Affinity Selection from Synthetic Peptide Libraries Enabled by De Novo MS/MS Sequencing

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Abstract
Recently, de novo MS/MS peptide sequencing has enabled the application of affinity selections to synthetic peptide mixtures that approach the diversity of phage libraries (> 10^8 random peptides). In conjunction with ‘split-mix’ solid phase synthesis to access equimolar peptide mixtures, this approach provides a straightforward means to examine synthetic peptide libraries of considerably higher diversity than has been feasible historically. Here, we offer a critical perspective on this work, report emerging data, and highlight opportunities for further methods refinement. With continued development, ‘affinity selection–mass spectrometry’ may become a complimentary approach to phage display, in vitro selection, and DNA-encoded libraries for the discovery of synthetic ligands that modulate protein function.

Keywords Synthetic peptide libraries · De novo sequencing · Mass spectrometry

Introduction
Synthetic peptide libraries arguably spurred the development of combinatorial chemistry (Lowe 1995), and developed in parallel with phage libraries (Scott 1992)—a powerful tool for discovering binding molecules from libraries of random peptides (Smith and Petrenko 1997; Sidhu et al. 2000; Kehoe and Kay 2005). As discussed in a classic review, central to the phage display technique are affinity selections, which separate binding molecules from non-binders in a single physical step, and yield mixtures of reduced complexity that are enriched for binding molecules (Clackson and Wells 1994). In contrast to screens, which measure the activity of many compounds individually, affinity selections require a method to sequence individual peptides from complex mixtures (Clackson and Wells 1994). For phage display, this task was facilitated by (1) use of sequential rounds of selection and propagation of selected phage, which serve to reduce mixture complexities by amplifying high-affinity clones; and (2) the ease with which individual clones can be isolated for Sanger sequencing, by plating at dilution. More recently, next-generation sequencing can be used to sequence many clones in parallel, after a single affinity selection step (Dias-Neto 2009; Matochko et al. 2012).

Historically, the challenge of sequencing individual peptides from mixtures precluded the use of affinity selections with libraries of synthetic peptides. Therefore, although many early peptide libraries were accessed by chemical synthesis, their use was more appropriate for screening—in arrayed formats (Fodor, et al. 1991; Frank 1993), while bound to solid support beads (the ‘one-bead one-compound’ approach) (Lam et al. 1991), or as a series of mixtures, each with a known fixed position (the ‘positional scanning’ approach) (Houghten et al. 1991). A number of early uses of synthetic peptide libraries in affinity selections are suggestive of interest in the concept (Flynn et al. 1991; Zuckermann et al. 1992; Zhou et al. 1993). However, analysis of selected peptides was limited to either pooled sequencing (yielding positional frequencies only), or to sequence inference by unique molecular mass (applicable to small libraries only).

A solution to the challenge of sequencing individual peptides from complex mixtures—such as those generated by affinity selection from synthetic libraries—is provided by liquid chromatography/tandem mass spectrometry. In the
mid 1990s, this approach was used by at least two groups to identify individual peptides post-affinity selection, by manual interpretation of tandem MS spectra (Chu et al. 1996; Kelly et al. 1996). However, despite early interest in the general use of mass spectrometry to characterize synthetic libraries (Metzger et al. 1993; Till et al. 1994), peptide sequencing by tandem MS was not routine. Moreover, when mass spectrometry has been applied to peptide libraries, it has generally been through ‘ladder sequencing’ (not applicable to peptide mixtures) (Chait et al. 1994; Youngquist et al. 1995), or by tandem MS-based sequencing of individual peptides (Paulick et al. 2006).

Here, we discuss how automated de novo peptide sequencing using nano-liquid chromatography/tandem MS has become a practical approach for sequencing synthetic peptide mixtures (Vinogradov et al. 2017), and enables the application of affinity selections to synthetic libraries of high diversity (up to $10^8$–$10^9$ random peptides) (Touti et al. 2019; Quartararo et al. 2020) (Fig. 1). These numbers are comparable to early phage libraries, and motivate a re-examination of synthetic peptide libraries as sources of novel binding molecules. In our view, success hinges on two key issues: (1) whether de novo sequencing provides sufficient coverage of selected mixtures with high accuracy, such that sequencing data is meaningful; and (2) whether affinity selections can recover rare binding molecules efficiently with high enrichment, such that determinants of binding affinity are discernable amongst sequenced peptides. We discuss both recently published results and new data that speak to these questions, provide the interdisciplinary background necessary to place the work in context, and highlight open questions. Emphasis is placed on the p53/MDM2 interaction as a system for benchmarking the performance of synthetic libraries.

