Y (0.47)             | HLA-B*07:02 (0.78) |
| WTAGAAAYY   | 0.82/0.54              | Y (0.63)             | HLA-B*15:01 (0.37) |

Abbreviations: 2019-nCoV, 2019 novel coronavirus; ANN/SVM, artificial neural network/support vector machine; CTL, cytotoxic T lymphocyte; HLA, human leukocyte antigen; MHC, major histocompatibility complex.

### FIGURE 1

Interactions between cytotoxic T lymphocyte binding epitopes and human leukocyte antigen (HLA) chains of corresponding major histocompatibility complex class I supertype representative during molecular dynamics simulations. A, Residue-residue contacts (X: epitope, Y: HLA) were mapped in contact map matrices. B, Salt bridge-forming residues were detected in three complexes. C, Continuous hydrogen bonds were observed between all epitopes and corresponding HLA chains throughout the simulations.
Complexes that scored the highest, based on interface atomic contact energy, shape complementarity, and electrostatics were selected for MD simulations in NAMD. Structures were solvated inside a TIP3P water box, and neutralized with Cl⁻ and/or Na⁺, as required. Energy minimization was carried out in a series of conjugate gradients to ensure appropriate structure geometry. Equilibration with restrained solute atoms was performed in an NPT ensemble (310 K, 1 bar) with periodic boundary conditions, and constant temperature and pressure dynamics via Langevin dynamics and Nose-Hoover Langevin piston respectively. Finally, free production runs, without positional restraints were carried out for 2 ns. Hydrogen bonds between epitopes and HLA chains were mapped in VMD with a two-way donor-acceptor distance of 3 Å and an angle of 20° as thresholds. Residue-residue contacts between epitopes and HLA chains were visualized in a contact map matrix, and salt bridge-forming residues were identified using the oxygen-nitrogen cut-off distance of 3.2 Å.

Sequential B cell epitopes within the surface glycoprotein were identified in BepiPred 2.0 (http://www.cbs.dtu.dk/services/BepiPred), and residues scoring above the default threshold of 0.5 were further confirmed in ABCPred, with default threshold of 0.51 and an overlapping filter (http://crdd.osdd.net/raghava/abcpred). BepiPred 2.0 uses a random forest algorithm trained on epitope and non-epitope amino acids validated from crystal structures, whereas ABCPred employs ANN using fixed-length patterns trained on 700 B cell epitopes and 700 random non-epitopes. To identify discontinuous B cell epitopes, we used Ellipro using the surface glycoprotein structure modeled in RaptorX, and the top five scoring epitopes were selected as significant. The modeled structure was validated by the Ramachandran plot generated in RAMPAGE. Antigenicity of all identified epitopes was predicted using VaxiJen 2.0. Conservation of the sequential epitopes within spike proteins across bat coronavirus (Bt-CoV, UniProt: A3EX94), SARS-CoV, and MERS-CoV (UniProt: K9N5Q8) were studied by multiple sequence alignment in Clustal Omega and visualized in NCBI Multiple Sequence Alignment Viewer.

### RESULTS

The surface glycoprotein of 2019-nCoV was found to have 76.3% identity and 87.3% similarity with the spike glycoprotein of SARS-CoV.

We identified five CTL epitopes in the surface glycoprotein, with all five epitopes classified as strong binders to their corresponding MHC class I supertype representative (Table 1). During MDs simulations, we observed the CTL epitopes to bind with the peptide-binding groove of MHC class I molecules, which is typically used for antigen presentation. The relevant residues of the epitopes and HLA chains in contact were mapped in a contact map matrix (Figure 1A). We observed salt bridge-forming residues positioned over the viral epitopes in three of the five complexes, as well as continuous hydrogen bonding between epitopes and HLA chains maintained throughout the simulation in all complexes (Figure 1B,C). Mutations in salt bridge-forming amino acids have been shown to weaken TCR recognition of MHC class I-antigen complexes. Also, the continuous formation of a set of hydrogen bonds between the MHC molecules and epitopes indicates an accommodating nature of the HLA chains to bind epitopes within the groove between the two helices. Our results are therefore optimistic of ideal antigen presentation to CTLs.

