Structure-based cross-docking analysis of antibody–antigen interactions

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Antibody–antigen interactions are critical to our immune response, and understanding the structure-based biophysical determinants for their binding specificity and affinity is of fundamental importance. We present a computational structure-based cross-docking study to test the identification of native antibody–antigen interaction pairs among cognate and non-cognate complexes. We picked a dataset of 17 antibody–antigen complexes of which 11 have both bound and unbound structures available, and we generated a representative ensemble of cognate and non-cognate complexes. Using the Rosetta interface score as a classifier, the cognate pair was the top-ranked model in 80% (14/17) of the antigen targets using bound monomer structures in docking, 35% (6/17) when using unbound, and 12% (2/17) when using the homology-modeled backbones to generate the complexes. Increasing rigid-body diversity of the models using RosettaDock’s local dock routine lowers the discrimination accuracy with the cognate antibody–antigen pair ranking in bound and unbound models but recovers additional top-ranked cognate complexes when using homology models. The study is the first structure-based cross-docking attempt aimed at distinguishing antibody–antigen binders from non-binders and demonstrates the challenges to address for the methods to be widely applicable to supplement high-throughput experimental antibody sequencing workflows.

Antibody–antigen interactions are an important component of our immune response to pathogens1, and understanding the structural basis of antibody–antigen interactions can help in designing more potent therapeutics. Recent experimental advances2 have enabled high-throughput sequencing studies of antibody repertoires in multiple organisms3–5, including perturbation effects of various antigens on the repertoire sequence frequencies6–8. Since antibody repertoire sizes for an individual range from 103 for zebrafish to 1010 for humans9, 10, computational approaches represent a practical structure-based method to study antibody specificity and selectivity. To understand the driving forces behind the generation and maturation of these antibody sequences, we need not just structural models, but knowledge of the antigen/epitope pairings and models of the bound antibody–antigen complexes. In this paper, we present a computational cross-docking study to discriminate binders from non-binders by identifying native antibody–antigen interaction pairs among cognate and non-cognate complexes. We also discuss the major remaining challenges for the methods to be useful to support high-throughput next-generation sequencing (NGS) pipelines or to support individual studies on particular antibody–antigen complexes of interest.

Cross-docking study efficacy relies on the accuracy of binding energy estimates. A team of researchers collected experimental binding affinity measurements for 179 protein–protein complexes as a benchmark dataset for training computational algorithms11, 12. Kastritis and Bonvin tested the affinity prediction ability of nine of the standard score functions used in the leading docking algorithms and found all score functions correlated poorly with the experimental results13. Some score functions were able to widely classify weak, medium and strong binders, but the standard deviations of the predicted binding energies were wider than the energy gap between the three categories. The ability to distinguish binding from non-binding interfaces was also tested in Critical Assessment of PRediction of Interactions (CAPRI) through a challenge to predict successful high-affinity binders from a set of designed protein–protein interfaces and distinguishing natural interfaces from unsuccessful Rosetta-designed interfaces14. Both these challenges were difficult, and no computational method was able to identify the design responsible for the successful binder.

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In this paper, we present a cross-docking study to discriminate antibody–antigen binders from non-binders using a dataset of 17 complexes with both bound and unbound antigen structures available. We first generate a dataset of cognate and non-cognate models for each antibody–antigen pair (see Methods for details).

We superposed each antibody onto the native antibody in 16 other antibody–antigen complexes, generating 289 total pairs to discriminate. The tightly packed interface residues in the native crystal complexes and the steric clashes in the non-cognate complexes generated using antibody superposition make the initial set trivial to separate. To construct a more realistic set, it is necessary to refine each candidate complex to (1) erase the memory of the crystal structure in cognate complexes, and (2) optimize the interface in non-cognate complexes. Furthermore, generation of an unbiased set of starting models is critical if the method is to be extendable to antibodies with no available experimental structures with the corresponding antigens. We used a customized version of the fixed-backbone RosettaDock high-resolution stage ("local refine") to generate an ensemble of 50 cognate and non-cognate refined models for each antibody–antigen pair (see Methods for details).