**‘Split-Mix’ Synthesis Enables Access to Synthetic Peptide Libraries of Defined Diversity**

Straightforward access to equimolar peptide mixtures for use in affinity selection–mass spectrometry is enabled by ‘split-mix’ synthesis (Lam et al. 1991; Houghten et al. 1991; Furka et al. 1991) on mono-sized resin beads (Rapp et al. 1992; Bayer 1991) (Fig. 2). Following chain assembly, the resulting peptide mixture is cleaved from the solid support, isolated, and ready for input to affinity selections (Fig. 1). A strength of this approach is its flexibility: both the number of unique peptides (the library diversity) and the quantity of individual peptides can be controlled by varying the synthetic program, the quantity of resin beads employed, and the resin bead particle size and loading capacity. How these variables interact to determine the outcome of ‘split-mix’ synthesis is essential background, as they define the range of libraries accessible to the approach.

In general, the number of peptides accessed by ‘split-mix’ synthesis is determined by either of two factors: (1) the theoretical diversity of the library, when small; or (2) the number of beads employed, when theoretical diversity...
is high (‘theoretical diversity’ is the number of unique compounds that would be present, if all possible combinations of amino acids were sampled, and is determined by the synthetic program—the number of varied positions, and the number of monomers used at each). For ‘low diversity’ libraries, each compound is represented on many individual beads, and the library is said to be ‘over-sampled’; using more beads increases the quantities of individual peptides generated, but not their total number. For ‘high diversity’ libraries, in contrast, only a fraction of possible sequence combinations are sampled, and each peptide is represented on a single bead only. The resulting libraries are said to be ‘under-sampled’ or ‘sparsely-sampled’; using more beads increases the number of unique peptides accessed, but not their individual quantities. Both library types will be discussed: low diversity libraries for benchmarking de novo sequencing (“Automated De Novo MS/MS Sequencing Provides Acceptable Sequencing Coverage of Peptide Mixtures, with Good Accuracy” section), and high diversity libraries for both benchmarking de novo sequencing and discovering binding molecules by affinity selection (“Single-Pass Affinity Selections Recover Binders from Random Peptide Libraries” section).

Originally, ‘split-mix’ synthesis was devised to avoid an issue with a simpler alternative for combinatorial synthesis, wherein a single portion of resin beads is treated with activated amino acid mixtures (Flynn et al. 1991; Ivanetich and Santi 1996). In this approach (originally used to prepare degenerate oligonucleotides) (Ike et al. 1983), every possible sequence combination is sampled, but in unequal proportion, due to differences in amino acid coupling rates. ‘Split-mix’ synthesis avoids this issue, since couplings occur in distinct vessels, such that competition between amino acids is avoided. Additionally, ‘split-mix’ synthesis provides a second important advantage: the ability to access libraries of high theoretical diversity by ‘sparse’ sampling, while still generating a sufficient quantity of each peptide for downstream use. In contrast, libraries prepared by coupling of reagent mixtures sample all sequence combinations, but in diminishingly useful quantities.

Figure 2c illustrates the number of beads per gram of resin, at different particle sizes, and the quantities of peptide contained in single beads. For context, low femtomole sample quantities have long been accessible to commercial mass spectrometers (Wilm et al. 1996; McCormack et al. 1997), suggesting that even the smallest commercial beads generate sufficient quantities of individual peptides for affinity selection and tandem MS sequencing. Using 10 micron beads at a reasonable lab scale (e.g. 1–10 g of beads), \(10^8–10^{10}\) unique peptides could be accessed, each in sufficient quantity for at least 10 affinity selections (based on a selection scale of 10 fmol/peptide).

Automated De Novo MS/MS Sequencing Provides Acceptable Sequencing Coverage of Peptide Mixtures, with Good Accuracy

As described elsewhere (McCormack et al. 1997; Eng et al. 1994; Steen and Mann 2004; Coon et al. 2005), mass spectrometry is ideally-suited for analysis of peptide mixtures, because of the ease with which individual ions can be isolated based on unique mass to charge ratio. When coupled to liquid chromatography, which is conveniently interfaced with electrospray ionization, an additional separation dimension is achieved, enabling individual ions to be isolated from mixtures containing many thousands of unique peptides. Once isolated, individual peptide ions are fragmented into ‘daughter ions’, which are analyzed as composite spectra in MS/MS. The differences between peaks in the resulting MS/MS spectra correspond to amino acid residue masses, characteristic of the peptide sequence.

