Broad epitope coverage of a human in vitro antibody library

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ABSTRACT
Successful discovery of therapeutic antibodies hinges on the identification of appropriate affinity binders targeting a diversity of molecular epitopes presented by the antigen. Antibody campaigns that yield such broad “epitope coverage” increase the likelihood of identifying candidates with the desired biological functions. Accordingly, epitope binning assays are employed in the early discovery stages to partition antibodies into epitope families or “bins” and prioritize leads for further characterization and optimization. The collaborative program described here, which used hen egg white lysozyme (HEL) as a model antigen, combined 3 key capabilities: 1) access to a diverse panel of antibodies selected from a human in vitro antibody library; 2) application of state-of-the-art high-throughput epitope binning; and 3) analysis and interpretation of the epitope binning data with reference to an exhaustive set of published antibody:HEL co-crystal structures. Binning experiments on a large merged panel of antibodies containing clones from the library and the literature revealed that the inferred epitopes for the library clones overlapped with, and extended beyond, the known structural epitopes. Our analysis revealed that nearly the entire solvent-exposed surface of HEL is antigenic, as has been proposed for protein antigens in general. The data further demonstrated that synthetic antibody repertoires provide as wide epitope coverage as those obtained from animal immunizations. The work highlights molecular insights contributed by increasingly higher-throughput binning methods and their broad utility to guide the discovery of therapeutic antibodies representing a diverse set of functional epitopes.

Introduction

The biopharmaceutical industry invests heavily in the discovery and clinical development of monoclonal antibodies (herein referred to as “antibodies”), which generates demand for higher throughput analytical tools that can meet this capacity. An antibody’s epitope specificity largely dictates its biological function, and, since it is an innate property that cannot be optimized rationally by engineering, it must be selected properly. In early-stage therapeutic antibody discovery, the aim is often to identify clones that target a broad range of epitopes on the specific antigen of interest and prioritize those with the desired biological function. The vast genetic diversity of state-of-the-art antibody libraries and optimized selection methods enable the discovery of functionally effective and mechanistically differentiated binders. Sourcing antibodies from both in vivo and in vitro platforms may maximize epitope diversity opportunities, given their complementary strengths and limitations.

High-throughput antibody generation methods must be used with advanced analytical methods to efficiently characterize their outputs and enable a more informed selection of antibodies with therapeutically relevant epitopes. High-throughput epitope binning methods on label-free biosensors are particularly useful in this regard because the results from these experiments allow antibodies to be sorted into epitope “families” or “bins” based upon their ability to cross-block one another’s binding to their antigen in a combinatorial and pairwise fashion. These methods can also discern fine specificities that can be used to discriminate antibodies with near-identical epitopes, which may manifest as functional differences, or offer intellectual property opportunities.

Here, we describe a collaborative effort in which we applied state-of-the-art label-free analytical methods developed at Rinat-Pfizer to characterize the epitope coverage of antibodies derived from Adimab’s yeast-based library and compare it with the collective epitope coverage of antibodies drawn from the literature against the same target. As a model antigen, we chose hen egg white lysozyme (HEL) because of the wealth of epitope information available on this target in the form of published co-crystal structures representing 20 unique antibodies derived primarily from in vivo sources (the “structural benchmark”), binding to different HEL epitopes. We also illustrate how high-throughput binning methods can be used to reveal the intricate epitope
diversity of antibody libraries, thereby informing early-stage discovery of therapeutic antibodies when it is advantageous to identify antibodies representing a diversity of functional epitopes for lead development.

**Results**

The results are presented in the following order. First, we outline the steps involved in library selections, kinetic characterization of HEL binders, epitope binning and selection of the literature controls. We then present the experimental cross-blocking data for 7 control antibodies considered as an isolated subset and retrospectively validate a structure-based metric that correctly predicts these results. Next, we deploy this metric to predict a cross-blocking matrix of the set of 20 literature controls to gain insight into the epitope coverage of the entire structural benchmark. Finally, we present the experimental binning data for the entire set of library clones and 7 controls, and compare the epitope coverage of the library to that of the structural benchmark.

**Binning study of 350 anti-HEL antibodies discovered via selections using a human in vitro library**

Selections were performed against monomeric HEL antigen using the Adimab synthetic IgG antibody library (see Methods). A panel of 350 antibodies was included in the binning study based on their confirmed binding in an Octet screening assay toward soluble HEL and sequence uniqueness. The antibody variable region sequences mapped to light chain germline families Vk1, Vk2, Vk3 and Vk4 and heavy chain germline families VH1, VH3, VH4, and VH5. Thirteen of the 16 possible VH/Vk germline family pairings are represented in this antibody panel. Five pairs, VH3 Vk1, VH4Vk3, VH4Vk1, VH3 Vk3 and VH1 Vk1, accounted for 77% of the panel (Table S1). The complementarity-determining region (CDR) H3 length (IMGT definition) ranged from 10 to 23 amino acids (AA) with a median of 15 AA (the library design comprised lengths 4 to 31 AA); L3 lengths ranged from 8–10 AA with 9-AA L3 sequences comprising the majority (85%) in line with the input library design. The polyreactivity of the antibody clones was assessed using a previously published protocol and categorized into low (72%), medium (8%) and high (20%) levels based on poly-specificity reagent mean fluorescence intensity (PSR MFI) cutoffs of 200, 500 and > 500, respectively. While low polyreactivity is generally desired, clones that exhibited medium to high PSR binding levels were retained because this property can be optimized by engineering, if clones have interesting enough epitopes to warrant their further development.

