Considering both small and large scale motions of vascular endothelial growth factor (VEGF) is crucial for reliably predicting its binding affinities to DNA aptamers†

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Vascular endothelial growth factor 165 (VEGF165), a predominant isoform of VEGF signal proteins, is an ideal target for developing drugs against various diseases. It is composed of a heparin binding domain (HBD) and a receptor binding domain (RBD), which are connected by a flexible linker. Among the two domains, RBD is utilized in binding with the signal transduction protein, the VEGF receptor (VEGFR). None the less for its pharmaceutical importance, structure-based studies for developing drugs has been severely hindered by the lack of its whole structure determination, mainly owing to the existence of the flexible linker. Fortunately, the utilization of computer simulation methods can offer a possibility to circumvent this difficult issue. Here, we employ ensemble docking in combination with the anisotropic network model analysis to examine the interactions between DNA aptamers and VEGF165. We model three-dimensional structures of aptamer variants based on their sequence information and perform docking calculations with the whole VEGF165 structure. Indeed, we show that we can closely reproduce the experimental binding affinity order among different DNA aptamer variants by inclusively considering the flexible nature of VEGF. In addition, we address how DNA aptamer that binds to HBD of VEGF 165 impedes the interaction between VEGFR and VEGF165 through RBD, even though HBD and RBD are rather distant. The present study illustrates that the flexible docking scheme employed here can be applied to tricky cases that involve flexible proteins with undetermined structures, toward effectively predicting ligand binding affinities to such proteins.

VEGF165 is composed of the heparin binding domain (HBD) and the receptor binding domain (RBD) which are connected by a flexible linker. The structure of each domain has been determined, but mainly due to the flexible linker, the whole VEGF165 structure has not been determined. The correct structural information on a target protein to start with is crucial for computational pharmaceutical studies such as docking, but proteins with undetermined structural parts due to flexible regions render the computation quite tricky. Therefore, despite the potential as a promising drug target against AMD, no docking study considering the whole structure of VEGF165 has been carried out, except one recent study that examined the interaction between VEGF165 and heparin. Even in that study, the flexibility of VEGF165 was not taken into account during docking.

Experimentally, aptamers are considered as important drug candidates against AMD. Aptamers are single stranded oligonucleotides that bind to target molecules with high specificity and affinity. They are generated through the systematic evolution of ligands by exponential enrichment (SELEX) process, during which aptamers are screened from large and randomized nucleic acid libraries based on their affinity to a specific
target molecule.\textsuperscript{13,14} Pegaptanib was the first of its kind approved by Food and Drug Administration (FDA) of the United States for treating AMD.\textsuperscript{15} Since it exhibits affinity only to VEGF\textsubscript{165} and not to other VEGF isoforms, it is ideal for the treatment.\textsuperscript{15} Recently, another VEGF\textsubscript{165} binding aptamer patented in 1998 (ref. 16) was studied by considering its variants toward characterizing their relative binding affinities.\textsuperscript{17} The variants were created by extension and substitution of the original aptamer, and their relative affinities were evaluated based on kinetics and competitive binding analyses. The biophysical characterization of these interactions provides a testbed to check whether a more advanced docking scheme can reproduce experimental observables even for a tricky system with an undetermined structure like VEGF\textsubscript{165}. Of course, through the characterization, we can also access new insights toward designing new drug candidates against AMD.

Here, in this regard, we carry out docking simulations between VEGF\textsubscript{165} and the reported DNA aptamer variants to test whether the experimentally measured affinity order can be reproduced by an advanced docking scheme. Based on molecular dynamics (MD) simulations in conjunction with docking, we also address the puzzling issue that DNA aptamers bind only to HBD but still inhibit the binding between VEGF\textsubscript{165} and VEGF receptor (VEGFR), even though the binding takes place via the epitopes in RBD which are rather far from HBD. We expect that our scheme can be a useful tactic for handling flexible protein binding and that the additional findings will provide a clue in designing new inhibitors targeting VEGF\textsubscript{165}.

