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Orientation of immobilized antigens on common surfaces by a simple computational model: Exposition of SARS-CoV-2 Spike protein RBD epitopes

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HIGHLIGHTS

\begin{itemize}
  \item Protein immobilization is a crucial in preparing immunosensors.
  \item Immobilization must ensure exposition of the epitope.
  \item A simple computational model can help predicting the exposition.
\end{itemize}

GRAPHICAL ABSTRACT

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ABSTRACT

The possibility of immobilizing a protein with antigenic properties on a solid support offers significant possibilities in the development of immunosensors and vaccine formulations. For both applications, the orientation of the antigen should ensure ready accessibility of the antibodies to the epitope. However, an experimental assessment of the orientational preferences necessarily proceeds through the preparation/isolation of the antigen, the immobilization on different surfaces and one or more biophysical characterization steps. To predict a priori whether favorable orientations can be achieved or not would allow one to select the most promising experimental routes, partly mitigating the time cost towards the final product. In this manuscript, we apply a simple computational model, based on united-residue modelling, to the prediction of the orientation of the receptor binding domain of the SARS-CoV-2 spike protein on surfaces commonly used in lateral-flow devices. These calculations can account for the experimental observation that direct immobilization on gold gives sufficient exposure of the epitope to obtain a response in immunochemical assays.

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1. Introduction

The activity and reactivity of an immobilized protein strongly depend on its orientation with respect to the surface of the support in - or on - which it is immobilized. This holds true for enzymes, as well as for antibodies and antigens. Therefore, the possibility to control and manipulate the exposition of the relevant residues and protein surfaces plays an important role in the rational design of devices based on immobilized proteins. Among these devices, immunosensors represent an expanding space for research and market opportunities.

While the path to reach a technologically relevant product must rely upon a strong experimental characterization [1,2], possibly relying upon atomic-level methodologies [3–9], the preparation/isolation of the protein of interest, its immobilization, and the characterization of the resulting composite are complex and time-consuming, therefore it is also true that guidelines for achieving optimal orientations could improve the efficiency of the R&D connected to protein immobilization [10,11]. However, simplified simulation models that would allow for a rapid prediction of the most plausible orientations are not particularly common. In this manuscript we apply a very simple method based on a united-residue modelling of protein-surface interactions, to specifically address the problem of determining the orientation of the SARS-CoV-2 Spike protein Receptor Binding Domain (RBD) on a few prototypical surfaces for biomedical use. United residue modelling of protein-surface interactions is a rather effective method to screen the poses of protein molecules with respect to surfaces [12,13]. The method we apply is based on the works by Jiang, Zhou and del Monte-Martinez [10,12–18], and encompasses van der Waals and electrostatic interactions, as well as covalent immobilization.

The choice of the target protein is motivated by the recent emergence of a new infectious disease (COVID-19) caused by a coronavirus (SARS-CoV-2) [19]. This infectious disease has spread significantly upon the world, counting 13.841.890 infected people and a death toll of 590.845 as of July 2020 [20]. Models suggest that it will remain circulating and active for several months [21,22], and there is a marked possibility that reinfeciton is possible [23,24], thus increasing the time of the circulation of the virus. This pandemic outbreak has had a major impact on world economics, with a very long outlook [25]. A capillary activity of the diffusion of the infection has proven crucial [26], and serological tests are expected to have a key role in mass screening [27,28].

2. Methods

The structures of the proteins were downloaded from the protein databank (PDB) [29], the pKa values of reactive groups were calculated using PROPKA [30,31], and the interfaces were calculated using the PDBe PISA server [32]. The non-bonded interaction of a residue of type \( i \) is represented with a Lennard-Jones (LJ) potential: [12,13].

\[
U(r) = 4\epsilon_i \left( \frac{\sigma_i}{r + \delta_i} \right)^{12} - \left( \frac{\sigma_i}{r + \delta_i} \right)^{6}
\]

where \( r \) is the nearest distance between the residue and the surface, \( \epsilon_i \) the energy at the minimum position, \( \sigma_i \) is the equivalent van der Waals radius of each residue and \( \delta_i \) is a size parameter taken from the literature (see tables S2-S7, parameters are taken from [14,16,18,33–37], as indicated in the table captions).