The panel of library anti-HEL antibodies exhibited diverse binding kinetics, informing the design of the epitope binning experiments

To assess the affinity range of the antibodies used in this study, we used a ProteOn XPR36 surface plasmon resonance (SPR) biosensor and characterized the binding kinetics of HEL toward a panel of 7 control antibodies, CTL-1, 2, 3, 4/4, and 5/5, chosen from the literature (discussed subsequently) and the panel of library clones. The apparent equilibrium dissociation constants (K_D values) for the controls ranged from ~20 pM to 1 nM, mostly showing very slow dissociation rate constants (k_d values of ~2 × 10^{-4} s^{-1}), approaching the resolution of the employed capture-based approach (Table 1). In contrast, the library was kinetically diverse, showing apparent K_D values ranging from 82 pM to 69 nM, with the majority of clones in the single digit nM range, and having relatively fast k_d values (4 × 10^{-3} to 0.13 s^{-1}). Representative examples of the global kinetic analysis are shown in Fig. 1A, and an isoaffinity plot of the kinetic rate constants obtained for the entire study is provided in Fig. 1B.

Characterizing the binding kinetics of the antibodies guided the design of our array-based SPR imaging epitope binning assays because clones with fast k_d values are generally ill-suited as capture antibodies (or “ligands”) when coupled to the chip. To mitigate this limitation, we employed a “co-injection” approach to sample delivery in our binning experiments whereby an injection of HEL was followed immediately by an injection of solution antibody (analyte), thereby minimizing the time allowed for HEL to dissociate from the ligand array during the sandwiching step. The single flow cell configuration employed by array-based SPR imaging relied on the use of a universal regeneration condition that did not work perfectly for all ligands. To compensate for ligand attrition in

| Antibody | k_a (1/Ms) | k_d (1/s) | K_D (nM) |
|-----------|------------|-----------|----------|
| D1.3      | 2.9 (1.3) × 10^6 | 3.2 (0.4) × 10^{-3} | 1.21 (0.36) n = 5 |
| D11.15    | 4.4 (1.6) × 10^6 | 5.5 (1.3) × 10^{-3} | 1.35 (0.42) n = 5 |
| cAb-Lys3  | 2.2 (0.9) × 10^6 | 1.1 (0.5) × 10^{-3} | 0.52 (0.10) n = 4 |
| HyHEL-8   | 2.6 (0.7) × 10^6 | 2.1 (0.7) × 10^{-4} | 0.083 (0.029) n = 5 |
| HyHEL-10  | 2.6 (0.1) × 10^6 | 1.7 (0.5) × 10^{-4} | 0.065 (0.019) n = 4 |
| F10.65    | 1.1 (0.3) × 10^7 | 2.5 (1.5) × 10^{-4} | 0.024 (0.014) n = 4 |
| HyHEL-5   | 9.5 (1.6) × 10^6 | 2.3 (0.6) × 10^{-4} | 0.025 (0.010) n = 5 |

| In vitro library (N = 350 antibodies) | |
|---------------------------------------|--------|
| 0% quartile                           | 7.7 × 10^0 |
| 25% quartile                          | 6.1 × 10^0 |
| 50% quartile                          | 1.2 × 10^1 |
| 75% quartile                          | 2.4 × 10^0 |
| 100% quartile                         | 3.0 × 10^0 |

Table 1. SPR affinity determinations of HEL/anti-HEL antibody binding interactions. A one-shot kinetic method was used on a ProteOn XPR36 biosensor to determine the affinity of solution HEL binding to immobilized antibodies; all antibodies were reformatted into human IgG1 and captured via goat anti-human-IgG (Fc-specific) reagent coated on NLC or GLC chips. The values reported for the literature controls are the mean (and standard deviation) of N replicates. See Fig. 1.
a given epitope binning campaign, we therefore address the entire 2-dimensional "analyte x ligand" interaction matrix. To bypass the need for any regeneration, we also performed lower-throughput epitope binning assays on a subset of clones using the Octet, a commonly used biolayer interferometry (BLI)-based biosensor technology that employs single-use disposable sensors. The complementary strengths and limitations of SPR imaging and Octet technologies in the context of label-free epitope binning assays have been discussed elsewhere.7 Further details of the kinetic characterization and epitope binning experiments are provided in the Methods section.

The collective epitope of the 7 controls covers 60% of HEL’s accessible surface area

To benchmark the epitope diversity of the library clones, we sought a panel of control antibodies from the literature with crystallographically-defined epitopes that collectively covered a large area of the HEL surface. Based on a comprehensive search of the Protein Data Bank (PDB),13 we identified 79 antibody:lysozyme complexes, 20 of which represent unique antibody sequences. Details of these 20 complexes, constituting the “structural benchmark,” are presented in Table 2. The formatted coordinate files for the complexes are provided as supplementary material. The majority of these antibodies originate from rodent and camelid immunizations (n = 16) and a small number (n = 4) from in vitro library designs. Based on solvent-accessible surface area calculations (data not shown) using the crystal structures, the binding epitopes of the 20 antibodies collectively cover 87% of the lysozyme solvent-accessible surface area of ~6100 Å². To capture the breadth of epitope coverage found in the literature, we selected a subset of 5 antibodies as primary experimental controls, namely D1.3, D11.15, cAbLys3, HyHEL-8, and F10.6.6, referred to here as CTL-1 through CTL-5, respectively. To probe fine specificities, we added to the control panel HyHEL-10 (CTL-4*) and HyHEL-5 (CTL-5*), which bind near-identical epitopes as their respective counterparts, CTL-4 and CTL-5. Antibodies CTL-1, 4 and 5 have been used in previous HEL epitope binning studies.14 The camelid antibody (CTL-3) displays an interesting binding mode to the HEL active site using a long CDR3 loop.15,16 The 7 controls, whose collective epitope covers 60% of the HEL solvent-accessible surface area, were prepared recombinantly for use in the binning experiments. An additional 12 antibodies in the set (not including Ab_4n1e) increase the coverage to 78%, implying rather low marginal increase per added control. The remaining 9% of the collective epitope is recognized by the VL:VL homodimer (antibody Ab_4n1e) corresponding to PDB entry 4n1e. While this antibody would have been a valuable addition to our control panel, it was deposited into the database after we had finalized the set of controls.