In proteomics, peptide sequences are typically inferred from MS/MS spectra by the method of database searching (Eng et al. 1994; Domon and Aebersold 2006; Zhang et al. 2013; Sinitcyn et al. 2018). In this approach, experimental spectra are matched to predicted spectra generated from sequence databases (Eng et al. 1994), or to sequence-annotated spectral libraries (Lam et al. 2007). Database searching routinely matches a high proportion of MS/MS spectra (50% or more) (Cox and Mann 2008; Michalski et al. 2011; Senko et al. 2013; Hebert et al. 2014) with high accuracy (e.g., such that ~99% of peptide-spectrum matches are correct, at a false discovery rate of 1%) (Elias and Gygi 2007; Nesvizhskii et al. 2007; Marx et al. 2013). The approach is suitable when relevant databases are available, and contain a reasonable number of largely unrelated peptides. However, it is less suitable for random peptide libraries, which comprise trillions of possible sequences with significant overlap in local sequence. De novo sequencing (Taylor and Johnson 2001; Seidler et al. 2010; Hunt et al. 1992) is an alternative approach to interpreting MS/MS spectra that is better-suited to peptide libraries, since it is applicable in principle to any peptide sequence, provided complete series of fragment ions can be assigned (with the exception that Ile and Leu are not routinely distinguished). But compared to database searching, the reliability of automated de novo sequencing continues to be viewed with skepticism (Muth et al. 2018; O’Bryon et al. 2020).

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1 In the early days, it was considered important for synthetic libraries to achieve complete sampling. But given that some of the earliest phage libraries were under-sampled, and the relative success of the phage approach, this point seems less important in retrospect.
In this section, we discuss evidence regarding the performance of automated de novo sequencing, and its suitability for analyzing peptide mixtures of unknown complexity. Broadly, performance evaluation involves performing de novo sequencing on a set of MS² spectra (often from a data repository), and comparing the resulting sequence assignments to reference peptides associated with the spectra (‘ground truth’ peptides). Typically, ground truth peptides have been taken as peptide-spectrum matches from database searching. As a complimentary approach, we focus on the use of synthetic libraries as a source of ground truth peptides. For both approaches, emphasis is placed on the value of precision-recall analysis (Davis and Goadrich 2006)—a concept borrowed from information science—for evaluating sequencing performance, and on the use of confidence score thresholds for obtaining sequencing results of improved accuracy.

### Precision and Recall Applied to De Novo Sequencing

In general terms, precision and recall report on the quality and quantity, respectively, of retrieved data. For de novo sequencing, recall is the fraction of ground truth peptides that are correctly sequenced, and precision is the fraction of de novo sequence assignments that are accurate (in other words, the fraction of total de novo assignments that correspond to ground truth peptides) (Fig. 3a). In some scenarios, precision and recall are equivalent—for example, when just a single spectrum/de novo assignment are associated with each ground truth peptide (Fig. 3b), and when all de novo assignments are considered regardless of confidence score. But in other scenarios—such as when the number of spectra/de novo assignments exceeds the number of ground truth peptides (Fig. 3c), or when some de novo assignments are excluded based on a confidence score threshold—precision and recall will differ.

By considering de novo sequence assignments with highest confidence scores only, sequencing precision may be improved, at the expense of recall. This tradeoff can be visualized in a precision-recall curve, where precision and recall are plotted as a function of confidence score. Precision-recall analysis has been applied to de novo sequencing, but generally at the level of amino acids only (Ma 2015; Tran et al. 2017; Yang et al. 2019). Recently, the analysis was extended...
to the level of peptide sequences (Vinogradov et al. 2017; Devabhaktuni and Elias 2016; Miller et al. 2018). This type of analysis is of considerable practical importance, since it provides guidance on appropriate confidence score thresholds to use when sequencing unknown samples, and the precision/recall to be expected at those thresholds. As discussed below, continued work along these lines will be important.

**Evaluating De Novo Sequencing: Ground Truth from Database Searching**

The use of test spectra with high-confidence sequence annotations is perhaps the best way to evaluate the performance of de novo sequencing algorithms, absent other factors (Fig. 3b). When evaluated in this fashion using proteome-scale datasets, and with ground truth peptides taken as peptide-spectrum matches from database searching, the best de novo algorithms have produced sequence assignments identical to database peptides for ~10–50% of matched spectra (Tran et al. 2017; Pevtsov et al. 2006; Muth and Renard 2018). This type of studies suggest that de novo sequencing performs significantly worse than database searching in terms of both precision and recall. However, they do not generally speak to the precision that may be obtained by considering high-confidence assignments only.