The electrostatic interaction is represented through the Gouy-Chapman potential [12,13,38].

\[
U(r) = \frac{q_i q_e e^{-\kappa r}}{4\pi\varepsilon_0}
\]

where \( r \) is the nearest distance between the i-th residue with charge \( q_i \) and the surface, \( \kappa \) is the surface charge density, \( \varepsilon \) is the inverse Debye Length calculated from the ionic strength \( I \) as \( \kappa = 0.304/\sqrt{I} \), and the relative permittivity of the medium is assumed to be distance-dependent \( \varepsilon(r) = r \) [13,38]. A 1:1 buffer salt concentration of 0.15 mol dm\(^{-3}\) is assumed.

For silica, the surface charge density is estimated to be \( -0.3\epsilon \) m\(^{-2}\) [37].

For self-assembled charged monolayers (SAM), the charge density is set to \( +0.02\epsilon \) m\(^{-2}\) for the amino-capped monolayer (SAM-NH\(_2\)) and to \( -0.02\epsilon \) m\(^{-2}\) for the carboxyl-capped monolayer (SAM-CO\(_2\)H) [14,18].

The formation of a covalent bond is treated with the following potential:

\[
U(r) = \begin{cases} 
\epsilon_b & \text{if } r \leq a_i \\ 0 & \text{elsewhere} 
\end{cases}
\]

where \( \epsilon_b \) is the bond energy and is set to 600 kJ mol\(^{-1}\) for imino bonds [39] and 100 kJ mol\(^{-1}\) for gold-thiol bonds [40–42], regardless of the starting oxidation state of the thiol [43]. The desolvation energy is already accounted for in the vdW term.

LJ parameters for epoxide-glyoxyl functionalization is assumed to be equal to SAM-CO\(_2\)H, whereas for gold the parameters have been adapted from reference [36].

The sampling of the relative protein-surface orientations is performed by rotating a plane around the center of mass of the protein. The plane is initially parallel to the z = 0 plane. Only two rotations are necessary, as all the rotations around the normal to the plane will yield the same energy. The first rotation by an angle \( \alpha \in [0,\pi] \) is applied around the y-axis, followed by another rotation of an angle \( \beta \in [0,2\pi] \) around the z-axis, and then a translation is applied to optimize the position, similarly to what is done in the popular PALES software [44–46]. The sampling of the \( \alpha, \beta \) pairs is made uniform by using REPULSION angular sampling [47–49]. The distance of the plane to the protein is then set by minimizing the energy terms described above.

3. Results and discussion

The most important feature of a composite thought for immunochemical applications is that the orientation of the antigen with respect to the surface must ensure the accessibility of the epitope to the antibodies, to guarantee the recognition. Therefore, we have selected the crystallographic structure of RBD in complex with a fragment (FAB) of the human antibody CR3022 (PDB ID: 6W41) [50], and identified the interface residues relevant for the interaction (Fig. 1 and Table S1). In the analysis of the orientations, we assume that the full length antibody will have the same accessibility as the FAB because of the high flexibility of the linkers of the heavy chains (see fig. S1) [51–54]. It is also important to note that, while the spike protein is highly glycosylated at N- and O- positions [55,56], the structured part of the RBD which is recognized by the antibody only carries one glycation at position 343 [55] (pink in Fig. 1), and the glycation site faces away from the spike protein.