Results and discussion

VEGF\textsubscript{165} exists as a homodimer, and each monomer is composed of HBD and RBD that are connected by a flexible linker. When it binds to the extracellular domains D2 and D3 of VEGFR, it promotes the reorientation of the VEGFR dimer, leading to a conformation suitable for activating the receptor tyrosine kinase (RTK) in the intracellular domain (Fig. 1).\textsuperscript{18} As already discussed, the flexible nature of VEGF\textsubscript{165} has rendered the determination of its whole structure impossible. Especially, docking with the flexible HBD has not been carried out yet to the best of our knowledge. To shed light on this issue, by employing docking and MD simulations, we examined the binding between DNA aptamer variants and VEGF\textsubscript{165} and inspected how this binding impedes the interaction between VEGF and VEGFR. We used macromolecular docking program DOT2 (ref. 19 and 20) for all docking calculations because of its excellence in describing protein–DNA interactions. A very recent study has also employed DOT2 for dockings between transmembrane protein and RNA aptamers and succeeded in discovering aptamer variants with improved binding affinities.\textsuperscript{21}

We attempted docking calculations of aptamers with VEGF\textsubscript{165}, starting from the experimental results reported by Potty \textit{et al.}\textsuperscript{17} They characterized biophysical properties of a DNA aptamer 5’-CGGTCTTCCAGACAAGAGTGCAGGG-3’ (referred to as the “initial” aptamer hereafter) and its variants on their binding to VEGF\textsubscript{165}. The variants were synthesized by elongation (“0C” and “extended”) or point mutations (A14G, G16A, and G18A). Their schematic two-dimensional structures are shown in Fig. 2. They measured dissociation constants for the initial aptamer and its elongated variants, and obtained the relative binding affinities of the point mutants.\textsuperscript{17} The aptamer binding affinities to VEGF\textsubscript{165} were measured to be in the order of extended, 0C, initial, A14G, and G16A/G18A, with G16A and G18A exhibiting comparable affinities. We built the structures of these aptamer variants based on their sequences, and the details can be found in the last section (Computational details).

Because there is no experimental VEGF\textsubscript{165} structure, the relative pose of HBD with respect to RBD is not fixed at equilibrium and will actually keep moving. This also suggests that the corresponding potential energy minimum for the motion of HBD will be very shallow or even flat in reference to the thermal energy. Therefore, diverse VEGF\textsubscript{165} conformations should be allowed at equilibrium. Although there is no information...

![Fig. 1](https://example.com/fig1.png)  
**Fig. 1**  Schematic representation of the structural aspects of VEGF\textsubscript{165} and its complex with VEGFR.
regarding the whole VEGF\textsubscript{165} structure, RBD X-ray crystal structure and HBD NMR structure are available (PDB IDs: 2VPF\textsuperscript{23} and 1VGH,\textsuperscript{24} respectively). Because the NMR structure contains 20 conformations, the same number of structures for the whole VEGF\textsubscript{165} are possible when concatenating the two domains (Fig. S1 in ESI†). The six aptamers (Fig. 2) that we created earlier were docked to these twenty VEGF\textsubscript{165} structures. The best docking energies of these twenty VEGF\textsubscript{165} to each aptamer are presented in Fig. 3. The results show that the docking scores with respect to the twenty VEGF\textsubscript{165} structures are merely inconsistent with the experimental binding affinity order, extended > 0C > initial > A14G > G16A ≈ G18A.\textsuperscript{17} These poor agreements are likely due to the flexibility of the protein that we did not include in our computations. Please note that, instead of computing binding affinities with the binding energies based on free energy calculations, we used docking scores here as a metric for predicting the binding affinity order. This was because conventional free energy methods require extensive conformational sampling, which will be prohibitively costly due to the multiple pose binding nature between VEGF\textsubscript{165} and DNA aptamers. Namely, in more conventional docking, the best pose tends to be predicted because a specific binding site normally exists for each ligand, and a free energy calculation method can be subsequently applied to the best pose toward computing the binding affinity. However, we could not find such a binding site for the DNA aptamer in HBD of VEGF, as multiple pose binding occurred. Therefore, if a free energy calculation was pursued, it would have cost an immense amount of computational time. In fact, the multiple pose binding nature between HBD of VEGF and its ligand driven by electrostatic interactions has also
been reported previously.\textsuperscript{12} In addition, we note that binding affinities have been reasonably reproduced by docking scores without actual free energy calculations.\textsuperscript{25,26}