A recent study demonstrated that with an appropriate confidence score threshold, the commercial software PEAKS recalled ~50% of accurate de novo assignments with a precision of 90% (Devabhaktuni and Elias 2016). As above, ground truth peptides were peptide-spectrum matches from database searching. The results clearly demonstrate the effectiveness of the PEAKS confidence score (‘average local confidence score’; ALC) at differentiating accurate assignments at the level of peptides, and of bringing the precision of de novo sequencing closer to that of database matching (99%). There is a tradeoff, as expected, with respect to recall. Based on a maximum recall of ~10–50% (above), and a reduction by ~half at 90% precision, a recall of ~5–25% can be expected, compared to database searching at 99% precision. Further studies of this kind would be helpful in generalizing the results for PEAKS, and in establishing similar guidelines for new software.

**Evaluating De Novo Sequencing: Ground Truth from Synthetic Libraries**

The performance of de novo sequencing on test spectra with matches from database searching may not reflect the performance on raw datasets, collected from peptide mixtures. Synthetic peptide libraries provide a means to test this situation, which is of direct relevance to affinity selection—mass spectrometry and differs in several respects from evaluation using database matches. Whereas for database matches each ground truth peptide is associated with a single test spectrum, for a library analysis the pairings of individual spectra and associated peptides are unknown, and the number of spectra may exceed the number of input peptides. For these reasons, de novo assignments are considered accurate if they match any ground truth peptide, rather than a specific ground truth peptide associated with each spectrum (Fig. 3c).

Recently, the performance of PEAKS was assessed by analysis of a high-diversity synthetic library (Vinogradov et al. 2017). The exact sequences of analyzed peptides were unknown, but the library design was constrained by alternating varied positions with distinct amino acid subsets, such that ground truth peptides were taken with some confidence as de novo sequence assignments consistent with the library design (Fig. 4a). Using design-compliant assignments as ground truth, precision and recall were then calculated for the partially-processed de novo output (all assignments of correct length and C-terminal identity, to simulate the degree of processing that would be possible for a fully random library with fixed C-terminus). Similar to above (“Evaluating De Novo Sequencing: Ground Truth from Database Searching” section), good recall was achieved at 90% precision (~60% here, vs. ~50% previously). In this case, 90% precision corresponded to confidence scores >75, which may provide guidance on the appropriate score threshold to use when analyzing unknown mixtures.

Here, we have applied a similar analysis to synthetic libraries of low diversity, where the sequences of analyzed peptides were known exactly. Two libraries were prepared by ‘combinatorial scanning’ of a K-Ras-binding peptide (Niida et al. 2017)—one of which contained a fixed C-term Lys, presumed to facilitate productive fragmentation for MS/MS (Fig. 4b). The two libraries were analyzed by nLC-MS (with cysteines in acetamide-modified form), and the total PEAKS de novo output was processed as above, keeping all assignments with correct length and fixed terminal residues for evaluation. Precision and recall were then calculated, using the known set of library sequences as ground truth. For the library containing C-term Lys, good recall was achieved at 90% precision (69%), in broad agreement with prior work. But for the second, lacking C-term Lys, significant recall was achieved at lower precisions only, and the maximum recall was lower (34%, at 59% precision). Further work is needed to determine the actual cause of this effect, which was reproduced on a second instrument (Fig. S1) and could potentially result from over-fragmentation of the non-Lys library (expected to fragment at lower energy) (Dongré et al. 1996).

Together, the results across three peptide libraries suggest that within individual datasets, PEAKS confidence scores are effective at differentiating correct de novo assignments, since for each library precision increased with increasing score threshold. However, the significance of confidence
scores is not absolute, since the precision achieved at a given score threshold varied considerably between libraries (for example, either 0.97, 0.94, or 0.80 at score threshold = 80; Fig. 4). Careful benchmarking will be important when applying de novo sequencing in new contexts (e.g., to peptoids), since a high confidence score alone does not imply assignment accuracy. But for libraries of relatively short canonical peptides (e.g. 10-mers), a sensible approach would be to keep design-compliant candidates above a score of 80, and to anticipate that at least 80% are accurate.