Fig. 1. Biological assembly from the crystallographic structure 6 W41. The SARS-CoV-2 receptor binding domain is shown in light green, with the interacting residues highlighted in red and the N-terminus highlighted in green. The glycation site 343 is highlighted in pink. The fragment of the human antibody CR3022 is shown in blue. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
3.1. Interaction with a hydrophobic surface

Hydrophobic adsorption occurs selectively on hydrophobic carriers at low ionic strength [57]. It is a rather common immobilization protocol, because of its simplicity. The interaction is here represented only through a simple Lennard-Jones (LJ) potential, the parameters of which have been defined according to the hydrophobicity index (table S2) [33–36]. The most probable orientation (5‰ relative population) is shown in Fig. 2, with the surface represented as a disk. In this, and in the following representations, the interaction is calculated for the antigen alone, and then the complex is shown for examining the interference of the surface with the binding. Of the 2000 considered orientations, 263 are within 10% of the probability of the orientation shown in Fig. 2. Most of those orientations involve contacts between the interface residues and the surface and are therefore expected to be poorly efficient for the recognition. This is not completely unexpected; as hydrophobic carriers mimic the interfaces formed by the naturally occurring interfaces of the proteins.

3.2. Interaction with charged surfaces

Also this immobilization strategy is rather common because of its simplicity. It is slightly less general, because the outcome strongly depends on the nature of the protein and of the surface. The electrostatic interaction of the i-th residue with the uniformly charged surface with a given charge density is estimated by the Gouy-Chapman potential [12,13], which is added to a LJ term. The parameters defining each system are listed in tables S3–S6.

We have considered the following surfaces:

1) silica - a common chromatographic support with high negative surface charge;
2) positively charged self-assembled monolayer (SAM), with amino capping of the chains [14,18];
3) negatively charged SAM, with carboxylic capping of the chains [14,18].

Supports #2 and #3 imply the possibility of colorimetric detection through gold [58], vide infra.

The most probable orientations are shown in Fig. 3.

The relative populations of the orientations shown in Fig. 3 are 100% for silica and SAM-NH₂. For SAM-CO₂H, the orientations in Fig. 3b is populated for about 25%, and there are other 6 orientations out of 2000 that have relative population above 2%, all within a few degrees from the one with highest relative population, except one that is more tilted, yielding a larger accessibility, with a relative population around 4% (Fig. S2).

It is apparent that only negatively charged surfaces allow for the exposition of the epitope, and this is anyway relatively marginal. These results suggest that it would be nontrivial to achieve a good orientation relying upon adsorption, either based on hydrophobic or on charge interactions. Therefore, we have considered directed approaches based on stronger interactions. In particular, we have considered epoxide-glyoxyl (directed at primary amine moieties) [59] and gold (thiols and disulfide bridges) [58].

The glyoxyl-based approach is quite popular for multipoint orientation-selective immobilization of proteins on surfaces. It involves a two-step mechanism, in the first step, the primary amine groups of the protein are allowed to react with the aldehyde groups to form Schiff base bonds, in the second step the bonds are reduced with sodium borohydride. This kind of immobilization has been simulated in a similar way as described by del Monte-Martínez et al. [11], assuming a working pH = 7.5, to maximize the reactivity of the N-terminus and at the same time limiting the reactivity of lysine residues (see table S7).

The choice of gold is also extremely popular, because of two reasons: the strong plasmonic response of gold, which causes a purple coloring of the bioconjugate, and because of the relatively easy manipulation required. Current SARS-CoV2 serological tests are indeed based on gold conjugates [60]. The conjugation to the surface is simulated in the same way as the amine-glyoxyl reaction, assuming that all cysteines are equally reactive towards gold (disulfide bridges can interact with gold to a comparable extent as thiols) [43]. The resulting orientation has 100% relative population. Colloidal gold has a net negative surface charge [61], but including the electrostatic term has no impact on the recovered orientation.

In the epoxide-glyoxyl strategy, the conjugation appears to be mostly directed at the N-terminus,¹ which is facing away from the recognition interface but is not topologically very remote. Therefore, the epitope will only be partially exposed, whereas for the gold conjugate, ample access to the epitope is possible in the most probable orientation (Figure 4).

Finally, a completely different strategy could be applied for conjugation to (e.g.) gold nanoparticles: the use of a avidin-biotin affinity system [62]. Biotinylation can be achieved through amine-specific reagents [63], and improvement in the selectivity can be achieved with minimal engineering of the sequence [64]. Given that there is a rather substantial difference in the calculated pKas for the different amine sites (see table S7), it can be expected that, for pH values lower than 7, all lysine residues will be protonated and thus less reactive with probability higher than 99%. The N-terminus is not facing the interaction site (see Fig. 1). Therefore, selective biotinylation at the N-terminus is expected to be possible. In this case, the accessibility of the epitope is warranted if the interaction between the antigen and streptavidin, if at all possible, is sufficiently weak.