As shown in Fig. 1, the cognate models are refined starting from the crystal complex, while the non-cognate models are refined after superimposing the $V_L-V_H$ framework of the non-cognate antibody on the cognate antibody in the crystal complex. The starting antibody ligand-RMSDs between cognate and non-cognate antibodies average around 1.2–1.4 Å (Supplementary Fig. S1). The primary goal is to predict the correct antibody that pairs with each antigen. To test the effects of antibody backbone accuracy on the binding predictions, we generated antibody and non-cognate models using bound–bound (henceforth referred to as simply "bound"), unbound–unbound ("unbound"), and homology–unbound ("homology") backbones, respectively, for the antibody–antigen

| No. | Antibody | Antigen          | Complex PDB | Antibody PDB | Antigen PDB | Exp. Kd (nM) |
|-----|----------|------------------|-------------|--------------|-------------|--------------|
| 1   | Fab B02C11 | Factor VIII domain C2 | 1HQP | 1HQC* | 1D7P | <0.014 |
| 2   | Fab N10   | Staphylococcal nuclease | 1JN5 | 1JN6* | 1KD6 | <0.10 |
| 3   | Fab D3H44 | Tissue factor | 1JPS | 1JPT | 1TT8 | 0.10 |
| 4   | FabF10.6.6 | HWE lysozyme | 1P2C | 2Q76 | 3JLT | 0.10 |
| 5   | Fab BV16  | Birch pollen antigen Bet V1 | 1FSK | 1FSK* | 1BV1 | 0.24 |
| 6   | Fab E8   | Cytochrome C | 1WEJ | 1QBL | 1HRC | 0.71 |
| 7   | Fab HC19  | Flu virus hemagglutinin | 2VIR | 1GIG | 2HMG | 1.00 |
| 8   | Fab Hyhel63 | HWE lysozyme | 1DQF | 1DQQ | 3JLT | 2.80 |
| 9   | Fab Jd42  | HPf | 2JEL | 2JEL* | 1POH | 2.80 |
| 10  | Fab 5p9   | Tissue factor | 1AHW | 1FGN | 1TFH | 3.40 |
| 11  | Fab A4.6.1 | VEGF | 1MJ1 | 1BJ1* | 2VPF | 3.40 |
| 12  | Fv DL3   | HWE lysozyme | 1VFR | 1VFA | 8LYZ | 3.70 |
| 13  | Fab NC41  | Flu virus neuraminidase N9 | INCA | INCA* | 2NN9 | 8.30 |
| 14  | Fv Hdyx11 | HWE lysozyme | 1BVK | 1BVL | 3JLT | 14 |
| 15  | Fab 13B5  | HIV-1 capsid protein p24 | 1E6f | 1E6O | 1A43 | 29 |
| 16  | Fab 44.1  | HWE lysozyme | 1MLC | 1MLB | 3JLT | 91 |
| 17  | Fab HC19  | Flu virus hemagglutinin T131I mutant | 2VIS | 1GIG | 2VIU | 4000 |

Table 1. Antibody–antigen complexes used for cross-docking. *In these six cases, the only antibody structure available is in the antigen-bound state.
complexes. About 35% (6/17) of the antibodies in the dataset lack unbound structures, so we used bound anti-
body backbones to generate their unbound cognate and non-cognate complex models. We used RosettaAntibody
3.018 to generate the antibody models required for homology complexes (see Methods). Based on CDR RMSDs,
most of the generated antibody models are approximately equidistant (1–3 Å) from the known bound and
unbound conformations (Supplementary Fig. S2).

Sample binding analyses: Fab NC41–influenza virus neuraminidase N9 and Fab 44.1–hen
egg-white lysozyme complexes. The main metrics of interest are the rank of the cognate complex relative
to the non-cognate complexes and the magnitude of the predicted binding scores. We used Rosetta's docking
interface score19 (based on the Talaris2013 force field20) to rank the complexes. Rosetta interface score is defined
as \( I_{sc} = E_{\text{bound}} - E_{\text{unbound}} \), where \( E_{\text{bound}} \) is the score of the bound complex and \( E_{\text{unbound}} \) is the sum of the scores
of the individual protein partners in isolation. The score function is a linear combination of several score terms
including a Lennard–Jones potential, an implicit solvation potential21, an orientation-dependent hydrogen bond-
ing potential22, a Coulomb electrostatic potential with a distance-dependent dielectric, a side-chain torsional
potential23, and a knowledge-based residue pair term based on the probability of proximity of two amino acids in
the PDB24.