**Summary and Outlook—De Novo Sequencing**

Existing de novo MS/MS sequencing tools can provide acceptable coverage of synthetic peptide mixtures with good accuracy (~ 50% recall at ~ 90% precision), and with continued refinement, additional performance gains may be expected (Tran et al. 2019; Gessulat et al. 2019). At the same time, the approach is not yet competitive with peptide identification by database searching. Toward this goal, benchmarking should focus on precision at the peptide level, and consider the effect of assignments to all MS/MS spectra in raw datasets, rather than subsets of spectra only.

Synthetic libraries provide a powerful tool for benchmarking de novo sequencing, but not without limitation. The sequencing precisions obtained for synthetic libraries benefit from removing a priori incorrect assignments (e.g. of incorrect length, or lacking the correct C-terminus) prior to evaluation (Fig. S2). Where this processing is not warranted (e.g., when peptides of varying length are analyzed), precision may be lower. Additionally, precision and recall may be overestimated, if erroneous assignments happen to match ground truth peptides incidentally.

With respect to sequencing coverage, good recall was obtained for the mixture complexities studied here (~ 500 peptides per sample). More work to test the effect of mixture complexity on sequencing coverage would be helpful, over a wider range of complexities. As the recovery of de novo assignments is an indirect measure of the number of peptides present in a mixture, an orthogonal method to determine mixture complexity would be valuable, when analyzing unknown mixtures.

**Single-Pass Affinity Selections Recover Binders from Random Peptide Libraries**

As discussed in “Automated De Novo MS/MS Sequencing Provides Acceptable Sequencing Coverage of Peptide Mixtures, with Good Accuracy” section, the available data suggest that de novo sequencing provides adequate coverage and accuracy for mixtures containing hundreds to perhaps thousands of synthetic peptides. The next question

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**Fig. 4** De novo sequencing provides acceptable coverage of synthetic peptide mixtures, with good accuracy. Each point corresponds to the precision/recall values obtained as described, for de novo assignments above a varied score threshold (from 50 to 99). The precision and recall values at score threshold = 80 are indicated (black points). Where applicable, the minimum score thresholds that achieved 90% precision are specified. a Adapted with permission from Vinogradov, A. A. et al. Library Design–Facilitated High-Throughput Sequencing of Synthetic Peptide Libraries. *ACS Comb. Sci.* 19, 694–701 (2017). Vinogradov et al. 2017. Copyright 2017 American Chemical Society. The number of input peptides was controlled by counting beads from split-mix synthesis, prior to cleavage from the solid support (Vinogradov et al. 2017). b This work. Because the library diversity was sufficiently low, the entire library was analyzed.
is whether mixtures of this complexity can be obtained by affinity selection from high-diversity libraries, and whether de novo sequencing can be applied reliably when the complexity of selected mixtures is unknown. Equally important is whether affinity selections provide efficient recovery of binding molecules with high enrichment, such that rare binding motifs can be discerned among sequenced peptides.

Here, we discuss recent evidence that at least in some cases, single-stage affinity selections can provide sufficient enrichment to identify binders from libraries of random synthetic peptides. We focus on selections using magnetic bead affinity reagents (Quartararo et al. 2020; Pomplun et al. 2020; Pomplun et al. 2021), which have identified a number of interesting ligands from synthetic peptide libraries (Fig. 5) and have been used in phage display for many years (Fowlkes et al. 1992). Recent foundational results are highlighted, as well as new data that replicates a key benchmark: recovery of p53-like peptides by selection for MDM2 binding (Quartararo et al. 2020). The new results are encouraging, but differ from the original report in several important aspects. Possible strategies to further improve selection performance are discussed, including proof of concept data for a multi-stage selection that illustrate the feasibility of recovering femtomole quantities of individual peptides over two sequential selection steps. The need for a measure of peptide mixture complexity that is ‘orthogonal’ to de novo sequencing is also discussed, along with proof-of-concept data for a possible solution.

**Establishing Selection Performance as a Function of Library Diversity**

Recently, selections for binding to a monoclonal antibody were used to test the performance of affinity selections, using random 9-mer libraries of increasing diversity (from $2 \times 10^6$ to $2 \times 10^9$ peptides) (Quartararo et al. 2020). The results are foundational in several respects: (1) in demonstrating further the reliability of de novo sequencing, since the expected binding epitope was identified among selected peptides; (2) in establishing the possibility of achieving very high enrichment by single-pass affinity selection, since the vast majority of selected peptides contained the binding epitope; and (3) in formally testing the benefit of library diversities beyond those accessible to standard synthetic combinatorial approaches.