The control antibody panel was critical here to compare and contrast the epitope coverage of the library clones vis-à-vis the structural benchmark. Each antibody was assigned a "blocking profile" (see Methods), which is a single annotation that merges the pairwise cross-blocking data against the control antibodies. The panel of 5 primary controls leads to 32 (25) such theoretical profiles, of which 25 were structurally compatible with the simultaneous cross-blocking of at least 2 controls. Since it intersects information related to cross-blocking of control antibodies binding to distinct HEL epitopes, the profile definition is generally

Figure 1. One-shot kinetic analysis of HEL/anti-HEL antibody binding interactions using the ProteOn XPR36 biosensor. (A) Global fits for select clones chosen from the literature (top panel) and the in vitro library (bottom panel), where the overlay plots show the measured data (noisy lines) and simulated fit (smooth, color-matched lines). (B) Isoaffinity plot graphing the kinetic rate constants for the entire set of clones used in this study. The inset shows the capture-based assay format employed.
Table 2. Literature anti-HEL antibodies used in theoretical structure-based cross-blocking predictions. Unless stated otherwise, the PDB code refers to the co-crystal complex with HEL; complex with 1) pheasant egg lysozyme, 2) turkey egg-white lysozyme, 3) guinea fowl lysozyme and 4) human lysozyme. Seven antibodies were selected as controls and prepared recombinantly for use in epitope binning experiments. See Methods and Table 3 for explanation of the blocking profile assignments. The type of the antibody format, i.e., whether standard VL:VH heterodimer (VL:VH), camelid heavy chain only antibody (VHH), heavy chain only antibody (VH) or light chain dimer (VL:VL) is indicated in addition to the source technology. The coordinates corresponding to the light (LC), heavy (HC) and lysozyme (Ag) chain IDs were extracted from each PDB entry and used to compute metrics described in the text.

| PDB code | Resolution (Å) | Antibody | Control name | Blocking profile | Construct | Source | LC | HC | Ag |
|----------|----------------|----------|--------------|------------------|-----------|--------|----|----|----|
| 1vfb     | 1.80           | D1.3     | CTL-1        | 12               | VL:VH    | Rodent immunization | A  | B  | C  |
| 1jhl     | 2.40           | D11.15   | CTL-2        | 123              | VL:VH    | Rodent immunization | L  | H  | A  |
| 1jto     | 2.50           | cAbLys3  | CTL-3        | 23               | VH       | Camelia immunization | A  | L  |    |
| 1ndg     | 1.90           | HyHEL-8  | CTL-4        | 4                | VL:VH    | Rodent immunization | A  | B  | C  |
| 1cd8     | 2.30           | HyHEL-10 | CTL-4*       | 4                | VL:VH    | Rodent immunization | A  | B  | C  |
| 1pzc     | 2.00           | F10.6    | CTL-5        | 5                | VL:VH    | Rodent immunization | A  | B  | C  |
| 1uyv     | 1.70           | HyHEL-5  | CTL-5*       | 5                | VL:VH    | Rodent immunization | L  | H  | Y  |
| 1zhb     | 2.00           | I19      | —            | 1234             | VL:VH    | Rodent immunization | A  | A  | X  |
| 1fb1     | 3.00           | F9.13.7  | —            | 4                | VL:VH    | Rodent immunization | L  | X  |    |
| 1qpt8    | 1.86           | HL6      | —            | 4                | VH       | Camelia immunization | A  | B  |    |
| 1ri8     | 1.85           | 1D2L19   | —            | 35               | VH       | Camelia immunization | A  | B  |    |
| 1rjc     | 1.40           | cAb-Lys2 | —            | 235*             | VH       | Camelia immunization | A  | B  |    |
| 1zv5     | 2.00           | D2-L29   | —            | 35               | VH       | Camelia immunization | A  | L  |    |
| 1zvh     | 1.50           | D2-L24   | —            | 5                | VH       | Camelia immunization | A  | L  |    |
| 1zvy     | 1.63           | D3-L11   | —            | 234              | VH       | Camelia immunization | A  | B  |    |
| 4ic5     | 1.95           | cAbHel-5 | —            | 12               | VH       | Camelia immunization | D  | A  |    |
| 4n1c     | 1.70           | Ab_4n1c  | —            | 123              | VL:VL    | In vitro library    | A/B| C  |    |
| 4n1e     | 2.23           | Ab_4n1e  | —            | 0                | VL:VL    | In vitro library    | A/B| I  |    |
| 4tsa     | 2.27           | Ab_4tsa  | —            | 123              | VL:VH    | In vitro library    | L  | H  | A  |
| 4u3x     | 2.26           | H04      | —            | 235              | VH       | In vitro library    | A  | B  |    |

Epitope binning studies confirmed that the 7 literature control antibodies fall into 5 bins when considered as an isolated subset