Thus, it is clear that we have to somehow include the flexible nature of the protein for our computations. As a matter of fact, properly considering various protein flexibilities has been a challenging task not just in free energy calculations but also in any computations including docking. Although the thermal fluctuations of the backbone and the surface residues can be taken into account to some degree by adopting ensemble docking,\textsuperscript{27,28} large scale motions such as HBD movement in VEGF\textsubscript{165} will be very expensive to sample statistically. To circumvent this issue, we carried out the anisotropic network model (ANM) analysis for the twenty concatenated VEGF\textsubscript{165} conformations created earlier. This will enable us to examine the principal movements based on the simplified normal mode analysis.\textsuperscript{29–31} Among the computed normal modes, we focused on the lowest frequency one because that will be the lowest energy vibration whose motion is energetically most favored. Furthermore, regardless of the conformation identified from the twenty possibilities used for the ANM analysis, the lowest frequency normal mode motion always appeared similar. In fact, the motion could be described as the scissoring bending vibration by the two HBD units, resulting in oscillations of the distance between the two units (Fig. 4A).

In order to reflect the effect of this lowest frequency normal mode in docking, we generated several different VEGF\textsubscript{165} conformations by varying the distance between the HBD units starting from the one that bears largely separated HBD units ("Structure 11" in Fig. S1†). For that, we carried out biased MD simulations with a restraint on the distance between the two units in the range of 30–90 Å at an interval of 10 Å, eventually extracting seven different VEGF conformations at as many inter-unit distances (Fig. 4B). In principle, computing a free energy profile along the inter-HBD distance can also be considered, but our system in combination with a large number of explicit water molecules is excessively large for conducting the required sampling. Instead, we then performed docking simulations with the six DNA aptamer variants to these seven VEGF\textsubscript{165} conformations, yielding a total of 42 docking calculations. The lowest energy from each docking and the average from the seven VEGF\textsubscript{165} conformations are plotted in Fig. 5A. Even after considering the VEGF\textsubscript{165} conformational flexibility from the lowest frequency normal mode, the predicted binding affinity order from docking simulations is still inconsistent with the experimental results. In addition, almost no correlations between the docking scores and the logarithms of the dissociation constants were found (Fig. 5B) for the three variants (initial, 0C, and extended) whose dissociation constants are available from experiments.

Fig. 4  (A) Lowest frequency normal mode obtained from ANM. (B) Seven VEGF\textsubscript{165} conformations generated from biased MD simulations.
Although we did not observe a satisfactory trend yet, meanwhile, we found that all DNA aptamer variants bound only to HBD in these 42 docking simulations. In fact, this binding specificity is in line with a previous experimental observation on a well-known DNA aptamer, Macugen. This correct aspect has motivated us to perform new docking simulations between the DNA aptamers and the HBD. Because the computational time for docking can be significantly reduced with this approach, at this time, we decided to carry out ensemble docking toward considering small-scale motions of the protein. Toward this end, we generated 50 snapshots of HBD using MD simulations. We then performed docking of the six DNA aptamers with the 50 HBD snapshots, obtaining 300 docking results in total. The energies computed from these dockings are shown in the leftmost part of Fig. 6A. Now the results reproduce the experimental binding affinity order well, except one slightly outlying mutant A14G. The correlation between docking scores and dissociation constants is also improved remarkably, although it is not yet quantitatively agreeing (Fig. 6B).

