

## Construction of Epitope-Based Peptide Vaccine Against SARS-CoV-2: Immunoinformatics Study

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### Abstract

Recently, a novel coronavirus (SARS-CoV-2) appeared which is conscientious for the current outbreak in China and rapidly spread worldwide. Unluckily, there is no approved vaccine found against SARS-CoV-2. Therefore, there is an urgent need for designing a suitable peptide vaccine constituent against the SARS-CoV-2. In this study, we characterized the spike glycoprotein of SARS-CoV-2 to obtain immunogenic epitopes. In addition, we used 58 SARS-CoV-2 isolates were retrieved from the Global Initiative on Sharing All Influenza Data (GISAID) and National Center for Biotechnology Information (NCBI), then aligned to obtain the conserved region of SARS-CoV-2 spike glycoprotein. The interaction between the conserved region with ACE2 receptor, a SARS-CoV-2 receptor on the host cell, has been evaluated through molecular docking approach. The B-cell epitope was identified using the immune epitope database (IEDB) web server. Interestingly, we recommend Pep\_4 ADHQPQTFVNTLH as a epitope-based peptide vaccine candidate to deal with the SARS-CoV-2 outbreak. Pep\_4 has a high level of immunogenicity and does not trigger autoimmune mechanisms. Pep\_4 is capable of forming BCR/Fab molecular complexes with the lowest binding energy for activation of transduction signal the direct B-cell immune response. However, further study is suggested for confirmation (*in vitro* and *in vivo*).

**Keywords:** Coronavirus, Epitope-based peptide vaccine, Immunoinformatics, SARS-CoV-2

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## INTRODUCTION

On 30 December 2019, Chinese government reported an outbreak of pneumonia disease in Wuhan (Lam *et al.*, 2020). Furthermore, the causative agent identified by International Committee on Taxonomy of Viruses (ICTV) and named as SARS-CoV-2 (Gorbalenya *et al.*, 2020). The new virus has rapidly spread across China and other countries (Wang *et al.*, 2020). Then, WHO announced a new name for the disease caused by SARS-CoV-2 as coronavirus disease 2019 (COVID-19) (Lai *et al.*, 2020).

Up until today, there are more than three millions people are infected by SARS-CoV-2 worldwide. This data referring to online interactive dashboard hosted by the Center for Systems Science and Engineering (CSSE) at Johns Hopkins University, Baltimore, USA. This interactive web-based dashboard is tracking the reported cases of COVID-19 in real-time (Dong *et al.*, 2020). Presently, there are three coronaviruses to cause disease in the human, such as severe acute respiratory syndrome coronavirus (SARS-CoV), Middle-East respiratory syndrome coronavirus (MERS-CoV), and SARS-CoV-2 (Walls *et al.*, 2020).

Coronaviruses belong to the *Coronaviridae* family in the *Nidovirales* order. In addition, the structural proteins are encoded by the four structural genes, including membrane (M), spike (S), nucleocapsid (N), and envelope (E) (Shereen *et al.*, 2020). Interestingly, the spike glycoprotein plays an essential role in binding receptors on the host cell and the major target of neutralizing antibodies (Phan, 2020; Walls *et al.*, 2020). Thus, the spike glycoprotein is recently considered as as the greatly expectant antigen formulation to the SARS-CoV-2 vaccine construction.

Vaccines have been proven to decrease morbidity and mortality levels of many infectious diseases (Ansori and Kharisma, 2020). In addition, Callaway (2020) stated that approximately 90 vaccines are being developed against SARS-CoV-2 by research groups in universities and companies worldwide. Furthermore, scientists are trialing diverse technologies and some of which haven't been used in a licensed vaccine before. On the other hand, Shang *et al.* (2020) demonstrated that inactivated virus vaccines and live-attenuated

virus vaccines have their own side effects and disadvantages. New vaccine designs based on the protective antigens derived from SARS-CoV-2 should be considered.