To establish confidence in the 3D structure-based interpretation of the binning data from the subsequent high-throughput experiments, we first binned the 7 control antibodies against one another using the Octet and validated the results against expectations based on the co-crystal structures. Fig. 2A shows an image of 5 control antibodies, CTL-1 through CTL-5, as ribbon diagrams superposed as an overlay plot on the HEL surface, shown as a white space-filled model. Collectively, it appears that the controls densely cover one face of the HEL surface and occupy 3 non-overlapping regions; (1) CTL-1, 2, and 3; (2) CTL-4; and (3) CTL-5. CTL-1 (blue) and CTL-2 (green) would be predicted to block one another because their epitopes appear to overlap significantly, but it is unclear whether the minimally overlapping epitope contacts between CTL-2 and CTL-3 (red) would be sufficient to manifest as a blocking pair (see Fig. 2A - inset). We employed a classical sandwich assay format, in which a solution antibody (analyte) is tested for binding to HEL that is first captured by an immobilized antibody (ligand). An example of the data obtained on amine-coupled coupled antibody D11.15 (CTL-2) is shown in Fig. 2B. Antibody analytes that showed significant binding responses were classified as “not blocked” (also known as “sandwich pairing” with the coupled antibody), whereas those that showed negligible binding, similar to that of a buffer analyte were classified as “blocked.” By addressing all pairwise permutations of the 7 control mAbs, we generated a comprehensive interaction matrix, as shown in the heat map of Fig. 2C, where the red and green cells indicate blocking and non-blocking (or sandwiching) pairs, respectively. Five epitope bins emerge from this analysis, populated by (1) CTL-1, (2) CTL-2, (3) CTL-3, (4) CTL-4 and CTL-4*, and (5) CTL-5 and CTL-5*, as graphed in the network blocking plot of Fig. 2D, where chords between antibodies indicate the blocking relationships, and bins are represented by the envelopes. Consistent with the structural data, the 7 control antibodies fall into 3 non-overlapping bin clusters, since bin 2 overlaps with bin 1 and bin 3, while bin 4 and bin 5 do not overlap with any other bin.

Theoretical predictions of cross-blocking for the structural benchmark returned a complex network of epitope bins

We attempted to predict the cross-blocking pattern of the 7 controls retrospectively using 2 in silico metrics (see Methods). The “Fv Ca metric” models the cross-blocking of a pair of antibodies on the basis of steric hindrance between the Fv regions in their respective binding orientations, while the “common epitope AA metric” does so on the basis of overlapping antibody binding footprints as suggested by antigen residues common to the pair of epitopes. The “Fv Ca metric” heat map for the 7 controls is shown in Fig. 3A, where the red and green cells represent the structure-based predictions for blocking and sandwiching pairs of antibodies, respectively. These predictions are consistent with the experimental data (Fig. 2C), i.e., antibody pairs with one or more Ca clashes defined using a 2.5 Å cutoff radius cross-block each other experimentally, while other pairs lacking such clashes are able to sandwich one another. In contrast, the “common epitope AA metric” returned false positives (Table S2A). For example, CTL-1 (D1.3) is predicted to cross-block CTL-3 (cAbLys3) and CTL-4/4* (HyHEL-8/HyHEL-10) on account of 2 and 5
common epitope amino acids, respectively, in contrast to the experimental "sandwiching" result for these pairs.

To assess the epitope coverage of the literature HEL antibodies as represented by the structural benchmark, we generated theoretical predictions on the 20 co-crystal structures (Fig. 3B) using the "Fv Ca metric" validated above, experimentally, for the 7 controls. With the caveat that the predictions featuring the 13 antibodies not included in the control panel must be verified experimentally, the information from all 400 pairwise permutations of the 20 antibodies creates a complex web of 18 distinct bins, 16 of which are populated by a single antibody. These 18 epitope bins and their interconnectivities are alternatively visualized in terms of a network blocking plot (Fig. 3C), colored by "blocking profile." Two bins are each populated by a pair of antibodies; HyHEL-8 and HyHEL-10 belong to the same bin, and 1D2L19 and D2-L29 belong to the same bin. In other words, no antibody within the panel of 20 is predicted to be able to discriminate the epitopes of 1D2L19 and D2-L29 or HyHEL-8 and HyHEL-10. In other cases, the larger interaction matrix introduces antibodies whose specificities fragment bins into sub-bins, thereby highlighting the increased epitope resolution that results from higher throughput cross-blocking analyses. For instance, 2 antibodies with near-identical epitopes (e.g., F10.6.6 and HyHEL-5) are discriminated by a single antibody (cAb-Lys2), resulting in F10.6.6 and HyHEL-5 occupying their own bins. The "structural benchmark" returns a total of 11 blocking profiles and includes examples of antibodies that simultaneously block just one control (HyHEL-8, profile 4 i.e., self-blocking), 2 controls (D2-L29; profile 35), 3 controls (D3-L11; profile 234), 4 controls (1f9; profile 1234) or none of the 5 controls (Ab_4n1e; profile 0).

**Cross-blocking data for the library clones reveals 2 main epitope communities**

To characterize the epitope coverage of the library-derived panel, we performed a series of epitope binning experiments by SPR imaging using a 384-ligand array format that provided throughput higher than previously published methods.7 By merging 350 library clones and 7 literature controls on a single chip, we aimed to probe finer epitope resolution than may be available by considering the library clones in isolation. Each experiment tested 96 antibody analytes over a 384-ligand array, as limited by the SPR imager’s autosampler capacity, thereby necessitating 4 consecutive experiments on the same chip to address all 350 antibodies as analytes. The merged cross-blocking results from this series of experiments are summarized in the heat map shown in Fig. 4A. These data are alternately visualized as a high-level community plot (Fig. 4B), revealing 2 main communities, one populated by 176 library clones and all 7 controls (community 1, shown in purple), and another populated by 167 library clones (community 2, shown in cyan blue). These results imply that community 1 accesses epitopes on the front face of HEL (see Fig. 2A), whereas community 2 accesses epitopes on the back face of HEL (see Fig. 8B). The heavy meshing between the communities indicates that they are...
intricately interconnected by clones that cross-block both, requiring a more granular analysis to tease out finer specificities. The library therefore appears to contain epitope diversity that overlapped with and extended beyond that of the controls.