Figure 2 compares the interface scores for antibody complexes involving influenza virus neuraminidase N9
(Cognate Fab NC41; 1NCA) and hen egg-white lysozyme (Cognate Fab 44.1; 1MLC) generated using the local
refine routine (Fig. 2a,b). Interface scores for the cognate and non-cognate complexes (turquoise and grey, respec-
tively; top-scoring complexes for each antibody against the target antigen are highlighted with a black outline)
generated starting from bound, unbound, and homology backbones are plotted as a function of the complex
interface size (ΔSASA in Å²). In general, interface scores are lower (better) for larger candidate interfaces, and
the cognate antibody–antigen complexes have both larger interfaces and lower scores. In both the antigens, the
calculated energy gap between the cognate and non-cognate antibodies is small, even though the non-cognate
complexes likely bind weakly or not at all.

In influenza virus neuraminidase N9, interface scores of the top-scoring cognate complexes (Fig. 2a) drop
from −9.7 to −9.2, and further to −6.2 REU when using bound antibody/antigen, unbound antigen/bound
antibody, and unbound antigen/homology antibody backbones, respectively, even though all the complexes
involve the same epitope (Fig. 2c). The cognate scores remain better than the corresponding interface scores
of all the top-scoring non-cognate complexes in each category (−5.5, −5.3, and −5.8 REU, respectively). In
hen egg-white lysozyme, interface scores of the top-scoring cognate complexes (Fig. 2b) are −7.7, −5.1, and
−5.1 REU when using antibody bound, unbound, and homology backbones against the same epitope (Fig. 2d).
Interface score can discriminate the cognate antibody using bound models (top non-cognate score: −7.2 REU),
but other non-cognate antibodies surpass even the top-scoring cognate antibody when using unbound and
homology models. For example, when using unbound backbones, the top-scoring non-cognate complex gener-
ated using the antibody from the Fab HC19–influenza hemagglutinin T131I complex (2VIS) scores better (−5.9
REU) than the top-scoring cognate complex (−5.1 REU) even though both antibodies target the same lysozyme
epitope (Fig. 2e). Finally, even the top-scoring bound cognate interface scores of −9.7 and −7.7 REU in the two
complexes underestimate the true binding affinities (~11 and 9.6 kcal/mol, respectively, mapping $^{13, 25, 26}$ REUs to kcal/mol).

In summary, interface score distinguishes the cognate antibody from the non-cognate antibodies in both the cases when using the bound antibody coordinates but fails in the case of Fab 44.1–hen egg-white lysozyme complexes.
complex when using unbound and homology-modeled backbones, likely due to the decrease in the accuracy of the antibody backbone conformations used for generating the models.

Antibody backbone accuracy is critical for accurate binding partner identification. We used interface scores (Isc) to rank the cognate and non-cognate models for each antibody–antigen pair in the dataset. Figure 3 shows the interface scores of the top-scoring cognate and non-cognate models for each antigen. Using bound backbones (Fig. 3a), the cognate antibody–antigen pair is the top-scoring model in 14/17 antigens, and it is one of the top three scoring models in 16/17 antigens. Using unbound backbones, the cognate pairs ranked at the top in six targets, and in the top three in ten targets (Fig. 3b), compared to two and eight cognate pairs at the top and top three when using homology backbones (Fig. 3c). Four of the six top-ranked cognate pairs using unbound backbones are from targets missing unbound antibody structures and were generated using bound antibody coordinates. The drop in the prediction accuracy correlates with the drop in the accuracy of the backbone coordinates moving from bound to unbound, and eventually homology complexes. The decline in the discrimination capability is not surprising as backbone conformation is a critical determinant of protein–protein docking accuracy. Inaccuracies in the backbone often translate to docking errors through sub-optimal arrangement of the interface residues in the docking models.

To assess the interface quality of the top-scoring cognate and non-cognate models, we first calculated the distribution of the number of antibody–antigen interface hydrogen bonds. As shown in Fig. 4a, the native crystal
complexes contain $7.7 \pm 3.3$ interface hydrogen bonds, compared to $6.9 \pm 2.4$ and $1.8 \pm 1.7$ hydrogen bonds in the top-scoring cognate and non-cognate models when using bound backbones. The average number of interface hydrogen bonds drop to $3 \pm 1.5$ and $1.7 \pm 1.4$ for the top-scoring cognate and non-cognate models using unbound backbones, respectively, and finally $2.4 \pm 1.5$ and $2.2 \pm 1.5$ with homology-modeled antibody backbones. The number of interface hydrogen bonds in the top-scoring cognate models thus decline moving from bound to unbound, and homology complexes, but are similar for the top-scoring non-cognate models across the three categories. As the number of interface hydrogen bonds in cognate complexes drop, there is a greater chance the resulting buried unsatisfied polar atoms contribute to unfavorable interfaces affecting the discrimination power of the interface score.