Encouraged by the fact that the docking with HBD ensemble considering small-scale motions showed noticeable improvements in predicting both binding affinity order and the energy correlation, we decided to further consider both the large and the small-scale motions simultaneously. To this end, 50 snapshots for all seven separations by the lowest frequency normal mode were taken into account, together with the six...
aptamers. This resulted in a total of 2100 docking results (i.e. 350 VEGF165 snapshots and six DNA aptamers). The results in Fig. 6A show that the averages from these are now in good agreement with the experimental binding affinity order except the slight affinity overestimation for A14G in comparison with the initial aptamer. Of course, the energy “resolution” of the docking result will not be good enough to discriminate the small affinity difference between the initial aptamer and A14G. Even still, the averages from 350 dockings at each distance (30 to 90 Å) predict significantly better binding affinity order compared to the single docking results shown in Fig. 3. This again substantiates the importance of ensemble docking. In addition, an almost perfect correlation between the docking scores and the experimental dissociation constants is observed at this time (Fig. 6C). Considering that this correlation is a large improvement over the ensemble docking results that involved only one HBD (Fig. 6B), we can infer that the whole structure of VEGF165 should be taken into account toward docking for a more reliable prediction of the binding affinity order. In fact, an earlier electrostatic potential isosurface analysis has shown that HBD is predominated by positive surface charges.12 Therefore, one HBD unit will likely influence the binding of a negatively charged DNA aptamer on the other HBD unit. This explains the necessity of adopting the whole VEGF165 for a reliable prediction. In addition, it is noteworthy that DNA aptamers bind to the HBD surface without any specificity, indicating that the binding is mainly driven by electrostatic interactions rather than shape complementarity. Such a multiple pose binding nature of HBD has been also reported previously against heparin ligand.12

Although we closely reproduced the binding affinity order between VEGF165 and DNA aptamers based on docking scores, there are still two issues to be clarified in order to substantiate the validity of our docking scheme. One is the validity of considering only lowest frequency normal mode in generating large-scale motions of VEGF165, and the other is the validity of examining only one initial structure (Structure 11 in Fig. 51†) among the twenty. To address the first issue, we additionally tested VEGF165 conformations based on the linear combination of six lowest frequency normal modes by using the NMSim web server.13,14 We generated 350 VEGF165 structures, which is the same number that we used for docking when considering only the lowest frequency normal mode. Since the number of VEGF structures for docking is identical, we can directly compare the results from our earlier docking scheme and this new one with NMSim. The docking results are shown in the leftmost part of Fig. 7A, together with the corresponding correlation given in Fig. 7B. Although we included as many as six normal modes, the results changed only marginally compared to the matching average from using one mode given in Fig. 6A. To address the second issue, we picked another VEGF165 structure (“Structure 17” in Fig. S1†), which is the most different from Structure 11 in terms of the root-mean-squared distance (RMSD). We then generated seven VEGF165 conformations along the distance between the HBD units (Fig. S2†). With these conformations, we repeated the same calculations as with Structure 11, resulting in

Fig. 7 (A) Average docking scores of the DNA aptamer variants to VEGF165 structures from NMSim and to additional structures derived from Structure 17, (B) the correlation between docking scores and the dissociation constants with NMSim, and (C) the same correlation for the docking results with Structure 17. Error bars represent statistical standard errors.
a total of 2100 docking scores (Fig. 7A and C). The results indeed show similar improvements in binding affinity prediction as with Structure 11. This surely indicates that our docking scheme will work well regardless of which initial structure we choose.

As described earlier, from our docking, we observed that all DNA aptamers interact with HBD only. An earlier study also reported this binding specificity of DNA aptamers. On the other hand, VEGFR binds to VEGF165 through its RBD, which is rather distant from HBD. Thus, the fact that aptamer binding on HBD can impede the VEGFR–VEGF165 binding may be puzzling. The answer to this puzzle is actually intriguing and may provide a new insight into the development of new VEGF/VEGFR antagonists to inhibit angiogenesis. This is also notable, as many researchers have tried designing new antagonist peptides by targeting on RBD. In order to address this, we performed MD simulations of the initial aptamer bound on VEGF165 without fixing any atoms in space. For VEGF165, we chose the conformation with the inter-HBD distance value of 30 Å as it would reflect the fact better that the negatively charged DNA aptamer interacts with both HBD units with positive surface charges. We performed the trajectory simulation until the RMSD from the initial structure converged (40 ns, Fig. S3†). As expected, during the simulation the DNA aptamer attached itself to the other HBD, and this additional binding caused the two HBD unit to twist somewhat. This twist in turn triggered one of the HBD units to strongly interact with the L1 loop of RBD (Fig. 8). Indeed, according to an X-ray crystal structure, the L1 loop of RBD is known to interact with the D3 domain of VEGFR1 when they bind together. Especially, the Glu44 residue of RBD was shown to interact with Gln263 of D3 in VEGFR1. Interestingly, in the complex of VEGF165 with DNA aptamer from the MD simulation, this Glu44 of RBD formed a strong hydrogen bond with Arg124 of HBD. Moreover, seven additional hydrogen bonds were observed between HBD and RBD besides the hydrogen bond of Glu44 with Arg124. Consequently, these strong interactions between HBD and RBD are expected to effectively block the interaction between RBD and VEGFR1.