The repertoire of library epitopes is represented by a subset of clones, providing “internal” controls to facilitate future comparisons

To obtain a higher-resolution analysis of the data, we focus on a subset of 51 clones that broadly represent the library’s diversity and for which clear pairwise cross-blocking data are available in both analyte and ligand orientations. Within this smaller subset, additional details emerge (Fig. 5A). The two main communities persist, but clones such as 3F2 and 3D2 (red bin) and 4B9, 3C4, 3E10, 4C2, and 3H9 (green bin) appear to segregate from their respective bin counterparts due to their cross-blocking of non-bin clones. We further distilled the panel to a subset of 7 clones whose overlapping epitopes spanned the breadth of the library’s epitope coverage, allowing their use as internal standards against which other clones could be compared in the absence of any literature controls (Fig. 5B).

The library clones returned a larger number of “blocking profiles” relative to the structural benchmark, suggesting broader epitope coverage

Table 3 summarizes the blocking profile distribution of a set of 243 library clones for which clear assignments were possible when the controls were injected as analyte over a 384-ligand array comprising the library and controls. Seventeen “blocking profiles” (defined in Methods) emerged in addition to an unexpected one that discriminated between the near-identical epitopes of CTL-5 and CTL-5/C3 (blocking profile 5'). The majority of the library fell into profile 1 (91 clones) or profile 1234 (60 clones). Five other profiles (0, 5, 123, 345 and 2345) are represented by at least 10 clones, while the remaining profiles are represented by 1–3 clones each (so-called “rare” profiles). To validate the profile assignments obtained by SPR imaging, we performed an independent set of smaller, more focused experiments on the Octet. Fig. 6 shows a side-by-side technology comparison for some of the library clones that populated rare profiles in our analysis. The binning outcomes were in excellent agreement across the complementary technology platforms, confirming that the 384-ligand array SPR imaging format was reliable as a high-throughput binning platform.

The Venn diagrams shown in Fig. 7 compare the epitope coverage observed in (A) the structural benchmark and (B) the library, in terms of their blocking profiles. Our structure-based theoretical analysis of 20 antibodies from the literature returned 11 predicted profiles, all but one of which (profile 235°) are represented in the collection of 17 profiles among the library clones. Judged by the metric of the number of cross-blocking profiles, the library appears to return expanded epitope coverage relative to the structural
benchmark, although we recognize that the library panel contains over 10 times more clones than the benchmark. Notably, antibodies that block CTL-1 exclusively (profile 1) are the most abundant (n = 91) in the library, but do not occur in the literature set. The next most abundant profile (1234, n = 60) in the library is, however, also observed in the literature set (antibody 1F9). Five other profiles, namely 13, 14, 34, 345 and 2345 with n = 2, 2, 3, 14 and 13 clones assigned, respectively, in the library are not represented in the literature. Based on protein-protein docking calculations, details of which are provided in the supplemental text, it appears that the 6 novel profiles are geometrically compatible with the binding footprint of an antibody on the HEL surface (Fig. S1).

Profile 0 and profile 5⁺ demonstrate the broad epitope coverage obtained from the library. Fig. 8A shows examples of the data obtained for library clones exhibiting profile 0 (2F9, top panel) and profile 5⁺ (1E4, bottom panel). In both cases, the assignments deduced from SPR imaging data were reinforced by Octet data. Clones belonging to profile 0 do not cross-block any of the controls. These so-called “universal sandwichers” represent 10% of the antibody panel. Antibody Ab_4n1e in the structural benchmark, which became available after we finalized the list of experimental controls, is also predicted to be a “universal sandwich.” In contrast to the front view of HEL, which shows dense coverage by the controls (Fig. 2A), the back view shows sparse coverage, revealing a bald spot that could offer novel epitopes (Fig. 8B) relative to these controls. Indeed, the universal sandwichers mutually blocked one another, upon detailed inspection of the binning analysis shown in Fig. 4 (cyan blue community) and Fig. 5 (red bin), suggesting that they targeted a common region on the HEL surface. While the definition of a “universal sandwich” depends on the controls being used, in Fig S2, we visualize the binding of Ab_4n1e and CTLs 1, 3, 4 and 5, and note that the “bald spot” represents a different epitope than the one in Fig. 8B.

Clones with profile 5⁺ display fine epitope specificity by cross-blocking F10.6.6 (CTL-5), but not HyHEL-5 (CTL-5⁺), despite the near-identical epitopes of these controls as compared in Fig. 8C. The set of 243 clones contained a single member (1E4) with this profile and further inspection of the full panel revealed a few more clones that belonged to profile 5⁺. Fig. 3 illustrates that the literature also contains a clone (cAbLys2) that is predicted to block HyHEL-5 (CTL-5⁺) and not F10.6.6 (CTL-5), but it is predicted to block other controls, D11.15 (CTL-2) and cAbLy3 (CTL-3), so it is assigned to profile 235⁺ (see Table 2).

Inspecting the germline family distributions within the most abundant individual blocking profiles and comparing them against the baseline distribution in the set of 243 clones with assigned profiles reveals a few trends (Table S4). Profile 1234 is dominated by VH3 sequences (54/60 clones; 90%), while VH1 and VH4 sequences account for the vast majority (76/91; 83%) of profile 1 antibodies. VH4 family antibodies are 1.4-fold over-represented in profile 0 (56% versus 40%), while VK4 sequences are 2-fold over-represented (23% vs. 11%) in profile 1.