We computed the number of buried unsatisfied polar atoms (ignoring contribution of structural waters) across the interfaces of the top-scoring cognate and non-cognate models. As shown in Fig. 4b, on average, crystal structures of the antibody–antigen complexes have $10.1 \pm 4.3$ buried unsatisfied polar atoms at the interface, with the top-scoring cognate and non-cognate bound models containing $10.5 \pm 4.2$ and $7.9 \pm 4.2$ buried unsatisfied interface polar atoms, respectively. The number of buried unsatisfied interface polar atoms in the top-scoring cognate and non-cognate complexes remain about the same at $11.2 \pm 5.7$ and $7.4 \pm 4.1$ when using unbound backbones,
but fall to $8.5 \pm 3.7, 6.7 \pm 3.6$, respectively, with homology-modeled antibody backbones. Surprisingly, the number of buried unsatisfied interface polar atoms in the top-scoring cognate complexes are always higher than the corresponding non-cognate complexes in each category. In addition, the reported drop in the number of interface hydrogen bonds between bound and unbound backbones does not increase the number of buried unsatisfied interface polar atoms (which remain about the same). Finally, the number of buried unsatisfied interface polar atoms drop from unbound to homology models correlating with a drop in the number of hydrogen bonds, more so in the top-scoring cognate models. One reason for this unexpected behavior is a simultaneous decline in the absolute interface sizes between i) bound, unbound, and homology, and ii) cognate and non-cognate docking models.

Finally, we investigated the interface sizes of the generated antibody–antigen complexes. Figure 4c shows the surface area change upon binding ($\Delta$SASA) in the top-scoring cognate and non-cognate models. The average $\Delta$SASA in the crystal complexes is $1622 \pm 290 \text{ Å}^2$, and it is $1670 \pm 299 \text{ Å}^2$ and $1212 \pm 317 \text{ Å}^2$ in cognate and non-cognate models, respectively, generated using bound backbone coordinates. $\Delta$SASA in the cognate and non-cognate models is $1467 \pm 321 \text{ Å}^2$ and $1187 \pm 332 \text{ Å}^2$ with unbound backbones, and finally, $1351 \pm 337 \text{ Å}^2$ and $1148 \pm 322 \text{ Å}^2$ using homology backbones. The interface sizes in the top-scoring cognate models decrease using bound, unbound, and homology backbones, but remain roughly the same in the top-scoring non-cognate models. This difference in the interface size trends between cognate and non-cognate models explains the drop in the number of interface hydrogen bonds across cognate but not non-cognate models, and the sharp drop in the number of buried unsatisfied polar atoms in the top-scoring cognate models.

In summary, the interface quality of the top-scoring cognate models declines when using bound, unbound and eventually homology backbones to model the complexes, while the interface quality of the top-scoring non-cognate models expectedly remains about the same (see Table 2 for summary). Since the structures generated by local refine comprehensively sample the antibody–antigen orientation space around the starting epitope, the decline in the prediction accuracy is primarily due to the inaccuracies in the backbone conformations in unbound and homology models.

### Expanding antibody–antigen model diversity reduces binding discrimination

The cognate and non-cognate models generated by the local refine protocol are typically under 5 Å C$_\alpha$ Irmsd from the native antibody–antigen complexes. That is, model diversity is limited because most model structures have interfaces around the epitope of the antigen. The limited diversity can hinder generation of native-like complexes with tightly-packed interfaces in non-cognate complexes or in cognate complexes where the homology models are not very accurate. To address this, we increased rigid-body sampling diversity using the standard RosettaDock local dock routine. Using local dock, we generated 1000 models starting from a random structure picked from the local refine ensemble for each antibody–antigen pair. In local dock, the starting structures are perturbed by about 3 Å translation and 8° rotation around the axis joining the centers of the two partners generating models up to 20 Å C$_\alpha$ Irmsd from the starting complex.