Based on the equilibrated structure of the VEGF165–DNA aptamer complex obtained from the MD simulations, we further examined how much this binding spatially interferes with the interaction between VEGF165 and VEGFR. To this end, we adopted the X-ray crystal structure of a complex between RBD of VEGF165 and VEGFR1 (PDB ID: 5T89), and overlaid our structure of VEGF165–DNA aptamer complex on it. The overlay was performed after aligning the structures with respect to the RBD units from our simulation result and the crystallography data. For comparison, we also overlaid free VEGF165 with closed and opened HBD units, respectively represented by the conformations at 30 and 90 Å inter-HBD separations displayed in Fig. 4B. We focused our attention especially on the change in HBD–VEGFR overlap induced by the introduction of the DNA aptamer binding to VEGF165. The overlaid structures are shown in Fig. 9. With free VEGF165, there is no significant spatial overlaps observed regardless of how widely the HBD pair opens. The observed overlaps are indeed quite small such that they can be avoided with the natural protein flexibility. What is noteworthy here is that, although a previous computational study predicted that VEGF165 with closed HBDS would bind to VEGFR more preferably than VEGF165 with opened HBDS, our results are not in line with that prediction. On the contrary, in our results, VEGF165 with closed HBDS (Fig. 9A) exhibits more spatial overlap with VEGFR1 than the one with the opened pair (Fig. 9B). Because VEGFR2 was used for the modeling in that previous study, the inconsistency may have arisen from the
structural differences between VEGFR1 and VEGFR2. At least, the binding of VEGF165 with opened HBDS to VEGFR1 is not disfavored according to our results. In the case of the binding between the VEGF165–DNA aptamer complex and VEGFR1, the aptamer-bound HBD completely overlaps with the D3 domain of VEGFR1, showing far larger spatial clash compared to the case with the free VEGF165 (Fig. 9C). This again confirms that the native interaction between VEGF165 and VEGFR1 will be severely hindered when the DNA aptamer is bound to VEGF165.

Conclusions

Due to its direct involvement in angiogenesis, VEGF165 has been a main target of many designed molecules that aim for inhibiting angiogenesis. Nevertheless, the flexible nature of HBD has rendered various structure-based analyses of VEGF165 very difficult. To make a step forward in this challenging task, we examined the binding of DNA aptamers to HBD through docking and show how this binding can eventually impede the interaction between VEGF165 and VEGFR. We found that ensemble docking with considerations on the backbone and side chain flexibility is crucial for correctly predicting the binding affinity order among different aptamer variants. In addition to small-scale motions, we found that properly considering a large domain motion of VEGF165 as obtained with the ANM analysis is essential in docking toward correctly reproducing the correlation between docking scores and dissociation constants. Once a DNA aptamer binds to one of the HBD units in VEGF165, it subsequently attaches itself to the other HBD. This in turn causes a strong interaction of HBD with the L1 loop of RBD through a number of hydrogen bonds. Because the L1 loop of RBD is a binding site for VEGFR, this will eventually lead to the inhibition of the interaction between VEGF165 and VEGFR. The present work demonstrates that each of two flexible HBD units of VEGF165 affects the aptamer binding as well as the interaction between VEGF165 and VEGFR. This may provide an important design rule to devise new antagonist molecules against VEGF165 to control angiogenesis.

Computational details

DNA aptamer modeling

Because the 3D structure of the aptamer variants are not available, we modeled each structure based on the sequence information. For this, we referred to a previously suggested approach and used mfold web server to generate the 2D structures of the aptamers using their sequences as inputs. When we tried an alternative, RNA structure web server to see the robustness of the prediction by mfold, we obtained exactly the same 2D DNA structures. Predicting the 3D DNA structure from such 2D information is not practically possible, and we built an equivalent 3D structure of an RNA sequence using the RNA composer first and then subsequently converted it into DNA by manually modifying corresponding atoms. We also tried other alternatives in predicting the 3D RNA structures, namely, 3dRNA, SimRNA, and MC-Sym. However, they gave either energetically unstable structures or even abnormal structures with extremely elongated covalent bonds. Therefore, we decided to use RNA composer for this step. After this, we refined each structure by an energy minimization for 500 steps.
with the ff14SB AMBER force field parameter set and the generalized Born solvent model.