The recovery of sequenced epitope-containing peptides from libraries of $10^6$, $10^7$, $10^8$, and $10^9$ random 9-mers is reproduced in Fig. 6a. The results show that higher diversity is generally beneficial, since epitope-containing peptides were identified in proportion to diversity over a wide range ($2 \times 10^6$ to $2 \times 10^9$), with no loss in enrichment. So long as the recovery of binders is maintained, as it was in this range, then additional diversity will help to identify a rare...
binding motif, since the number of times a motif is sampled will increase with library diversity. Beyond diversity of $\sim 10^8$, however, recovery of epitope-containing peptides suffered (Fig. 6a). Similar results were obtained by selection for MDM2 binders from libraries of random 12-mers (Fig. 6b). Therefore, $\sim 10^8$ was taken as a practical limit to library diversity amenable to the ‘affinity selection–mass spectrometry’ approach, and subsequent applications focused on libraries of this size (Pomplun et al. 2020, 2021).

The underlying cause of poor recovery of sequenced peptides from libraries beyond $10^8$ members has not been definitively proven, and should be revisited, since its correction could lead to further gains in accessible diversity, and the chances of identifying rare binding motifs. It was hypothesized that low recovery was caused by ‘peptide interference’, when selections yielded more peptides than are compatible with tandem MS sequencing (Quartararo et al. 2020). This is a plausible hypothesis, but would be strengthened by a direct measurement of the number of retained peptides, independent of de novo sequencing. Absent such a measure, the recovery of de novo assignments cannot distinguish between changes in the recovery of binders by selection, and changes in the coverage of selected binders by de novo sequencing (changes in sequence recall).

Another open question is the absolute recovery of sequenced binders, based on the total number input (which can be estimated based on statistical motif frequencies, but is not known with certainty). The proportionality of sequenced epitope peptides and total input peptides establishes that relative recovery was constant across conditions (Fig. 6a), but does not speak to the fraction of epitope peptides that were recovered and sequenced from the individual libraries. As we will discuss below, there is reason to believe that absolute recoveries could be improved.

**Selections for MDM2 Binding: Replicating a Benchmark Experiment**

Based on findings from the anti-haemagglutinin system, and to reproduce an important result from phage display, recent work tested libraries of $2 \times 10^8$ random 12-mers in selections for MDM2 binding, with the goal of recovering p53-like peptides (Quartararo et al. 2020). Remarkably, these selections yielded just a handful of library design-compliant de novo assignments, with PEAKS software score $> 80$ (‘sequenced peptides’). About half of sequenced peptides contained the ‘FxxxWxxL’ motif, in essence reproducing the phage display outcome. But as for the anti-haemagglutinin system, the absolute recovery of peptides from these selections was unclear, as was their coverage by de novo sequencing.

We have replicated selections for MDM2 binding from synthetic libraries of similar design (Fig. 7). The results are encouraging, in that recovery of sequenced motif peptides was ~ tenfold higher than reported originally (23 ‘FxxxWxxL/V’ peptides here, vs. 2 previously). Of the motif peptides, many contained a tyrosine residue not present in the p53 protein but that emerges from phage display (‘FxxY-WxxL’) (Böttger et al. 1996). Additionally, many peptides containing a partial ‘FxxxW’ motif were identified, which may be legitimate binders (117 total). At the same time, the enrichment achieved by the selection was poorer than
reported previously, since p53-like peptides comprised a smaller fraction of total sequenced peptides (1666 here, vs. 5 previously). Potentially, some of the non-motif peptides may be true binders, but these would be without strong precedent from phage display (Böttger et al. 1996; Kay et al. 1998; Hu et al. 2007; Pazgier et al. 2009).

We have not identified a single causal variable that explains the differences between these and other recent results. However, the results here demonstrate that absolute recovery of binders in recent work was not optimal, and motivate continued refinement of selection procedures that will increase the odds of success in more challenging applications, where binders are less frequent in a library. The key challenge will be to improve the recovery of binders, ideally from libraries of $10^9$ or higher, while maintaining the high enrichment necessary to distinguish binding motifs from unrelated ‘background’.

### Quantifying Peptides Recovered by Selection

As discussed above, the number of sequenced peptides obtained by de novo sequencing may not correspond to the number actually present in a mixture. If the number of peptides becomes sufficiently high—as it may when library diversity is increased, or when a high degree of non-specific binding occurs—then sequencing coverage is expected to drop, as the isolation of individual precursor ions becomes more challenging and ‘composite’ tandem spectra interfere with de novo sequencing (Michalski et al. 2011). Therefore, a means of measuring sample complexity independent of de novo sequencing would be informative.