We compared the interface scores of the top-scoring cognate and non-cognate models for each antigen (Fig. 5). Using interface scores, the cognate antibody–antigen pair is the top-scoring model in nine of the 17 target antigens, and is one of the top three scoring models in 13/17 targets using bound backbones to generate the complexes (Fig. 5a). The cognate pairs ranked at the top in four targets, and in the top three in six targets using both unbound and homology models (Fig. 5b,c). Two of the four top-ranked cognate pairs in unbound models are from targets missing unbound antibody structures and thus generated using bound antibody coordinates. As expected, the number of correctly identified cognate antibody–antigen pairs using local dock is lower than local refine when using bound, unbound backbone coordinates. Encouragingly, when using homology complexes, local dock places more cognate pairs at the top (four) compared to local refine (two).

We again calculated distributions of the three interface quality metrics: (i) number of interface hydrogen bonds, (ii) number of buried unsatisfied interface polar atoms, and (iii) surface area change upon binding for the top-scoring cognate and non-cognate models generated by local dock (see Table 2 for summary). The top-scoring cognate models generated starting from bound backbones contain $6 \pm 3$ interface hydrogen bonds, $11.7 \pm 3.7$ buried unsatisfied interface polar atoms, and $1636 \pm 290 \text{ Å}^2$ $\Delta$SASA, compared to $3.2 \pm 1.8$ hydrogen bonds, $8.4 \pm 3.7$ unsatisfied atoms, $1309 \pm 258 \text{ Å}^2$ $\Delta$SASA, respectively, in the top-scoring non-cognate models (Fig. 6).

Using unbound backbones, cognate and non-cognate models contain $3.1 \pm 2.1$ hydrogen bonds, $9.2 \pm 4$ unsatisfied atoms, $1394 \pm 316 \text{ Å}^2$ $\Delta$SASA, and $3.4 \pm 1.8$ hydrogen bonds, $8.3 \pm 3.5$ unsatisfied atoms, $1328 \pm 251 \text{ Å}^2$ $\Delta$SASA, respectively. Finally, the top-scoring models cognate and non-cognate models generated using homology backbones contain $3 \pm 1.6$ hydrogen bonds, $8.4 \pm 3.5$ unsatisfied atoms, $1332 \pm 213 \text{ Å}^2$ $\Delta$SASA, and $3.2 \pm 1.9$
hydrogen bonds, 7.8 ± 3.3 unsatisfied atoms, 1292 ± 265 Å² ΔSASA, respectively. The interface quality of the cognate models generated by local dock drops drastically moving from bound backbones, but is similar across models generated using unbound and homology backbones correlating with the same trend observed in the binding discrimination performance. For the non-cognate models, the average interface quality is similar across all the three categories.

In general, the binding discrimination performance in local dock generated models is lower compared to the models generated using local refine because the greater model diversity increases the chance of false positive interfaces picked up by the score function. One exception is models generated from homology-modeled antibody backbones where the greater antibody–antigen orientation diversity in local dock compensates for the backbone inaccuracies resulting in higher quality interfaces in both cognate and non-cognate models. Local dock achieves more top-ranked cognate complexes compared to local refine using homology-modeled antibodies.

Effect of experimental binding affinities and antibody backbone modeling errors. To determine the effect of absolute binding affinities of the antibody–antigen complexes on the binder predictions, we plotted correlations of the ranks of the top-scoring cognate models vs. the absolute experimental ΔG values of the native antibody–antigen complexes. As shown in Fig. 7a, binder discrimination of the models generated using local refine and local dock is independent of the absolute binding affinities for bound, unbound, and homology
complexes. For example, the cognate antibody against Factor VIII domain C2, Fab B02C11 (PDB: 1IQD) which is the stronger binder in the dataset (experimental $K_d < 0.014 \text{nM}$) ranks first both in local refine and local dock against its antigen when using bound backbone, but ranks fifth and eleventh, respectively, when using unbound backbone, and third and first when using homology-modeled backbone. The weakest binder in the dataset, Fab HC19 against influenza virus hemagglutinin T131I mutant (2VIS), ranks second both in local refine and local dock when using bound, but ranks third and first when using unbound, and eighth and fourteenth when using homology-modeled backbones. Therefore, when using unbound backbones, prediction accuracy is higher in the case of the weakest binder compared to the strongest binder in the dataset. It is not surprising binding discrimination is not strongly dependent on absolute binding affinities of the native antibody–antigen complexes, as the score function is not calibrated for absolute binding affinities but to distinguish native vs. non-native interfaces.

