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Author: Jirka, Silvana

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Phage display screening without repetitious selection rounds

Peter A.C. 't Hoen, <u>Silvana M.G. Jirka</u>, Bradley R. ten Broeke, Erik A. Schultes, Begoña Aguilera*, Kar Him Pang*, Hans Heemskerk, Annemieke Aartsma-Rus, Gertjan B. van Ommen, Johan T. den Dunnen

From the Center for Human and Clinical Genetics and Leiden Genome Technology Center, Leiden University Medical Center, Leiden, The Netherlands, and *Prosensa Therapeutics B.V., Leiden, The Netherlands

Abstract

Phage display screenings are frequently employed to identify high affinity peptides or antibodies. While successful, phage display is a laborious technology and notorious for identification of false positive hits. To accelerate and improve the selection process, we have employed Illumina next generation sequencing to deeply characterize the Ph.D.-7™ M13 peptide phage display library before and after several rounds of biopanning on KS483 osteoblast cells. Sequencing of the naïve library after one round of amplification in bacteria identifies propagation advantage as an important source of false positive hits. Most importantly, our data show that deep sequencing of the phage pool after a first round of biopanning is already sufficient to identify positive phages. While traditional sequencing of a limited number of clones after one or two rounds of selection is uninformative, the required additional rounds of biopanning are associated with the risk of losing promising clones propagating slower than nonbinding phages. Confocal and live cell imaging confirm that our screen successfully selected a peptide with very high binding and uptake in osteoblasts. We conclude that next generation sequencing can significantly empower phage display screenings by accelerating the finding of specific binders and restraining the number of false positive hits.

1. Introduction

Phage display is a powerful technique for the identification of peptides, proteins or antibodies with affinity for a specific target [1]. Many different phage libraries have been created and used for different applications, ranging from libraries with short random peptide inserts, to cDNA and classical and VHH antibody libraries [2-4]. In several rounds of affinity selection or "biopanning", the phage library is gradually enriched for high affinity binders.

Traditionally, peptides enriched after several rounds of selection are identified by DNA sequencing of the inserts of a limited number (tens to hundreds) of clones. Depending on the sequence diversity remaining in the library after selection, the analysis of a such a limited number of clones does not necessarily result in the discovery of the most promising candidates. Moreover, phage display screenings are notorious for their identification of false positive hits. These emerge for two important reasons: binding to non-target related materials used during the selection (such as plastics or albumin) and propagation advantages [5]. A well-known example in the latter category is the greatly accelerated propagation of phages displaying the HAIYPRH peptide in the Ph.D.-7™ library, as a consequence of a mutation in the Shine-Dalgarno box of the phage protein gllp in this clone [6]. This peptide has been identified in at least 13 independent biopanning experiments [5]. To aid the identification of potential false positives, several

web-based tools have been constructed: PepBank can be used to search for peptides already published in other experiments [7], while SAROTUP searches for peptides binding to unintended materials [8].

With the advance of next generation sequencing (NGS), it is now possible to sequence millions of inserts in parallel [9;10]. Thus, NGS permits a more expedient and higher resolution characterization of the library [11-13]. In this paper, we use NGS to examine the contents and the enrichment process of the Ph.D.-7™ library, the most popular, commercially available phage library for peptide ligand screening. In this library, random 7-mer peptides are displayed at the tip of the pIII minor coat protein of the M13 phage. We employed NGS after each round of selection to carefully characterize the enrichment process and show that positive hits can already be found after one round of selection. By comparison of the content of different libraries and by sequencing of the naïve library, we have found an efficient way to discriminate true binders and false positives such as target-unrelated peptides.