### TABLE 2  Sequential B cell epitopes identified in 2019-nCoV surface glycoprotein

| Sl. no | Epitope:position | Score |
|--------|------------------|-------|
| 1      | H1271, Y1272, T1273 | 0.998 |
| 2      | E1258, D1259, D1260, S1261, E1262, P1263, V1264, L1265, K1266, G1267, V1268, K1269, L1270 | 0.985 |
| 3      | G838, D839, C840, L841, G842, D843, I844, A845, A846, R847, D848, L849, I850, C851, A852, Q853, K854, F855 | 0.866 |
| 4      | Y1215, I1216, L1218, G1219, F1220, I1221, A1222, G1223, L1224, I1225, A1226, I1227, V1228, M1229, V1230, T1231, I1232, M1233, L1234, C1235, C1236, M1237, T1238, S1239, C1240, C1241, S1242, C1243, L1244, K1245, G1246, C1247, C1248, S1249, C1250, G1251, S1252, C1253, C1254, K1255, F1256, D1257 | 0.832 |
| 5      | M1, F2, V3, F4, L5, V6, L7, L8, P9, L10, V11, S12, S13, Q14, C15, V16, N17, L18 | 0.786 |

### TABLE 3  Discontinuous B cell epitopes identified in 2019-nCoV surface glycoprotein

| Sl. no | Epitope:position | Score |
|--------|------------------|-------|
| 1      | H1271, Y1272, T1273 | 0.998 |
| 2      | E1258, D1259, D1260, S1261, E1262, P1263, V1264, L1265, K1266, G1267, V1268, K1269, L1270 | 0.985 |
| 3      | G838, D839, C840, L841, G842, D843, I844, A845, A846, R847, D848, L849, I850, C851, A852, Q853, K854, F855 | 0.866 |
| 4      | Y1215, I1216, L1218, G1219, F1220, I1221, A1222, G1223, L1224, I1225, A1226, I1227, V1228, M1229, V1230, T1231, I1232, M1233, L1234, C1235, C1236, M1237, T1238, S1239, C1240, C1241, S1242, C1243, L1244, K1245, G1246, C1247, C1248, S1249, C1250, G1251, S1252, C1253, C1254, K1255, F1256, D1257 | 0.832 |
| 5      | M1, F2, V3, F4, L5, V6, L7, L8, P9, L10, V11, S12, S13, Q14, C15, V16, N17, L18 | 0.786 |

Abbreviations: 2019-nCoV, 2019 novel coronavirus; Bt-CoV, bat coronavirus; CTL, cytotoxic T lymphocyte; MERS-CoV, Middle East respiratory syndrome coronavirus; SARS-CoV, severe acute respiratory syndrome coronavirus.
We also identified three sequential B cell epitopes, and following modeling of 98% residues of the surface glycoprotein ($P = 1.03 \times 10^{-15}$, uGDT = 946), we further identified five discontinuous B cell epitopes in the structure (Tables 2 and 3). Ramachandran plot indicated 94.2% (1194 AA) residues in the favored region, 5% (63 AA) residues in the allowed region, and 0.8% (10 AA) residues in the outlier region (Figure 2).

With the exception of CTL epitope—VVNQNAQAL, which was found to have 100% identity with SARS-CoV, all identified sequential CTL and B cell epitopes were at least partially unique to 2019-nCoV, compared with Bt-CoV, SARS-CoV, and MERS-CoV. CTL epitope—EPVLKGVKL did not match any corresponding sequences of the other coronaviruses (Figure 3).

**FIGURE 2** Discontinuous B cell epitopes in 2019-nCoV surface glycoprotein and structure validation by Ramachandran plot. Left—five discontinuous B cell epitopes identified in the modeled structure, detailed in Table 3. Right—ramachandran plot indicated a high proportion of residues to be located within the favored and allowed regions. 2019-nCoV, 2019 novel coronavirus

**FIGURE 3** Visualization of frequency-based differences in residues of sequential CTL and B cell epitopes across bat coronavirus, MERS-coronavirus and SARS-coronavirus. CTL, cytotoxic T lymphocyte; MERS, Middle East respiratory syndrome; SARS, severe acute respiratory syndrome
We found one overlapping CTL epitope between MERS-CoV and SARS-CoV.\textsuperscript{22,23} We found one overlapping CTL epitope between MERS-CoV and 2019-nCoV with one gap and one mismatch (YLQPRTFLL/YKLQPLTFLL), and no comparable epitopes with SARS-CoV. In addition to the activation of CTLs, successful immunogens need to generate a persistent humoral immunity and in our study, we identified three sequential and five discontinuous B cell epitopes in the surface glycoprotein. There are a few limitations of our study. Epitope detection in glycoproteins is often complicated by the presence of covalently attached glycans. Modeling viral glycoproteins with the associated glycans is especially challenging due to the lack of related structures, although attempts have been made in hepatitis C and rabies.\textsuperscript{24,25} Second, similar to other RNA virus, coronavirus replication is error-prone, with a reported mutation rate of $4 \times 10^{-4}$ substitutions per site per year in SARS-CoV.\textsuperscript{26} This necessitates urgency in antiviral vaccine development before the identified epitopes are potentially rendered obsolete.

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