We next tested the dependence of the ranks of the cognate complexes on their interface scores (Fig. 7b). The cognate ranks are good (rank 1 or 1–3) when interface scores $I_{sc} < -6 \text{ REU}$ (i.e. tight binders), and cognate rank quality decreases (higher ranks) for $I_{sc}$ above $-6 \text{ REU}$. The increase in the cognate rank is especially evident in unbound and homology complexes where only three complexes from the dataset have $I_{sc} < -6 \text{ REU}$, compared to bound complexes where 76%, 88% of the complexes have $I_{sc} < -6 \text{ REU}$ when using local refine, lock dock,
respectively. Since interface score is a measure of the complex interface quality, the increase in the cognate rank highlights the drop in the interface quality when using unbound and homology backbones.

We also calculated the effect of antibody backbone accuracy on binding discrimination by plotting correlations of the ranks of the top-scoring cognate complexes vs. RMSD of the antibody CDRs from the bound backbone coordinates (Fig. 7c). As expected, binder discrimination accuracy in general drops at higher CDR RMSDs. Specifically, when using local refine with unbound and homology-modeled backbones, the cognate rank progressively worsens with increasing antibody CDR RMSD. The weak correlation with the antibody backbone accuracy in local refine but not local dock hints that the limited antibody–antigen orientation diversity in local refine is inadequate to compensate for the backbone errors. This observation is consistent with the small increase in the number of top-scoring cognate models when using local dock (vs. local refine) in models generated using homology-modeled backbones.

Finally, we evaluated the impact of the deviation of the final top-scoring cognate and non-cognate complexes from the original cognate crystal complexes on binding discrimination for the local refine and local dock cases. Supplementary Fig. 3 shows the antibody ligand-RMSD (Ab_L_RMSD, see Methods) versus the rank of all cognate and non-cognate top-scoring models for all 17 targets and various docking cases. Like the CDR RMSDs,
it is necessary for the antibody ligand-RMSD to be small for the cognate antibody to have a top rank relative to the non-cognate antibodies. Conversely, top-ranked non-cognate antibody complexes are sometimes close to the cognate complex structure (Ab_L_RMSD < 5 Å), and sometimes quite different (up to ~30 Å).

Binder discrimination summary: Antibody–antigen complexes are challenging targets. To compare the binding discrimination power of the interface score across various categories, we calculated the Receiver Operating Characteristic (ROC) curves using the interface score of the top-scoring model as the classifier in bound, unbound, and homology complexes for both local refine and local dock routines (Fig. 8). We also calculated the ROC curves for the complexes using four other widely-used scoring potentials computed using the CCharPPI 28 web server: DFIRE2 29 interaction energy, total FireDock 30 energy (antibody–antigen energy function), OPUS-PSP 31 all-atom potential, and ZRANK2 32 scoring function. We considered the native cognate antibody–antigen interaction pairs as the only true positives.

In ROC curves, the area under the curve (AUC) values represent the performance of the scoring potential as a binary binder vs. non-binder classifier. Using Rosetta interface score as the classifier for models generated using the local refine routine, the AUC for the bound complexes is 0.97, followed by 0.78 and 0.73 for unbound and homology complexes, respectively (AUC 0.5 = random, 1 = perfect discrimination, Fig. 8a). For local refine models, Rosetta interface score is the best discriminator in bound complexes, but FireDock and OPUS-PSP potential perform better when using unbound and homology complexes. Using interface score as the classifier in local dock routine, the AUC values for bound, unbound, and homology complexes are 0.87, 0.65, and 0.59, respectively (Fig. 8b). For local dock models, Rosetta interface score is one of the top two discriminators in bound and unbound complexes, behind OPUS-PSP and ZRANK2 potentials, respectively. In homology complexes, FireDock outperforms all the other potentials including the Rosetta interface score. Overall, discrimination by the interface score is better than random in all cases, and as expected, drops with an increase in antibody backbone inaccuracy and/or model diversity.