2. Materials and Methods

Cell culture

KS483 cells (murine preosteoblast) were cultured in α-MEM (1x) with glutamax (Gibco BRL, Breda, the Netherlands) supplemented with 10% Fetal Bovine Serum (Gibco BRL, Breda, the Netherlands) and 1% penicillin/streptomycin (Gibco BRL, Breda, the Netherlands) under 5% $\rm CO_2$. Cells were passaged by 0.05% trypsin-EDTA (Gibco BRL, Breda, the Netherlands) treatment at 3-4 day intervals. The cultured cells were grown to subconfluency. For differentiation, KS483 cells were seeded at a density of 15,000 cells/cm² in a 8.6 cm petridish. Every 3-4 days the medium was changed. From day 4 of culture, full confluence was reached and L(+) ascorbic acid (50 μg/ml, VWR International, the Netherlands) was added to the culture medium. When compact cell nodules appeared (from day 11 of culture onward), β-glycerolphosphate (5 mmol/L, Fluca) was added to the culture medium. At day 18 of differentiation, the control cells were stained with 3% Alizarin red solution (Sigma–Aldrich, St. Louis, MO) to confirm successful differentiation.

Biopanning

A heptapeptide phage display library (Ph.D.- 7^{TM} Phage Display Peptide Library kit, New England Biolabs, Beverly, Massachusetts) was used for the *in vitro* biopanning experiments. KS483 cells, at 4 and 18 days of differentiation, were washed gently with phosphate buffered saline (PBS) and incubated with 5 ml of α -MEM, containing 0.1% (w/v) bovine serum albumin (BSA), for 1 hour at 37°C under 5% CO $_2$. The cells were gently washed once with 5 ml of PBS before adding the phage library. Phages

(2x 10¹¹) were added in 3 ml α-MEM containing 0.1% BSA. Cells were incubated with the phage for 1 hour at 37°C, while shaking at 70 rounds per minute. After the incubation, the cells were gently washed 6 times by incubating with 5 ml of ice cold α-MEM, containing 0.1% BSA, for 5 minutes. Subsequently, the cells were incubated for 10 minutes on ice with 3 ml of 0.1M HCl pH 2.2 to elute cell-surface bound phage. This solution was neutralized by addition of 0.6 ml 0.5M Tris. The cells were then lysed for 1 hour on ice in 3 ml of 30mM Tris.HCl, 1mM EDTA, pH8, to recover the cell-associated phage fraction. Phages from each fraction were titrated and amplified according to the manufacturer's protocol. Each subsequent round of selection employed 2x 10^{11} phage derived from the phage library recovered from the previous round.

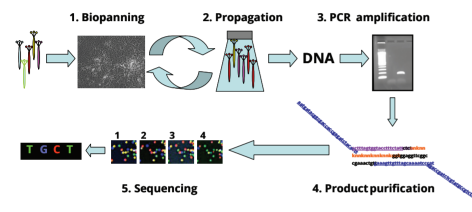


Figure 1: Overview of method. (1) In the biopanning phase, cells are incubated with the phage library. After washing away non-bound phages, binders are isolated and amplified in bacteria. The amplified libraries can be reused in a subsequent round of biopanning. (2) Alternatively, the DNA of the phages can be isolated and (3) amplified with PCR primers complementary to sequences flanking the variable region of the M13 phage DNA. The PCR primers contain tails with the adapter sequences necessary for Illumina sequencing. (4) A 160 bp fragment is purified from gel (lane 1: size marker; lane 2: negative PCR control; lane 3: phage DNA) and sequenced by the Illumina sequencer (5).

DNA preparation and sequencing

Sequencing was performed with the Illumina Whole Genome Analyzer WG2. Phage DNA was isolated from the amplified phage stocks and the naïve (unselected) library. For this, 10 μ l of a 1000 times diluted phage stocks were added to 1 ml of a 100 times diluted overnight culture of ER2738 bacteria and grown for 4.5 hours at 37°C while shaking at 200 rpm. Bacteria were centrifuged for 30 sec at 15700g. Then, 500 μ l of the top 80% of the supernatant was precipitated with 200 μ l of PEG/NaCl for 10 minutes at room temperature and the DNA was further isolated according to the manufacturer's protocol. The final pellet was dissolved in 25 μ l of milliQ

water and DNA concentration determined by Nanodrop before freezing at -20°C. Phage DNA was amplified with the following PCR primers:

Forward: AAT GAT ACG GCG ACC ACC GAG ATC TAC ACT TCC TTT AGT GGT ACC TTT CTA TTC TC*A

Reverse: CAA GCA GAA GAC GGC ATA CGA GAT CGG TCT ATG GGA TTT TGC TAA ACA ACT TT*C

where * indicates phosphorothioate bond. PCR primers used to amplify the phage DNA contained a subsequence that recognized the sequence flanking the 21 nucleotides long, unknown insert sequence and the adapters necessary for binding to the Illumina flow cell. The final product of the PCR was 160 base pairs long. The PCR protocol applied was the following: 1 ng of phage DNA was incubated with 2.625 U high fidelity Tag polymerase (Roche Diagnostics, The Netherlands), 20pM of primers in 1x High fidelity PCR buffer containing 15mM MgCl₂ and amplified for 20 cycles, each consisting of an incubation for 30 sec at 94°C, 30 sec at 60°C and 30 sec at 72°C. The PCR was stopped in exponential phase to mitigate PCR-induced sequence biases. The final PCR product was purified with the Qiaquick PCR purification kit (Qiagen, Valencia, CA). Concentrations as well as the correct length of the PCR product were established with an Agilent 2100 Bioanalyzer DNA 1000 assay. Each PCR product was applied to a single lane of an Illumina flow cell and subjected to solid phase amplification in the cluster station following the manufacturer's specification (Illumina, San Diego, CA). Single end sequencing for 27-35 cycles (27 cycles are sufficient but runs were sometimes extended to 35 cycles due to requirements for samples in other lanes of the same flow cell) was performed with a custom sequencing primer that started exactly at the first position of the unknown insert sequence (ACA CTT CCT TTA GTG GTA CCT TTC TAT TCT CAC TC*T)

Data analysis

All sequenced lanes were run through the initial Illumina Genome Analyzer Pipeline (Firecrest Bustard Gerald) for image analysis, quality control and base calling. Only sequences with the expected 6 nucleotide sequence after the insert (GGTGGA) were used (~95% of the sequences remaining after this filter step). DNA sequences were translated to amino acid sequences with a custom perl script using conventional amino acid codon tables. All reported numbers and sequences refer to the translated peptide sequences. *symbols indicate the presence of a stop codon, while – symbols indicate the presence of an unknown nucleotide in the triplet.

For plotting of phage abundance, a square root transformation was applied on the number of counts in the library, a commonly applied data transformation to stabilize the variance in count data [14].

Simulation of random clone picking

We randomly selected 50 amino acid sequences from the round 1, round 2, round 3, round 4 phage libraries selected for internalization into KS483 cells at day 18 of differentiation. We counted how frequently we selected the most abundant or 10 most abundant peptide sequences, as identified after complete sequencing of the round 4 library. We repeated this 20 times and report the average percentage and standard deviations for the 50 random picks in Figure 3D.

Peptide synthesis

7-mer peptides were synthesized by standard Fmoc solid phase peptide chemistry on a PS3 or Tribute Peptide synthesizer (Protein Technologies), using HCTU (5 eq.) as activating reagent and DIPEA (10 eq.) as base. Resin bound peptides were manually coupled to the fluorescent label by treatment with 5(6)-carboxyfluorescein N-hydroxysuccinimide ester and triethylamine in DMF. After cleavage [TFA/TIS/H $_2$ O 95/2.5/2.5 (v/v) or TFA/ thioanisol/TIS/H $_2$ O 90/2.5/2.5/5 (v/v) for peptide sequences containing Met], filtration, precipitation over cold-ether and centrifugation, crude FAM-labeled peptides were obtained. Peptides were purified by RP-HPLC on a Shimadzu Prominence HPLC [Alltima C $_{18}$ column (5mm, 10 x 250 mm); solvent A (0.1% TFA CH $_3$ CN/H $_2$ O 5/ 95 ; solvent B (0.1% TFA CH $_3$ CN /H $_2$ O 80/20)]. Peptides were analyzed by ESI-MS (positive mode) on a Agilent LC-ion trap mass spectrophotometer. Fluorescent peptide concentrations were determined by spectrophotometric analysis at 490 nm at pH=7.5.