The development of an effective vaccine against the SARS-CoV-2 infection is urgently required. The progress in molecular biology and biotechnology is driving the construction of the novel concepts in vaccinology. Synthetic recombinant proteins containing epitopes could be produced efficiently by using modern biotechnology methods (Oyarzún and Kobe, 2016). Therefore, the epitope-based peptide vaccine is suggested as new potential candidates for effective vaccine of SARS-CoV-2. In this study, we characterized the spike glycoprotein of SARS-CoV-2 to obtain immunogenic epitopes.

## MATERIALS AND METHODS

### Sample Retrieval from Database

The SARS-CoV-2 spike glycoprotein sequences were obtained from the Global Initiative on Sharing All Influenza Data (GISAID), Germany and National Center for Biotechnology Information (NCBI), USA.

### The Screening of Conserved Domain

The conserved domain in the SARS-CoV-2 viral spike glycoprotein sequences were identified through protein alignment with Molecular Evolutionary Genetics Analysis (MEGA) X, then the 3D structure was modeled using the SWISS-MODEL (<https://swissmodel.expasy.org>) with homology modeling approach (Schwede *et al.*, 2003; Adianingsih and Kharisma, 2019). Thus, the results are 3D models displayed in PyMol software (Himmah *et al.*, 2016; Putra *et al.*, 2020).

### Immunoinformatics Prediction

This study predictions based on linear B-cell epitopes, including BepiPred, Emini Surface Accessibility, Parker Hydrophilicity, and Karplus & Schulz Flexibility on the Immune Epitope Database Analysis Resource (IEDB-AR) web server (<http://tools.iedb.org/main/bcell>) (Hilyatuz *et al.*, 2016; Adianingsih and Kharisma, 2019; Ansori and Kharisma, 2020). Then we predicted the characteristics of candidate epitopes that can act as protective antigens using VaxiJen v2.0 web server (<http://www.ddgpharmfac.net/vaxijen/VaxiJen/VaxiJen.html>) (Hilyatuz *et al.*, 2016; Ansori and Kharisma, 2020). Prediction of similarity with

proteins in the cells of *Homo sapiens* is performed on vaccine candidate peptide via the Basic Local Alignment Protein Search Tool (BLASTp) (<https://blast.ncbi.nlm.nih.gov/Blast.cgi>). Peptides must have a low score below 20% to allow for the prevention of autoimmune (Himmah *et al.*, 2016; Adianingsih and Kharisma, 2019).

#### Protein Docking

Molecular interaction simulation between the host cell receptor, angiotensin-converting enzyme 2 (ACE2), and SARS-CoV-2 spike glycoprotein during viral attachment and mechanism for binding peptide vaccine candidate with B-cell receptors (BCR), using the protein docking on the Cluspro 2.0 web server (<https://cluspro.org/login.php>). Docking is used to determine the binding energy of molecular complexes and biological responses (Kozakov *et al.*, 2017; Adianingsih and Kharisma, 2019).

### RESULTS AND DISCUSSION

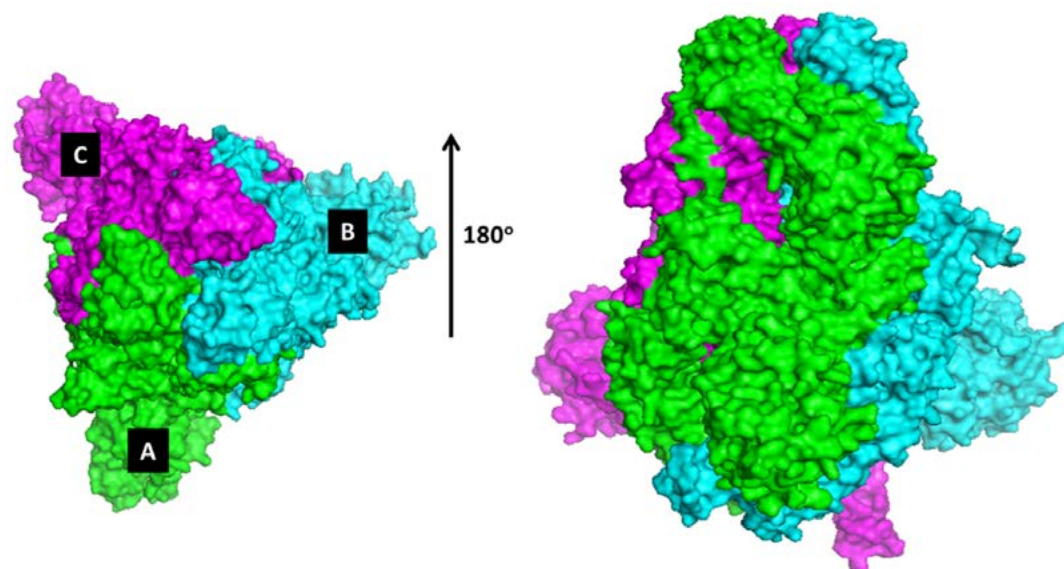
#### Conserved Identification from SARS-CoV-2 Spike Glycoprotein