**Discussion**

In this work, we characterized the epitope coverage obtained using output from a naïve human in vitro library and HEL as a model antigen. We demonstrated that emerging SPR imaging methods employing a 384-array format.
can be used to perform epitope binning assays with superior throughput compared to those published previously, enabling a deeper interrogation of a library’s epitope diversity. As judged by their ability to cross-block a panel of control antibodies selected from the literature, the epitopes of the library clones appear to cover > 75% of the HEL accessible surface, with an abundance of antibodies exhibiting particular binding footprints, such as those that exclusively block CTL-1 (profile 1). In addition, the library contains clones with a diversity of binding footprints across the entire HEL solvent-accessible surface, as judged by the 17 observed blocking profiles compared to 11 for the structural benchmark. It is noteworthy that, our structural-based analysis predicted a blocking profile of 235° for the literature clone cAb-Lys2, but, upon empirically testing its ability to cross-block the controls, we found that it clearly blocked CTL-3, CTL-5 and CTL-5°, whereas its blockade of CTL-2 was unclear (data not shown). Therefore, its empirically determined profile was 35 (or perhaps, 235), and contrary to our prediction, it did not discriminate between CTL-5 and CTL-5°. We did not empirically test of all 20 literature controls, and thus our structural predictions may have over-granularized the bin count in the literature, as opposed to the empirically confirmed bin count for our library. This example highlights that caution should be applied when predicting cross-blocking relationships based on structural data.

The germline sequence diversity of the library contributes to its broad epitope coverage since no individual light- or heavy-chain germline is observed across all 17 profiles. Antibodies with VH3 or Vk sequences are found in 14 and 15 profiles, respectively, but the VH3:Vk pairing itself returns only 11/17 observed profiles, and 3/6 profiles novel relative to the structural benchmark. Though no counter-selection against polyreactivity was employed during the selections, each of the 17 observed cross-blocking profiles was populated by one or more clones with low assayed polyreactivity. Thus, the deep pool of selected binders and the high-throughput epitope binning methods limit attrition in the epitope coverage due to polyreactive clones.

The library studied in this work has produced broad epitope coverage to various therapeutic targets, which has translated into tangible benefits for some of the corresponding programs. MM-151, a clinical-stage oligoclonal cocktail of 3 antibodies targeting epidermal growth factor receptor (EGFR), is an example. The enhanced therapeutic efficacy of MM-151 has been attributed to binding of novel epitopes located on EGFR domain I and domain III that are distinct...
from those of cetuximab and panitumumab. Another example is HT-19, an anti-human epidermal growth factor receptor (HER)2 antibody being developed as an antibody-drug conjugate, which binds an epitope distinct from that of pertuzumab and trastuzumab, and indeed from other known domain I17 and domain III18 HER2 epitopes. A third example is the broadly neutralizing, cross-reactive antibody against a-hemolysin and 4 additional leukocidins, with superior effectiveness in combating S. aureus infections compared to antibodies targeting a-hemolysin alone.8,19 The cross-reactivity results from the specific binding to an epitope that includes a small conserved patch of amino acids, present in the background of the overall low (26%) sequence homology among the 5 targets.8,19

The present study extends the scope of an earlier one carried out by Newman et al in 1992.14 In that study, a panel of 49 anti-HEL antibodies derived from a series of BALB/c mice immunizations were localized to different epitopes on HEL by testing their ability to cross-block a panel of control antibodies, including 3 with crystallographically-deined epitopes (D1.3, HyHEL-8, and HyHEL-5) which were also used in our control panel (CTL-1, 4 and 5/C3). Overall, the Newman et al study concluded that the epitopes of the murine antibodies collectively covered at least 80% of the lysozyme surface and speculated the availability of at least 6 non-overlapping epitopes for antibody binding. The multiple antibody:lysozyme crystal structures determined since 1992 reinforce these conclusions, as
shown by our theoretical analysis with 20 co-crystal structures, suggesting a collective coverage of 87% of HEL’s solvent-accessible surface. Furthermore, antibodies D1.3 (CTL-1), cAbLys3 (CTL-3), HyHEL-8 (CTL-4), F10.6.6 (CTL-5) and Ab_4n1e represent 5 structurally verified non-overlapping epitopes, leaving room for a sixth, as proposed by Newman et al (Fig. S2).

Our work demonstrates how high throughput epitope binning analyses can reveal deep insight into the epitope landscape of a given antibody campaign, and, by merging panels of
antibodies from different sources, how these assays can serve to benchmark libraries and selection strategies against one another. Ultimately, broad epitope coverage provides options for convergence upon the most promising lead molecules based on target biology, species cross-reactivity, developability characteristics, and intellectual property opportunities, and is thus critical to the success of therapeutic antibody programs.

Methods

Profile definitions based on cross-blocking of CTL-1 through CTL-5

Each antibody was assigned a “blocking profile” based upon its combinatorial blockade of the 5 primary controls, CTL-1 through CTL-5. Since CTL-4" and CTL-5" share the same bin as their respective counterparts CTL-4 and CTL-5 when the set of 7 controls are binned against one another (Fig. 2D), we reasoned that only 5 antibodies are required to represent the collective diversity of the 7 controls. These assignments were made using the experimental data for the library clones and the set of 7 controls, and on the basis of theoretical predictions for other antibodies in the structural benchmark (summarized in Table 3). The primary controls, CTLs 1–5, are schematically represented by the 5 lobes, following the coloring scheme used in Fig. 1. The number of antibodies assigned to a particular profile is provided in the overlapping regions between the relevant lobes, except for improbable profiles featuring simultaneous blockade of CTL-1 and CTL-5 (see Methods), which are left blank. Six profiles (1, 13, 14, 34, 345, and 2345) that are observed in the library-derived panel, but not predicted in the "structural benchmark," have been highlighted. Additional profiles that are not shown include profile 235", which is predicted in the structural benchmark, and profile 5", which was observed in the library-derived panel. The tool provided at http://bioinformatics.psb.ugent.be/webtools/Venn/ was used to generate the diagrams.

