Letter to the Editor

Infection-enhancing anti-SARS-CoV-2 antibodies recognize both the original Wuhan/D614G strain and Delta variants. A potential risk for mass vaccination?

The aim of the present study was to evaluate the recognition of SARS-CoV-2 Delta variants by infection enhancing antibodies directed against the NTD. The antibody studied is I052 (pdb file #7LAB) which has been isolated from a symptomatic Covid-19 patient. Molecular modeling simulations were performed as previously described. Two currently circulating Delta variants were investigated, with the following mutational patterns in the NTD:

- G142D/E154K (B.1.617.1)
- T19R/E156G/del157/del158/A222V (B.1.617.2)

Each mutational pattern was introduced in the original Wuhan/D614G strain, submitted to energy minimization, and then tested for antibody binding. The energy of interaction (ΔG) of the reference pdb file #7LAB (Wuhan/D614G strain) in the NTD region was estimated to −229 kJ/mol⁻¹. In the case of Delta variants, the energy of interaction was raised to −272 kJ/mol⁻¹ (B.1.617.1) and −246 kJ/mol⁻¹ (B.1.617.2). Thus, these infection enhancing antibodies not only still recognize Delta variants but even display a higher affinity for those variants than for the original SARS-CoV-2 strain.

The global structure of the trimeric spike of the B.1.617.1 variant in the cell-facing view is shown in Fig. 1A. As expected, the facilitating antibody bound to the NTD (in green) is located behind the contact surface so that it does not interfere with virus-cell attachment. Indeed, a preformed antibody-NTD complex could perfectly bind to the host cell membrane. The interaction between the NTD and a lipid raft is shown in Fig. 1B, and a whole raft-spike-antibody complex in Fig. 1C. Interestingly, a small part of the antibody was found to interact with the lipid raft, as further illustrated in Figs. 1D-E. More precisely, two distinct loops of the heavy chain of the antibody encompassing amino acid residues 28–31 and 72–74, stabilize the complex through a direct interaction with the edge of the raft (Fig. 1F). Overall, the energy of interaction of the NTD-raft complex was raised from −399 kJ.mol⁻¹ in absence of the antibody to −457 kJ.mol⁻¹ with the antibody. By clamping the NTD and the lipid raft, the antibody reinforces the attachment of the spike protein to the cell surface and thus facilitates the conformational change of the RBD which is the next step of the virus infection process.

This notion of a dual NTD-raft recognition by an infection enhancing antibody may represent a new type of ADE that could be operative with other viruses. Incidentally, our data provide a mechanistic explanation of the FcR-independent enhancement of infection induced by anti-NTD antibodies. The model we propose, which links for the first time lipid rafts to ADE of SARS-CoV-2, is in line with previous data showing that intact lipid rafts are required for ADE of dengue virus infection.

Neutralizing antibodies directed against the NTD have also been detected in Covid-19 patients. The 4A8 antibody is a major representative of such antibodies. The epitope recognized by this antibody on the flat NTD surface is dramatically affected in the NTD of Delta variants, suggesting a significant loss of activity in vaccinated people exposed to Delta variants. More generally, it can be reasonably assumed that the balance between neutralizing and facilitating antibodies may greatly differ according to the virus strain (Fig. 2).

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Current Covid-19 vaccines (either mRNA or viral vectors) are based on the original Wuhan spike sequence. Inasmuch as neutralizing antibodies overwhelm facilitating antibodies, ADE is not a concern. However, the emergence of SARS-CoV-2 variants may tip the scales in favor of infection enhancement. Our structural and modeling data suggest that it might be indeed the case for Delta variants.

In conclusion, ADE may occur in people receiving vaccines based on the original Wuhan strain spike sequence (either mRNA or viral vectors) and then exposed to a Delta variant. Although this potential risk has been cleverly anticipated before the massive use of Covid-19 vaccines\(^6\), the ability of SARS-CoV-2 antibodies to mediate infection enhancement in vivo has never been formally demonstrated. However, although the results obtained so far have been rather reassuring\(^7\), to the best of our knowledge ADE of Delta variants has not been specifically assessed. Since our data indicate that Delta variants are especially well recognized by infection-enhancing antibodies targeting the NTD, the possibility of ADE should be further investigated as it may represent a potential risk for mass vaccination during the current Delta variant pandemic. In this respect, second generation vaccines\(^8\) with spike protein formulations lacking structurally-conserved ADE-related epitopes should be considered.

References

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