A total of 58 isolates of SARS-CoV-2 spike glycoprotein sequences in the FASTA format were successfully obtained from the NCBI database with the prefixes "MN" and "EPI\_ISL\_" GISAID (Table 1). Then, protein alignment performed with MEGA X software to identify conserved domains until 229-mer sequences were identified

**MFVLLLSVPRDVLVLTGRVLVYIRGFNCGNVY  
FYLFKNVKHILDFLIRLSAYVLKALTVFLNRVY  
NTFKSLTASVEAQATCGGLFDIYPCGNLSGPV  
LSFAGTNVKCTGTGTFGDTDPTCGVPQLDLTY  
YSYQAYTNSPISMYECSLTNVTFQLSDFLID  
RCALLLITSLLMITYLISSNVFIVLRLSLILVGLA  
DHQPQTFVNTLHLIKLQLQLLICCVCVH**  
in SARS-CoV-2 spike glycoprotein as a conserved domain.

In this study, the identification of the conserved domain in 3D conducted via two stages, consisted of SARS-CoV-2 spike glycoprotein modeling and visualization in PyMol software with a representative structure (Adianingsih and Kharisma, 2019). Homology modeling on the SWISS-MODEL web server has a positive predictive score of approximately 20% (Schwede *et al.*, 2003; Waterhouse *et al.*, 2018). This study obtained the model of glycoprotein from a query protein sequence (MN908947.3) with a template (ID 6ACD) around 76.47% identified as a spike spike glycoprotein, so that the spike glycoprotein model was similar to the template and homologous. The 3D structure of SARS-CoV-2 glycoprotein is displayed in the form of rigid surfaces with a color selection based on its protein chains (Fig. 1).

The process of docking proteins in Cluspro is to determine the binding energy that is formed between spike glycoprotein and ACE2 receptor



**Fig. 1.** 3D model of SARS-CoV-2 spike glycoprotein. Protein structure consists of A (green), B (cyans), and C (magenta) chains.

when the SARS-CoV-2 virion attachment occurs in the host cell. The results showed that the molecular complex arrangement was identified to have the lowest binding energy of 1152.6 kcal/mol to achieve a achieve of stability (Table 2), in the glycoprotein\_ ACE2 binding formation. The more negative binding energy values can trigger the formation of stable molecular complexes (Kozakov *et al.*, 2017; Adianingsih and Kharisma, 2019). This

stability can trigger the activation of biological responses to target proteins such as mediated endocytosis during the SARS-CoV-2 viral entry (Cao *et al.*, 2020). The complex of molecular docking displayed in PyMol with representative structures and color selections to identify between receptors (ACE2) and ligands (SARS-CoV-2 spike glycoprotein) (Fig. 2).