![Venn diagrams](image)

Figure 7. Venn diagrams, showing the distribution of "blocking profiles" (defined in Methods) for (A) the 20 antibodies used as our "structural benchmark" (based on structural predictions) and (B) the set of 243 library antibodies (based on empiric cross blocking data; see Table 3). The primary controls, CTLs 1–5, are schematically represented by the 5 lobes, following the coloring scheme used in Fig. 1. The number of antibodies assigned to a particular profile is provided in the overlapping regions between the relevant lobes, except for improbable profiles featuring simultaneous blockade of CTL-1 and CTL-5 (see Methods), which are left blank. Six profiles (1, 13, 14, 34, 345, and 2345) that are observed in the library-derived panel, but not predicted in the "structural benchmark," have been highlighted. Additional profiles that are not shown include profile 235", which is predicted in the structural benchmark, and profile 5", which was observed in the library-derived panel. The tool provided at http://bioinformatics.psb.ugent.be/webtools/Venn/ was used to generate the diagrams.

According to this scheme, the control antibodies CTL-1, 2, 3, 4/4", and 5/5" were defined by the blocking profiles 12, 123, 23, 4, and 5 (see heat map in Fig. 2C for their empiric cross-blocking results). Eight of the theoretical blocking profiles (15, 125, 135, 145, 1245, 1345, and 1345) were deemed improbable because they involved simultaneous cross-blockade of CTL-1 and CTL-5, whose epitopes are too far apart to be bridged by a third antibody, according to structural calculations (see Table S3). To illustrate the utility of the profile definition, consider the alternate possibilities for a clone to block CTL-1 exclusively (profile 1), or in conjunction with blocking of other controls (profiles 12, 13, 14, 123, 124, 134, 1234). Each of these profiles represents a different mode of blocking CTL-1, with partial overlap between the modes. Thus, in general, a greater number of observed profiles for a set of library clones implies better epitope coverage.

3D structure metrics for predicting antibody blocking

From a structural perspective, 2 antibodies can block one another directly, by competing for common epitope amino acid contacts, or indirectly, via steric clashes upon binding to adjacent epitopes. When co-crystal structures are available, a theoretical prediction for cross-blocking can be made by quantifying the extent of such epitope overlap or steric clashes, as described here. For each antibody:antigen complex, we computed the epitope amino acids in each co-crystal structure using a 5 Å heavy atom contact definition. Further, for each pair of co-crystal structures, we superposed the HEL coordinates and computed, 1) the number of the Fv Cα atoms in one antibody
within a distance cutoff of 2.5 Å of the Fv Cα atoms in the second (Fig. 2); 2) the number of common epitope amino acids between the complexes (Table S2); and 3) the distances between the heavy-atom centroids of the respective Fv regions (Table S3). These three metrics were named the “Fv Cα metric,” “common epitope AA metric” and “Fv centroid metric,” respectively. Antibodies with one or more so-defined Fv Cα - Cα contacts or one or more common epitope AA are predicted to “block” each other, and otherwise “sandwich” each other. The matrix of the “Fv Cα metric” values is asymmetrical since the value for a pair of antibodies \( ij \) depends on the number of Fv Cα atoms in antibody \( i \), but this does not affect the “block” or “sandwich” predictions using this metric.

As an example, consider antibodies D1.3 (CTL-1) and D11.15 (CTL-2), which recognize epitopes adjacent on the HEL surface. Each antibody returns 223 Cα atoms in the Fv region, of which \( \sim 40 \) Cα atoms share a contact (2.5 Å), indicating their close approach in binding orientations results in steric clashes (Fig. 2A); the D1.3 and D11.15 epitopes comprise 21 and 15 AA, respectively, of which 9 are common to both antibodies (Table S2A); and the Fv centroid-centroid distance is 16 Å (Table S3A). In contrast, the D1.3:F10.6.6 (or CTL-1: CTL-5) pair have zero shared Fv Cα - Cα or epitope amino acids and a centroid separation of 64 Å.

**Antibody library selections against HEL**

HEL-specific antibodies were isolated from a full-length human IgG1 antibody library using an in vitro yeast presentation system and associated methods as previously described. The library sequences, especially those of CDR H3, are designed to mimic features of the human pre-immune repertoire. Specifically, a yeast population harboring IgG diversity of \( > 10^9 \) was subjected to 3 rounds of enrichment. **Round 1:** \( 1 \times 10^{10} \) yeast cells from each of 8 individual sub-libraries were selected against 3 mL of 100 nM biotinylated HEL (bHEL) for 10 min at 30°C, washed, and then incubated with 500 µL of streptavidin microbeads (Miltenyi Biotec) for 15 minutes at 4°C. Yeast populations of \( 2.9 \times 10^6 - 1.0 \times 10^7 \) were rescued after capture on an LS magnetic column (Miltenyi Biotec). **Round 2:** For negative selection, \( 1 \times 10^8 \) cells were incubated with 500 µL of streptavidin microbeads for 15 minutes at 4°C. Following incubation, the yeast-microbead suspension was applied to an LS magnetic column and the flow-through was retained. For positive selection, non-captured yeast were incubated with 750 µL of 100 nM biotinylated HEL for 10 min at 30°C, washed and incubated with 125 µL of streptavidin microbeads for 15 minutes at 4°C. Yeast populations of \( 9 \times 10^5 - 1 \times 10^6 \) were rescued after capture on an LS magnetic column. **Round 3:** For negative selection, \( 5 \times 10^9 \) cells were incubated with 500 µL of streptavidin microbeads for 15 minutes at 4°C. For positive selection, \( 1 \times 10^8 \) non-captured yeast were incubated with 500 µL or 5 mL of 100 nM or 10 nM biotinylated HEL for 10 min or 30 min, respectively, at 30°C. Following incubation, cells were pelleted, washed and incubated with 1 mL of secondary labeling mix (Extravidin-R-PE, anti-human LC-FITC, and propidium iodide) on ice protected from light for 20 min and run on FACSARia II (BD Biosciences) to record 100,000 PI negative (viable) yeast events. The data were used to sort 5,000–20,000 yeast cells with PtdIns negative, LC-FITC positive (IgG presenting), and EA-R-PE positive (bHEL-binding) phenotypes. A total of 2940 unique clones were produced and...
characterized for HEL binding by Octet, leading to a final selection of 350 antibodies included in this binning study.