Fluorescent imaging of KS483 cells

KS483 cells were seeded at a density of 15,000 cells/cm² in 6 wells plates with glass cover slides 21x26 mm (Menzel Glaser, Germany) or in Mattek glass bottom dishes with a diameter of 14mm and a glass thickness of 1.5 mm (Mattek corporation) in culture media described above. Cells were gently washed with PBS before adding 2.25 µM of FAM-labeled peptide in medium without serum for 24 or 48 hours. After incubation, cells at 4 days of differentiation were washed three times with PBS, fixed in ice cold methanol for 5 minutes and air dried for 5 minutes. Subsequently, the cells where embedded on microscope slides with Vectashield (Vector laboratories Inc.) and the slides where analyzed with a Leica TCS SP5 DMI6000 confocal microscope (Leica Microsystems, HCX PL APO 63×/1.4 oil-immersion objective, 8 bit resolution, 512×512 pixels, 400Hz speed). Cells at day 18 of differentiation were gently washed with PBS (three times) and supplied with fresh medium before analysis with live cell imaging using a Leica Af6000LX inverted microscope (Leica Microsystems, HCX PL FLUOTAR 63.0x1.25 oil-immersion objective, 12 bit resolution, 1392x1040 pixels).

3. Results

In the current study, we employed NGS technology to characterize the phage display screening process during successive rounds of selection. We used the combination of phage display and NGS to select for peptides that are binding to the surface of and/or internalized by KS483 cells in different stages of differentiation. KS483 are osteoblastic cells that can be efficiently differentiated into mature osteoblasts and form nodules depositing mineralized calcium material within 18 days [15;16]. The identified peptides may ultimately be used for the targeting of pharmaceutical formulations to bone or for enhancing the intracellular uptake of drugs into osteoblasts.

Sequencing of phage display libraries

The preparation of the phage libraries for sequencing is a simple and short procedure, as depicted in Figure 1. After DNA isolation from the entire phage pool, the fragment containing the insert was amplified and equipped with the linkers necessary for NGS in a one-step PCR reaction. The unknown inserts of 21 nucleotides (7 amino acids) were sequenced on the Illumina Whole Genome Analyzer. After quality control and translation of the DNA sequences to amino acid sequences, we obtained around 13 million peptide sequences for each phage display library (Table 1). This provides an ultradeep profiling of the content of the phage display libraries.

Table 1: Overview of sequencing results phage display experiments

| Selection Round | Differen- tiation | Internalized /surface | #sequences | #unique sequences | most abundant sequence | # counts for most abundant |
|--------------------|----------------------|--------------------------|------------|----------------------|------------------------|----------------------------|
| 3 | 4 days | surface | 11,192,802 | 274,666 | GETRAPL | 1,279,913 |
| 4 | 4 days | surface | 12,858,902 | 123,972 | GETRAPL | 3,113,643 |
| 3 | 4 days | internalized | 12,357,279 | 358,844 | GETRAPL | 397,669 |
| 3 | 18 days | surface | 13,146,497 | 196,615 | GETRAPL | 3,599,322 |
| 1 | 18 days | internalized | 15,595,055 | 1,913,785 | HAIYPRH | 41,257 |
| 2 | 18 days | internalized | 17,092,798 | 1,318,281 | GETRAPL | 547,210 |
| 3 | 18 days | internalized | 13,217,869 | 350,379 | RHEPPLA | 543,337 |
| 4 | 18 days | internalized | 13,880,199 | 282,266 | RHEPPLA | 1,413,696 |
| no selection | - | - | 6,688,401 | 3,887,498 | HAIYPRH | 36 |

Characterization of the naïve library

Before analyzing the phage display libraries selected against biological targets, we screened for potential sequence biases introduced by the propagation of the phages in bacteria by sequencing the naïve (unselected) library after one round of bacterial amplification. The library of 2x10¹¹ phages theoretically contains all of the 2x10⁹ heptapeptide sequences. By sequencing millions of peptides only a fraction of the entire library was analyzed. However, we confirmed that the peptide diversity in the

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library was high, since more than fifty percent of the peptide sequences in the naïve library were found only once (Table 1). Nevertheless, some peptides were found at higher frequencies than expected by chance. The peptide HAIYPRH was found most frequently (36 times) and is known for its accelerated propagation in phage display experiments due to a mutation in the Shine-Dalgarno box of the phage protein gllp [6]. Hence, growth advantages unrelated to the target selection emerge even after a single round of bacterial amplification of the naïve phage library. A high number of additional peptides were found more than twice in the naïve library, while the chance on this event based on a Poisson distribution with the current number of sequenced peptides would be only 5.5x10⁻⁹. Presumably, these peptides also have growth advantages. We provide a list of these nearly 700,000 peptides and their frequencies in the naïve library in Supplementary Table 1. It seems wise not to choose peptides that are found more than twice in the naïve library for follow-up studies, even when they demonstrate high enrichment after several rounds of selection.