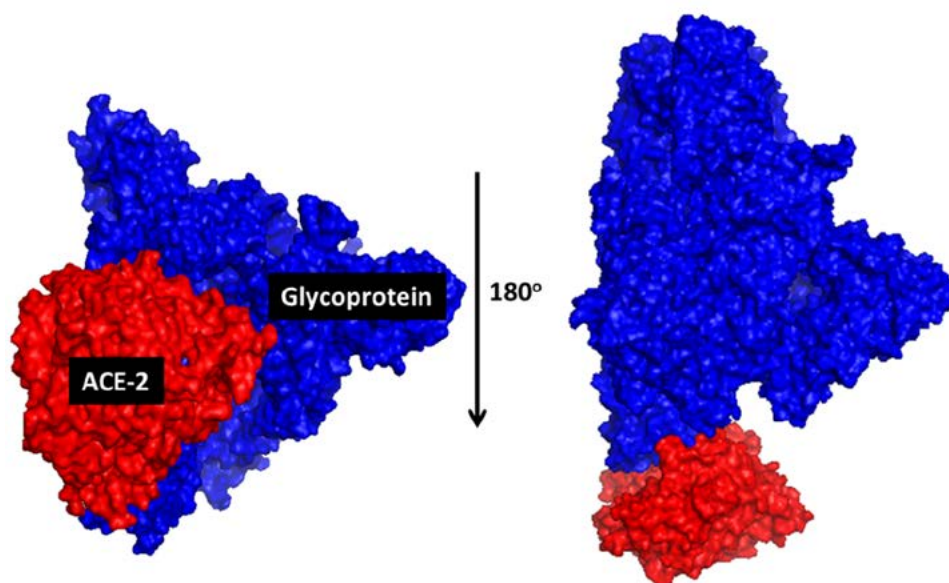
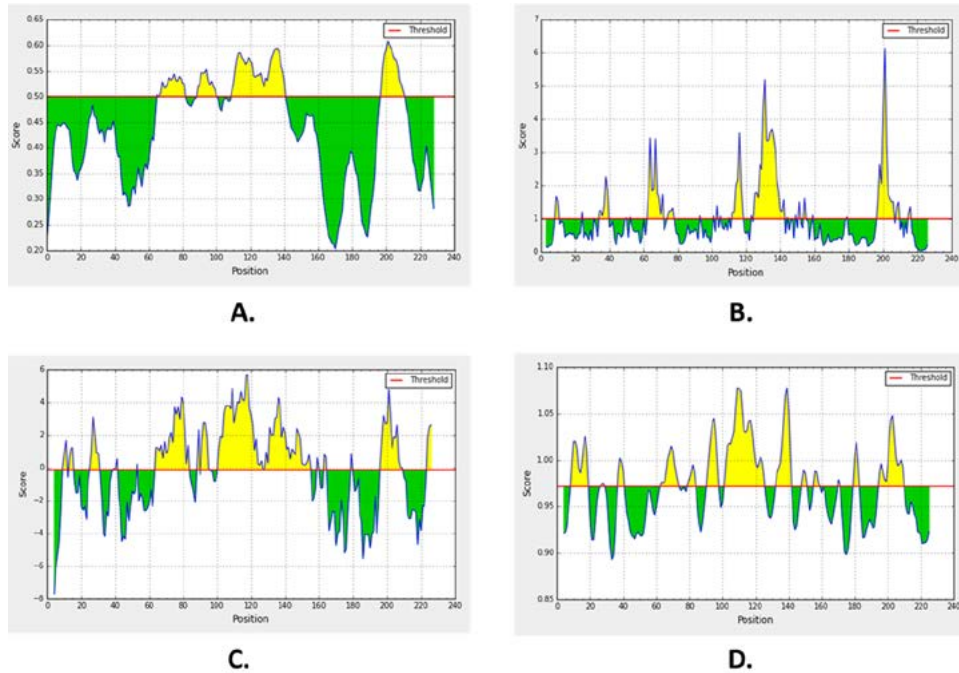


Fig. 2. Molecular visualization of the ACE2 (red) molecular complex and SARS-CoV-2 spike glycoprotein (blue). Both are displayed in a rigid surface structure with specific colour selection.