### Binding kinetics and affinities

The binding kinetics of soluble HEL (Sigma product number L6876) toward immobilized anti-HEL antibodies, were determined at 25°C and in a running buffer of phosphate-buffered saline (PBS) pH 7.4 + 0.01% Tween20 using a ProteOn-XPR36 surface plasmon resonance (SPR) biosensor (BioRad Inc.). A capture-based one-shot kinetic method was used, as described by Nahshol et al. Goat F(AB')2 anti-human IgG Fc (Cappel product number 55053, lot number 04459) served as the capture reagent and was coated onto the ligand channels of a GLC chip by standard amine-coupling or was biotinylated (using Pierce EZ-link LC-LC linker) and captured via an NLC (neutravidin) chip; final levels of the capture reagent were ~3,000 RU (GLC) or 2,000 RU (NLC). Anti-HEL antibodies (all formatted as human, aglycosylated, IgG1-Fc) were captured at 5 μg/ml along the ligand channels to low levels (100 - 300 RU) and then HEL was injected along the analyte channels at concentrations of 70, 14, 2.8, 0.56, 0.11, and 0 nM (or, in some experiments, 100, 20, 4, 0.8, 0.16, and 0 nM). Association and dissociation times of 3 min and 15 min were allowed. The capture surfaces were regenerated with 75 mM phosphoric acid detergent times of 3 min and 15 min were allowed. The reaction spot data were "double-referenced" by subtracting the interspot (reference) data and the buffer analyte data, and the resulting sensorgrams were fit globally to a simple Langmuir model using the ProteOn Manager software.

### Epitope binning experiments

An SPR imager was used (IBIS MX96) for the epitope binning experiments as described previously, but with a modified continuous flow microspotter (CFM) and immobilization routine designed to increase the number of reaction surfaces addressed to 384. A printhead device with different geometry flow cells (130 μm × 435 μm) than previously employed (400 μm × 500 μm) was used to amine-couple the entire set of 380 antibodies (comprising 350 antibodies from the library and 7 controls in triplicate) on a single Xantec 200 M prism (Xantec GmbH, Germany). The unattended print routine immobilized the 380 antibodies in 8 consecutive prints by cycling activating reagent for 5 mins, the antibodies for 7 mins, and finally rinsing with running buffer (PBS + 0.01% Tween-20). The chip printed with a 384-antibody array was then docked into the SPR imager (IBIS MX96) for online quenching with 1.0 M ethanolamine.HCl pH 8.5. Epitope binning experiments were performed using a classical sandwich assay format, where a binding cycle involved capturing 100 nM HEL for 240 s, immediately injecting one of the competing antibody analytes at 20 μg/ml for 240 s, allowing a 30 s dissociation phase, and finally injecting a single 20 s regeneration pulse of 75 mM phosphoric acid. A plate of 96 antibody analytes was addressed per unattended run, as limited by the autosampler’s capacity. Buffer analytes were drawn from containers or vials and interspersed throughout the assay after every 12 antibody analytes to provide the blanks used in the data analysis. The entire panel of 380 antibody analytes was addressed in 4 consecutive experiments on the same chip taking 6 d and consuming a total of 60 μg HEL and 3 μg antibody for each analyte.

The resultant 153,664 binding sensorgrams from the 4 experiments were then merged in the binning module of ECTO (Wasatch Microfluidics, US), a software tool for processing and analyzing epitope binning data. ECTO includes a series of advanced data analysis routines that were used to help identify non-ideal or problematic behaviors from analytes and ligands. This toolkit was especially useful when curating the 144,400 analyte/ligand pairs, representing the 380 × 380 (analyte x ligand) interaction matrix, that made up the final merged data set (presented in Fig. 4). For example, ~10% of the antibodies behaved poorly as ligands, either due to their inefficient coupling under the conditions used or their inactivation upon acid regeneration, but were highly active in solution. The software was able to correctly identify unusable ligand data based on its signal level and blocking result frequency while preserving the solution binding information from those same species for further analysis. By applying user-defined threshold settings, analyte/ligand pairs are classified as blocked or not blocked (sandwiching) and results are graphed in various ways. For example, to construct a heat map, ligands are represented in rows, analytes are represented in columns, and cells are colored according to whether a given analyte/ligand pair showed a blocked response (red) or a sandwiching response (green). An analyte/ligand pair that gives an intermediate response falling between these threshold settings is shown by a yellow cell. The heat map is sorted by advanced clustering routines and the self-blocking interactions (representing the use of the same antibody in the role of analyte and ligand) are shown along the diagonal with thick box borders in black. The results from a heat map are alternately graphed in terms of a network blocking plot where antibodies are represented as nodes and connected to other antibodies by chords to represent the blocking relationships. A dotted chord indicates a blocking relationship observed in only one orientation. Antibodies displaying similar blocking relationships are inscribed by an envelope to show that they belong to the same “bin.” A community plot is a low-resolution version of a network blocking plot using less stringent threshold settings for comparison criteria.

Confirmatory binning experiments were performed on smaller panels in lower throughput assays using BLI technology (Octet-Red384 equipped with amine-reactive sensors, Pall-Fortebio). Similar conditions were used as described above, except that antibodies were coupled at 30 μg/ml in a coupling buffer of 100 mM 2-(N-morpholino)ethanesulfonic acid (pH 4.0) and the running buffer for the binning was PBS + 0.05% Tween-20 + 5 g/l bovine serum albumin. Octet data were processed and displayed as overlay plots using in-house scripts.

### Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.
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