VEGF$_{165}$ modeling

In order to build the whole VEGF$_{165}$ structure, we used RBD taken from an X-ray crystal structure (PDB ID: 2VPF) and HBD from an NMR structure (PDB ID: 1VGH). Because two amino acids, Asp109 and Arg110, in a flexible linker connecting RBD and HBD are missing in both 2VPF and 1VGH, we first built a peptide chain Lys108–Asp109–Arg110–Ala111 using Avogadro. This chain was used as a bridge to connect Lys108 of RBD and Ala111 of HBD. Similarly to the DNA aptamer case, the concatenated VEGF$_{165}$ structure was further refined by 500 steps of minimization with the ff14SB force field and the generalized Born solvent model. The ANM analysis of VEGF$_{165}$ was performed using ANM web server 2.0.

MD simulations

The modeled structures were first solvated in a truncated octahedral TIP3P water box with the minimum distance of 10 Å from any atom in the solute to the edge of the water box. The solvated structures were subsequently neutralized by adding counter ions and minimized for 500 steps to avoid any possible steric clashes. The minimized structures were then gradually heated up to 300 K with positional restraints on the solute. These restraints were removed gradually, and the structures were equilibrated for 2 ns prior to any further calculations. Periodic boundary condition was employed for all MD simulations, and the cut-off distance of 8 Å was used for non-bonded interactions. Particle Mesh Ewald (PME) summation was employed for handling long range electrostatic interactions while the continuum model was adopted for treating long range van der Waals interactions. The time step of 2 fs was utilized, and the SHAKE algorithm was used to fix the bonds including hydrogen atoms. To keep the temperature constant during MD simulations, the Langevin thermostat was employed, and the pressure was kept at 1 bar by using Berendsen’s weak-coupling algorithm.

In order to generate seven discrete conformations of VEGF$_{165}$ along the distance between the two HBD units, we performed MD simulations with a biasing potential represented by a force constant of 40 kcal mol$^{-1}$ Å$^{-2}$. The 50 snapshots to carry out ensemble docking were extracted from each biased trajectory (10 ns) with the different inter-HBD distances. All MD simulations were carried out using AMBER 15.

Docking with DOT2

All dockings performed in this work used the DOT2 software. DOT2 is a docking program for rigid-body docking between two macromolecules. The notable strength of DOT2 is its capability of computing the electrostatic potential of molecules thoroughly based on Poisson–Boltzmann methods, so it is particularly superior in the estimation of electrostatic interactions between highly polar molecules such as protein–DNA binding. Due to this advantage, DOT2 has been employed for docking for various protein–DNA interactions, such as DNA–DNA repair enzymes, the nucleosome, and viral DNA-integrase. DOT2 calculates intermolecular energies as the sum of electrostatic and van der Waals (VDW) energy terms. We used the AMBER library (uhbd.amber84.prot.dna.rib) as the residue library to assign partial atomic charges for both VEGF$_{165}$ and DNA aptamer. The grid points with the spacing of 1 Å were generated for VEGF$_{165}$, and the electrostatic potential of VEGF$_{165}$ was computed for each grid by an outside program Adaptive Poisson–Boltzmann Solver (APBS) based on the linearized Poisson–Boltzmann equation. To compute the VDW interaction energies, the shape of VEGF$_{165}$ was determined first by its volume surrounded by a 3 Å thick layer using MSMS, and then each heavy atom of the DNA aptamer inside this layer was computed to contribute 0.1 kcal mol$^{-1}$ to the VDW interaction energy. At each grid, the interaction energy between VEGF$_{165}$ and the aptamer was calculated for 54 000 evenly spaced orientations (6$^2$ spacing) of the DNA aptamer under the potential of VEGF$_{165}$ at 300 K. During docking calculations, the atoms of the DNA aptamer were not allowed to spatially overlap with VEGF$_{165}$ within the distance of their VDW radii.

Conflicts of interest

There are no conflicts to declare.

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