### The B-cell Immunogenicity Predictions of Peptide Vaccine Candidate

The conserved domain on SARS-CoV-2 spike glycoproteins analyzed for the potential of B-cell immunogenicity to obtain specific peptides as vaccine candidates. Prediction of B-cell immunogenicity consists of BepiPred ( $\geq 0.222$ ), Emini Surfaces Accessibility ( $\geq 1.000$ ), Parker Hydrophilicity ( $\geq 2.068$ ), and Karplus & Schulz Flexibility ( $\geq 1.017$ ) parameters, each has a threshold score for predicting B-cell epitopes in linear sequences (Sanchez-Trincado *et al.*, 2017; Adianingsih and Kharisma, 2019; Ansori and Kharisma, 2020). B-cell epitope prediction shows on BepiPred parameters, it has 0.608 threshold points, Emini Surfaces Accessibility 1.000, Parker Hydrophilicity 6.138, and Karplus & Schulz Flexibility 1.078 (Fig. 3). The

scores show positive B-cell epitopes prediction because no one has a threshold score below the standard, peptides are obtained Pep\_1 FKSILTASVEAQATCG, Pep\_2 CGNLSGPVLSFA, Pep\_3 TGTFGDTPD CGVPQLDLTYYSYQAYTNSPI, and Pep\_4 ADHFFQQHHF. We predicted the antigenicity of the peptides in VaxiJen v2.0, and obtained peptides number 1, 3, and 4 have antigenic properties. Furthermore, the peptide was analyzed for its similarity to the cell surface receptors of the body of *Homo sapiens*, via the BLASTp server. Peptides number 1, 3, and 4 did not have similarities to the cell surface receptors of the body of *Homo sapiens* with a score of less than 20%, so it is predicted that it will not initiate autoimmune occurrence when produced in vaccine form (Himmah *et al.*, 2016; Adianingsih and Kharisma, 2019).

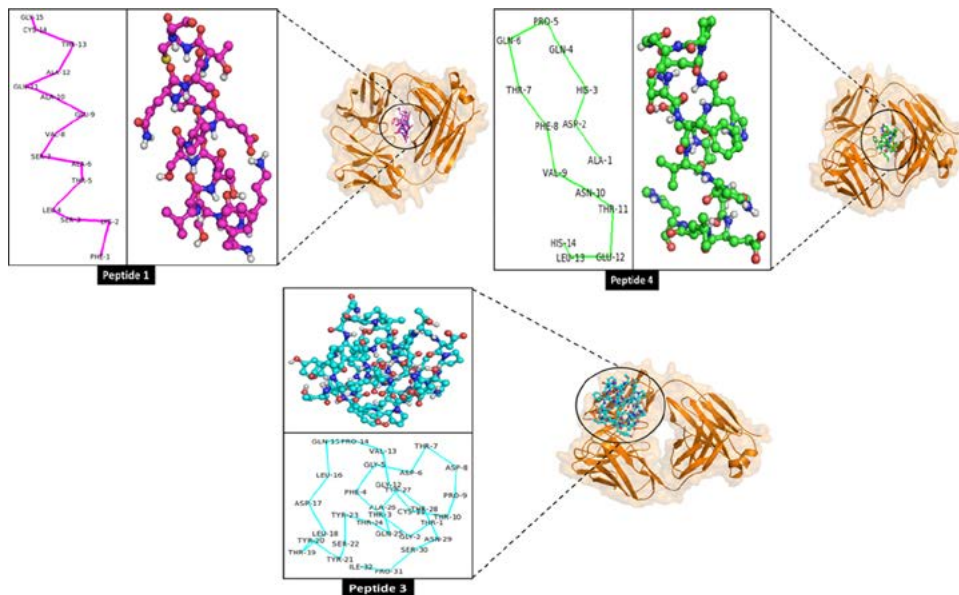


**Fig. 3.** The results of B-cell epitope prediction. Green areas showed negative prediction and yellow areas for positive prediction. (A) BepiPred; (B) Emini Surfaces Accessibility; (C) Parker Hydrophilicity; and (D) Karplus & Schulz Flexibility.

**Molecular Interaction between Peptide-BCR/Fab**

The positive predicted peptide results on B-cell immunogenicity, then the 3D structure is modeled via PEP-FOLD using the fold recognition modeling

(Hilyatuz *et al.*, 2016). Then, the structure is obtained and stored in the .pdb format, then the 3D sample of the antigen-binding fragment BCR receptor (Fab) (ID 5IFH) is obtained from the PDB



**Fig. 4.** Structure visualization of sticks, spheres, and transparent surfaces of molecular complexes consisting of peptide-based vaccine candidate with BCR/Fab (orange).