Primary MS ‘feature extraction’ is a potential solution, which extracts MS signals from raw data and could be used to estimate the number of peptides in a mixture, if features correspond reliably to peptides. We tested this approach using data from the MDM2 selection, and a commercial feature extraction algorithm. By comparing the extracted features to sequenced peptide precursor ions, the correspondence of features with sequenced peptides was evaluated. As shown in Fig. 8, extracted features significantly outnumbered precursors, and the majority were singly-charged. Therefore, when overlaid with precursor ions, little overlap was found, since MS$^2$ spectra were collected on ions of $z = 2$ and higher only. For higher charge states (especially $z = 3$ ions), extracted features clustered with sequenced peptide precursors, suggestive of correspondence. However, even for $z = 3$ ions, the total features still exceeded sequenced precursors by > tenfold. Similar results were found for a sample of known complexity, obtained by cleaving a ~2000 bead aliquot from ‘split-mix’ synthesis of a random 12-mer library (‘library QC’, Fig. S8).

The significance of extracted features was studied further by determining the proportion of features that could be ‘matched’ to sequenced peptide precursors ($z = 3$ ions only). To ensure that each feature matched to at most a single unique precursor, each list was filtered for uniqueness, such that precursors were internally unique within 2 ppm, and features within 4 ppm. Then, for each precursor, the feature list was searched for matches ($\Delta m/z < 2$ ppm). This analysis was repeated considering the most abundant features only, at each of 5 different peak area thresholds (Fig. 9).

At lower area thresholds, most of the sequenced precursors were identified among the features, demonstrating...
the ability of feature extraction to correctly identify sequenced peptides. However, since total features exceeded sequenced precursors, the vast majority of features remained unmatched. Of the highest-abundance features, a significant fraction matched with sequence peptide precursors (10–25%; Fig. 9), and the significance of these matches was supported by the agreement of retention times for matched feature-precursor pairs (Fig. S9). But even here, the proportion of matched features underestimated sequencing coverage for the ‘library QC’ sample, where the number of input peptides was known approximately. Based on the similar proportion of matched features between the ‘library QC’ and affinity selection samples, we find no evidence of dramatically different sequencing coverage, and conclude that de novo sequencing did not grossly underestimate the total peptides recovered by affinity selection. However, more work is warranted before feature extraction is used as a quantitative measure of sample complexity.

Possible Routes to Improved Enrichment—Parallel Selections with Unrelated Targets

In the anti-haemagglutinin system, it was shown that motif-containing peptides were recovered by the target of interest only, whereas non-motif peptides were also isolated by an unrelated selection target (Quartararo et al. 2020). Therefore, it was concluded that parallel selections might be useful in distinguishing specific vs. non-specific binders, and improving enrichment for binders at the level of data analysis. We re-investigate this topic here, using parallel selections between MDM2 and streptavidin.

Overlap between sequenced peptides from MDM2 and streptavidin selections is shown in Fig. 10a. In contrast to the anti-haemagglutinin system, the majority of total peptides isolated in the MDM2 conditions (> 90% of which lack a binding motif) were unique to MDM2 and not isolated by streptavidin. While some degree of overlap is evident, it is less than the overlap between replicate MDM2
selections. This result suggests that ‘subtracting’ peptides isolated by unrelated targets is unlikely to be a general strategy to improve enrichment for binders.

The overlap between replicate selections for MDM2 binding was examined further, to test whether motif-containing peptides were more likely to be isolated in replicate (Fig. 10b). Overlap between the total sequenced peptides from each replicate was visualized, alongside overlap between ‘FxxxW’ and ‘FxxxWxxL/V’ peptides. The similar degree of peptide overlap in each case demonstrates that peptides recovered from replicate selections are no more likely to contain a binding motif, compared to peptides isolated from a single replicate only. Therefore, when evaluating a selection outcome, isolation from replicates should not be an absolute criteria for confidence.