To explore this possibility we have performed an initial-stage docking using ZDOCK [65], and inspected the first two elements that had a significantly higher ZDOCK score (Fig. S3). The possible interaction between the RBD and streptavidin was investigated also using HADDOCK2.4 [66]. The protein-protein interface residues were

¹ 80 orientations out of 2000 account for 99.9% of the relative population. All of these orientations involve binding of the N-terminus.
predicted with CPORT [67], and then used as “active” and “passive” residues in the HADDOCK calculation. About 10 lowly populated clusters with weak energy were obtained; the most significant three with the lowest HADDOCK-scores are reported in Fig. S3 and their energies in Table S8. Both dockings indicate that, should the interaction occur, it would occur in a position that does not interfere with the antigen-antibody recognition.

4. Conclusions

In this work, we describe the use of united-residue modelling for the prediction of the orientation of the receptor binding domain of the spike protein of the novel coronavirus SARS-CoV-2, a protein of high immunological relevance at the most commonly used surfaces for the preparation of lateral-flow immunochemical devices. With this simple, yet very flexible approach, we find that immobilization on silica, or through glyoxyl reaction of amine residues, or on gold yield orientations compatible with antibody recognition, with gold granting the highest exposition. In this way, we can explain why random conjugation of the RBD to a gold surface yields responsive immunosensors, which are now routinely used. A more detailed experimental verification of the predictions of protein orientation at surfaces represents a significant challenge for the current biophysical methodologies [1]. One can expect that cryo-electron transmission microscopy will be limited by the fact that, in most cases, the surface has higher electron density than that of the protein. Confocal laser scanning microscopy can be used to assess the positioning of the protein with respect to the support, and super-resolution microscopic techniques, such as total internal reflection fluorescence microscopy also allow for the detection of discrete molecular events (e.g., desorption, unfolding, lateral

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**Fig. 3.** Antigen-FAB complex shown in superposition with the most probable positioning with respect to charged surfaces: (a) silica, (b) SAM-CO₂H (negative) and (c) SAM-NH₂ (positive).

**Fig. 4.** Antigen-FAB complex shown in superposition with the most probable positioning with respect to covalently bound surfaces (a) epoxide-glyoxyl and (b) gold. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
diffusion, ... \cite{2}, but the orientation is still a high-hanging fruit by these methodologies. Conversely, the interaction between the protein and the interface can be probed at the atomic level through the application of solid-state NMR \cite{4,7–9,68–70}, effort which is being started in our lab. Our results suggest that very simple modelling approaches can provide significant hints towards rationally orienting antigens in a way to maximize the exposition of epitopes, and therefore help in the initial moments of the design of conjugates for immunologic applications, when a rapid response to emergency is vital. This is also testified by the emergence of theoretical modelling of several molecular aspects of viral infection and inhibition mechanisms \cite{71–76}. Overall, the expected short-time impact of our work is to provide guidelines to avoid the experimental exploration of immobilization pathways that are less promising.

Declaration of Competing Interest

The authors declare no competing financial interest.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.bpc.2020.106441.

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[6] D. Tian, T. Li, R. Zhang, Q. Wu, T. Chen, P. Sun, A. Ramamoorthy, Conformations of solid-state NMR \cite{4,7–9,68–70}, effort which is being started in our lab. Our results suggest that very simple modelling approaches can provide significant hints towards rationally orienting antigens in a way to maximize the exposition of epitopes, and therefore help in the initial moments of the design of conjugates for immunologic applications, when a rapid response to emergency is vital. This is also testified by the emergence of theoretical modelling of several molecular aspects of viral infection and inhibition mechanisms \cite{71–76}. Overall, the expected short-time impact of our work is to provide guidelines to avoid the experimental exploration of immobilization pathways that are less promising.

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