Discussion
We have presented the first structure-based cross-docking study focused on discrimination of protein binders from non-binders by identifying native antibody–antigen interaction pairs among cognate and non-cognate
complexes. After generating an unbiased dataset of cognate and non-cognate complexes, we used Rosetta interface scores of the top-scoring models to rank the complexes. The binding discrimination results are encouraging when using bound backbones with the cognate pair ranked at the top in more than 80% of the antigen targets, compared to 35% and 12% when using unbound and homology-modeled antibody backbones, respectively. Using RosettaDock’s local dock routine to increase antibody–antigen model diversity helps achieve more top-ranked cognate complexes when using homology-modeled antibody backbones, but decreases binding prediction accuracy in bound and unbound backbone-generated models because of the higher false positive rate.

Interface quality of the cognate models declines between using bound and unbound, homology-modeled antibody backbones to model the complexes, but remains about the same in non-cognate models. The drop in the cognate interface quality and hence binding discrimination highlight the sampling limitations of the current docking algorithms when using unbound and homology modeled backbones. Therefore, accurate modeling of the bound structures starting from the unbound structures or homology models is critical for improving accuracy. Flexible backbone algorithms such as EnsembleDock39 and SnugDock35 based on the conformer selection (CS) binding models previously shown to help improve interface quality40 of the models generated using unbound and homology complexes can be of immense value. However, existing flexible docking algorithms may increase false positive rates as the additional backbone flexibility will also help accommodate the energetically unfavorable interface residues at the epitopes in non-cognate models. Moreover, recent results27 show that none of the current backbone flexibility generation algorithms successfully sample the bound conformation in a sizable fraction of proteins. Therefore, improving prediction accuracy will require development of novel flexible backbone sampling approaches to model the bound conformation with score functions specially catered to antibody binder discrimination.

Antibody–antigen cross-docking is the only in silico approach that offers a structural perspective in epitope-targeted antibody screening studies. It is useful to screen antibody binders raised from animal immunization campaigns to pick an epitope-diverse selection of antibodies increasing the chances of in vivo efficacy that is often a complex function of various interactions involving the antigen. With the rapid rise in the scale of the computing resources, computational structure-based methods are already playing a role in understanding the biological process of generation of antibody repertoires and how antibodies are selected for recognizing pathogens6. Further improvements in binding predictions with unbound and homology-modeled backbones will pave the way for studies parsing antibody sequence repertoires for antigen-specific binders and rational engineering of antibodies to minimize off-target activity. This study is the first step towards development of an efficient structure-based cross-docking framework to support high-throughput experimental antibody pipelines.

Methods

Homology modeling. Rosetta’s antibody homology modeling protocol, RosettaAntibody 3.018 is used for constructing homology models for the antibodies in the dataset. V_L,V_H homologs with more than 80% sequence identity, and CDR homologs with more than 98% sequence identity are excluded from the database during the template selection stage. Since the antibodies used in the study are part of the RosettaAntibody database, filtering is necessary to avoid picking structural components from the native structures during modeling. Additionally, the sequence identity cutoffs help simulate homology modeling of newly-determined antibody sequences that do not have existing highly homologous structural templates. The coordinate files of the 17 homology-modeled antibodies are provided as Supplementary Dataset 1.

Starting complex structure generation. For bound complexes, the crystal structure of the antibody–antigen complex is used as the starting structure. For unbound complexes, the starting structure is generated by superimposing the antibody and antigen unbound structures on the bound crystal complex, and for homology complexes, the antibody homology model and antigen unbound structure on the bound crystal complex.

Local refine. The local refine routine is derived from RosettaDock’s high-resolution refinement stage. Since superposing the V_L,V_H framework of the non-cognate antibody on the cognate antibody to generate the starting non-cognate structure often results in steric clashes with the antigen, the standard refinement routine moves the antibody away from the epitope to lower the total score of the complex. To generate models focused around the epitope, we created a customized local refine antibody–antigen refinement routine. The routine starts minimization of the antibody–antigen orientation with a score function with a low Van der Waals repulsive weight (w_rep = 0.02) to ignore starting steric clashes. The weight is gradually ramped up to its full value (w_rep = 0.186) in increments of 0.33 each over three minimization cycles allowing sampling of conformations that are otherwise inaccessible due to the high-energy barriers.

Local dock. We used the standard RosettaDock local docking routine35 involving a sequence of low-resolution and high-resolution steps to generate local dock models.