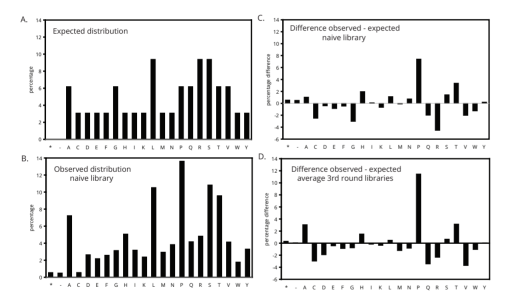


Figure 2: Analysis of amino acid composition of PhD7 library. Overview of the average amino acid composition (in %) of all unique peptides sequenced. A) Theoretical amino acid composition based on the translation of random (NNK)7 inserts. B) Observed composition in naïve library. C) Difference between the observed composition in the naïve library and the theoretical composition. D) Difference between the average composition in all 3rd round libraries and the theoretical composition. Standard amino acid one letter codes are used; *: one of the nucleotides in the triplet is unknown (N); -: stop codon.

Amino acid composition of peptides from naïve and enriched libraries

The naïve library is generated from a degenerate (NNK), oligonucleotide library where K represents an admixture of 50% T and 50% G. The expected amino acid frequencies resulting from this degenerate code is given in Figure 2A. Already after sequencing 70 random clones, the manufacturer noticed considerable differences between the actual and the expected amino acid composition of the naïve library (see manufacturer's documentation). A much more refined distribution of amino acid frequencies was obtained after sequencing >6 million inserts (Fig. 2B and C). The naïve library suffers from a considerable depletion of cysteine residues (frequency less than 1% of all amino acids). Glycine and arginine residues are also underrepresented but still found at frequencies higher than 2.5%. Proline is the most overrepresented amino acid. After selection, these trends were even stronger (Fig. 2D), suggesting that cysteine residues impede but proline residues enhance the propagation of phages. Some major positional effects were also observed (Supplementary Fig. 1): the most important is the restriction of the overrepresentation of proline residues to positions 2-7, while the first position is actually depleted of proline residues, as noted before [17].

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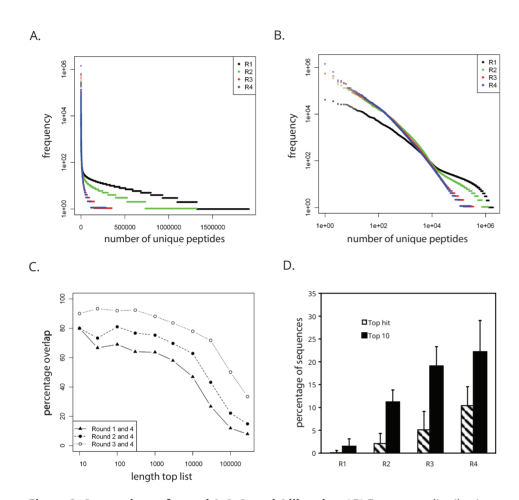


Figure 3: Comparison of round 1, 2, 3, and 4 libraries. AB) Frequency distribution (A: linear scale; B: 10log scale) of all unique peptide sequences (ordered from high to low abundance) after one (black), two (green), three (red), or four (blue) rounds of selection for phages internalizing into differentiated KS483 cells. The horizontal lines at the bottom of the plot represent the peptide sequences that occur once, twice, three times etc, with higher frequencies being more difficult to discriminate. The length of such a line represents the number of peptides with a particular frequency. C) Percentage overlap in the lists of most abundant peptides obtained after different rounds of selection. D) Simulation of traditional picking of 50 clones after round 1, 2, 3, or 4. The percentage of sequences belonging to the most abundant (hashed bars) or 10 most abundant (black bars) phages identified by next generation sequencing of the 4th round library are shown. Standard deviations refer to the variance observed in 20 independent random picks of 50 clones.