**Possible Routes to Improved Enrichment—Multi-stage Selections**

Two-stage selection procedures are commonly employed with DNA-encoded libraries (Goodnow et al. 2017), which are similar to synthetic libraries in that they cannot be propagated for sequential rounds of selection. Compared to DNA sequencing, however, the feasibility of tandem MS sequencing after two sequential selections (and accompanying material losses) is less clear. As a test of feasibility, we piloted a two-stage selection for MDM2 binding. The quantity of library employed was doubled, such that after the first selection, half of the eluate could be sequenced directly, and the remaining carried through a second selection stage. In this way, the enrichment for ‘FxxxWxxL’ peptides achieved at each stage could be compared.
The results of the two-stage selection are shown in Fig. 11. Encouragingly, ‘FxxWxxL’ peptides were identified after each stage, proving the approach’s feasibility. However, the enrichment conferred by stage 2 was only ~2.5-fold, compared to ~2000 for stage 1.² For the 10⁶-member library studied here, an optimized procedure would probably use less selection target in the second stage, to maintain stringency as the quantity of input peptides is decreased. Libraries of even higher diversity may be amenable to an optimized two-stage selection, if enrichment of ~10³ could be achieved in each stage (~10⁶ overall).

Summary and Outlook—Affinity Selections

Even with confidence in the performance of de novo sequencing under controlled conditions, its use in characterizing unknown mixtures—such as those generated by affinity selection from peptide libraries—remains an analytical challenge. The sophistication of commercial MS instrumentation is considerable, and the quantity of data generated necessitates its automated handling. However, when differences between experimental outcomes arise at the level of sequenced peptides, sound conclusions cannot be made without reference to the underlying data. As discussed, a reliable means of inferring sample complexity from primary MS data would be helpful, and should ideally supplement de novo sequencing. Additionally, analytical standards and metrics tailored specifically for de novo sequencing may help to standardize results across laboratories.

An optimal affinity selection would recover all of the binders present in a library, with high enrichment. The enrichment for MDM2 binders obtained here is in-line with expectation from phage display (~10⁴), but lower than a recent application of synthetic libraries (binders comprised 1% or 12% of total sequenced peptides after single stage selection here, compared to 40% previously). At the same time, many more sequenced binders were recovered here (up to 73 from a single stage selection, compared to only 2 previously). Optimizing selection performance in systems where the frequency of binders is measurable will be essential before applying the method in new contexts, especially where binders are less frequent. In our view, an optimized procedure will likely involve two sequential selections.

Conclusion

Affinity selections have long been an attractive means to examine the high-diversity peptide libraries accessible by ‘split-mix’ synthesis. As discussed here, automated de novo sequencing is a solution to the challenge of sequencing individual peptides from selected mixtures, and opens up new possibilities for examining synthetic libraries of diversities not conventionally accessible. Discovery of binding molecules from libraries of synthetic polymers is a topic of longstanding interest (Cho et al. 1993). However, absent a practical approach to examine such libraries for binding molecules with sufficient throughput, the ability of non-proteogenic structures (e.g. beta-amino acids, N-Me-amino acids) to fundamentally alter library ‘fitness’ has remained largely a question in principle. Going forward, ‘affinity selection–mass spectrometry’ is poised to make an impact.

‘Affinity selection–mass spectrometry’ (Muchiri and Breenen 2021; Prudent et al. 2021) accurately conveys the experimental workflow described here, and accordingly, the ‘AS–MS’ designation is not inappropriate. However, we note that discovery of binding molecules by selection from synthetic peptide mixtures is a longstanding goal, which predates the formal designation ‘AS–MS’. Because the enabling aspect of ‘AS–MS’ applied to peptide libraries is reliable de novo sequencing and not MS detection per se, and because a typical ‘AS–MS’ approach does not involve a general sequencing strategy, we suggest this term be used with caution. We look forward to continued progress in the space, driven by applications, unusual libraries, optimized methods, and continued progress in de novo sequencing.

Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s10989-022-10370-9.

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² By definition, enrichment is the fold-change in motif-peptide frequency that accompanies selection. The enrichment achieved in the second stage was measured directly, since the proportion of motif-peptides was determined both before and after selection (p53-like peptides increased in proportion from 12% of total sequenced peptides after stage 1, to 32% after stage 2). The enrichment achieved in the first stage was estimated based on the statistical frequency of motif-peptides in the library, and the measured proportion of sequenced motif-peptides after stage 1. The statistical frequency of ‘FxxYWxxL’ peptides was used, since most ‘F-W-L’ peptides from each stage contained a tyrosine, and was taken as 6.0 × 10⁻⁵ (based on a statistical frequency of 1.2 × 10⁻⁵ ‘FxxYWxxL’ 8-mer motifs, using 17 randomized residues, and 5 possible motif registers in a library of 12-mers).
Data Availability  The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

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