Metrics. Antibody ligand-RMSD (Ab_L_RMSD) of a model antibody–antigen complex is defined as the RMSD between the antibody framework Cα atoms (receptor) in the model and reference complexes calculated after aligning the antigen backbones (ligand) in the both the complexes.

\[
\text{Ab}_L\_\text{RMSD} = \sqrt{\frac{\sum_{\text{C}_\alpha} \left(\text{C}_\alpha^{\text{Ab}_L\_\text{Frw}} - \text{C}_\alpha^{\text{Ref}\_\text{Ab}_L\_\text{Frw}}\right)^2}{N_{\text{Ab}_L\_\text{Frw}}}}
\]
where \( C_{\alpha C} \), \( C_{\alpha AbFrwk} \), \( C_{\alpha RefAbFrwk} \) are the \( C_{\alpha} \)-atom coordinates of the antibody framework residues in the model and reference antibodies, respectively. The RMSDs are calculated over all the antibody framework residues \( (N_{AbFrwk} \cdot \rangle) \).

Antibody ligand-RMSDs are analogous to the commonly used ligand RMSDs in docking calculations.

**Rosetta command line.** The algorithms in the manuscript are implemented using the Rosetta molecular modeling suite. The Rosetta command-line arguments used for the calculations are as follows:

**Cognate and non-cognate model generation.**

(a) Local refine

\[ \text{ab\_binding.<exe>} \]

\[ -s \ 1BVK.1AHW.pdb \]

\[ -\text{docking:partners LH\_C} \]

\[ -\text{docking:docking\_local\_refine} \]

\[ -\text{pH:pre\_process} \]

\[ -\text{pH:cognate\_pdb 1AHW.b.pdb} \]

\[ -\text{pH:cognate\_partners LH\_C} \]

\[ -\text{ex1} –\text{ex2aro} -\text{use\_input\_sc} \]

\[ -\text{nstruct} \ 50 \]

where \( 1BVK.1AHW.pdb \) is the structure generated after superposing the non-cognate antibody from \( 1BVK.pdb \) on the cognate antibody in \( 1AHW.pdb \). The –docking:partners and –pH:cognate_partners arguments identify the receptor and ligand chains in non-cognate and cognate complexes, respectively. –pH:cognate_pdb provides the cognate bound complex.

(b) Local dock

\[ \text{docking\_protocol.<exe>} \]

\[ -s \ 1BVK.1AHW.pdb -native 1BVK.1AHW.pdb \]

\[ -\text{dock\_pert 3 8 –spin –partners LH\_C} -\text{ex1} –\text{ex2aro} \]

\[ -\text{nstruct} \ 1000 \]

**Homology modeling.**

(a) RosettaAntibody homology modeling

\[ \text{antibody\_H3.<exe>} \]

\[ -s \ 1AHW.h.pdb \]

\[ -\text{antibody:remodel perturb\_kic} \]

\[ -\text{antibody:snugfit true} \]

\[ -\text{antibody:refine refine\_kic} \]

\[ -\text{antibody:cter\_insert false} \]

\[ -\text{antibody:flank\_residue\_min true} \]

\[ -\text{antibody:bad\_nter false} \]

\[ -\text{antibody:h3\_filter false} \]

\[ -\text{antibody:h3\_filter\_tolerance 5} \]

\[ -\text{antibody:constrain\_cter} \]

\[ -\text{antibody:constrain\_vlvh\_qq} \]

\[ -\text{constraints:cte\_file 1AHW\_cter\_constraint} \]

\[ -\text{loops:legacy\_kic false} \]

\[ -\text{loops:kic\_min\_after\_repack true} \]

\[ -\text{loops:kic\_omega\_sampling} \]

\[ -\text{loops:allow\_omega\_move true} \]

\[ -\text{kic\_bump\_overlap\_factor 0.36} \]

\[ -\text{loops:ramp\_fa\_rep -loops:ramp\_rama} \]

\[ -\text{loops:outer\_cycles 5} \]

\[ -\text{corrections:score:use\_bicubic\_interpolation false} \]

\[ -\text{ex1} –\text{ex2aro} -\text{extrachi\_cutoff 0} \]

\[ -\text{nstruct} \ 2000 \]

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Author Contributions
Conceived and designed the experiments: K.P.K. and J.J.G. Performed the experiments: K.P.K. Analyzed the data: K.P.K. Wrote the manuscript: K.P.K. and J.J.G.

Additional Information
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