**Table 1.** SARS-CoV-2 isolates from the biological databases (GISAID and NCBI).

Sample ID	
MN908947.3	EPI_ISL_403928
MN938384.1	EPI_ISL_402132
MN975262.1	EPI_ISL_403963
MN996527.1	EPI_ISL_403962
MN996528.1	EPI_ISL_402120
MN996529.1	EPI_ISL_402121
MN996530.1	EPI_ISL_402119
MN996531.1	EPI_ISL_406031
MT007544.1	EPI_ISL_406531
MN994467.1	EPI_ISL_406533
MN994468.1	EPI_ISL_406534
MN997409.1	EPI_ISL_406535
MN988668.1	EPI_ISL_406536
MN988669.1	EPI_ISL_406538
EPI_ISL_404253	EPI_ISL_406592
EPI_ISL_404895	EPI_ISL_406593
EPI_ISL_403937	EPI_ISL_406594
EPI_ISL_403936	EPI_ISL_406595
EPI_ISL_403935	EPI_ISL_406596
EPI_ISL_403934	EPI_ISL_406597
EPI_ISL_403933	EPI_ISL_406862
EPI_ISL_403932	EPI_ISL_406798
EPI_ISL_404228	EPI_ISL_406800
EPI_ISL_404227	EPI_ISL_406801
EPI_ISL_402123	EPI_ISL_406970
EPI_ISL_403931	EPI_ISL_406973
EPI_ISL_403930	EPI_ISL_407071
EPI_ISL_403929	EPI_ISL_407073
EPI_ISL_407084	EPI_ISL_407193

**Table 2.** Binding energy of ACE-2 with SARS-CoV-2 spike glycoprotein

Molecular Complex	Cluster	Member	Lowest Energy (kcal/mol)
ACE-2_Glycoprotein	0	36	-993.2
ACE-2_Glycoprotein	1	35	-1152.6
ACE-2_Glycoprotein	2	31	-933.5
ACE-2_Glycoprotein	3	29	-1042.0
ACE-2_Glycoprotein	4	26	-912.2
ACE-2_Glycoprotein	5	24	-925.0

database. Protein-peptide docking simulation using Cluspro to determine the binding energy is needed in the formation of stable complexes and activation of biological responses at BCR receptors (Kozakov *et al.*, 2016; Adianingsih and Kharisma, 2019). The results show that the binding

energy produced by the Pep\_1-BCR/Fab molecule complex is -572.9 kcal/mol, Pep\_3-BCR/Fab -661.3 kcal/mol, and Pep\_4-BCR/Fab -686.5 kcal/mol, the results are displayed in PyMol software by the structural and color selection (Figure 4). The analysis showed that Pep\_4 ADHQPQTFVNTLH was predicted as a SARS-CoV-2 vaccine candidate with a length of 14-mer. Pep\_4 has the potential as an epitope-based peptide vaccine candidate because it has the lowest binding energy score. The lowest binding energy allows the formation of molecular complexes (Fu *et al.*, 2018). Binding energy is influenced by weak bonds such as hydrogen and hydrophobicity (Pourseif *et al.*, 2019). The formation of a stable molecular complex between Pep\_4-BCR/Fab allows for the activation of a direct immune response by B cell (Bros *et al.*, 2019). Specific antibodies for neutralization of SARS-CoV-2 and the role of memory cells to keep the body formed from future infections.

**CONCLUSION**

In conclusion, we recommend Pep\_4 ADHQPQTFVNTLH as epitope-based peptide vaccine candidate to deal with the SARS-CoV-2 outbreak. Pep\_4 has a high level of immunogenicity and does not trigger autoimmune mechanisms. Pep\_4 is capable of forming BCR/Fab molecular complexes with the lowest binding energy for activation of transduction signal the direct B-cell immune response.

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**CONFLICT OF INTEREST**

The listed authors declare no conflict of interest in any capacity, including competing or financial.

**AUTHORS' CONTRIBUTION**

All listed authors have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

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## ETHICS STATEMENT

This article does not contain any studies with human participants or animals performed by any of the authors.

## AVAILABILITY OF DATA

Not applicable.

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