Characterization of the enrichment process

To identify phages with high binding affinity for undifferentiated and differentiated osteoblast (KS483) cells, we sequenced phage libraries

extracted from the surface of the cells. Internalizing phages were identified by the sequencing of libraries extracted from cell lysates after removing surface-bound phages. We initially sequenced libraries after 3 or 4 rounds of selection (Table 1), since we observed a significant enrichment, based on titration of output/input ratios, after these rounds (data not shown). To further characterize the enrichment process and the potential for identification of interesting phages from earlier rounds of selection, we also analyzed the phage libraries isolated after one, two, three and four rounds of selection for the internalization of phages on differentiated KS483 cells.

During subsequent rounds of selection the overall diversity decreased, while the frequency of the most enriched peptides steadily increased (Table 1, Fig. 3A and 3B). To illustrate this: the number of unique sequences decreased from 1,913,785 in round 1 to 282,266 in round 4, while the frequency of the highest abundant peptide increased from 0.26% to 10%, and the number of sequences that were found only once decreased from 594,379 to 170,592 (Fig. 3A and 3B). Thus, libraries converge towards certain peptide sequences in later rounds, consistent with expectations behind (traditional) phage display experiments.

There is a high correlation between the counts observed after the different selection rounds (Supplementary Fig. 2). In all comparisons, the correlations are high (Pearson correlation ranging from 0.47 (round 1 vs. 2) to 0.94 (round 3 vs. round 4)). Moreover, the correspondence in the top 1000 most abundant peptides between the different rounds is high, ranging from ~60% between round 1 and 4 to >80% between round 3 and 4 (Fig. 3C). Eight out of the ten most abundant peptides from round 4 are also in the top-10 after the first or second selection round (Fig. 3C). This means that, with the current sequencing depth, (i) further rounds of selection will not lead to the identification of peptides that could not have been found in earlier rounds; (ii) high affinity peptides can already be identified after the first round of selection.

We performed a simulation study to illustrate that this result is in striking contrast with a phage display experiment with traditional clone picking. In Figure 3D, we show that, after the fourth round, around 10% (say 5 out of 50 clones picked) would be derived from the most abundant phage, as identified by sequencing of the complete round 4 library. A further 12% of sequenced clones (say 6 out of 50 clones picked) would be derived from other phages from the top 10 of the complete round 4 library. When sequencing only 50 clones, these top 10 peptides would most likely be sequenced only once, and some of them would not be sequenced at all. The remainder of the 50 randomly picked sequences (78%) are from phages that are not found in the top10 after NGS analysis of the round 4 library. It is clear from Figure 3D that these results are even worse when randomly picking 50 clones from the round 3 library, when only 5% of the sequences

would be derived from the top phage, while sequencing 50 randomly selected phages after round 1 and 2 would be completely uninformative.

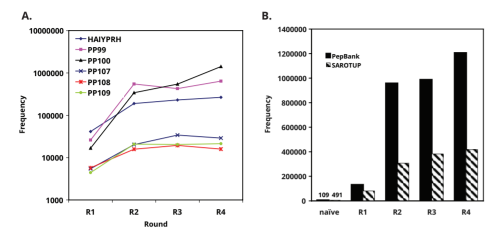


Figure 4: Counts of non-target and target-specific peptide sequences during subsequent rounds of selection. A) Counts for non-target specific peptides HAIYPRH, PP99, and PP100, and target-specific peptides PP107, PP108 and PP109 after four subsequent rounds of selection for phages internalizing into differentiated KS483 cells. B) Percentage of sequences recorded in PepBank (black bars, database of short peptide sequences previously reported in literature) and SAROTUP (hashed bars, database of peptides which bind unintended targets) identified after sequencing of the naïve library and the round 1, 2, 3, and 4 libraries.

Identification of false positives

The selection of one or a few sequences from a library of 10¹¹ molecules is an inherently noisy process. There are two principle causes for the high abundance of irrelevant peptide sequences in phage display experiments. On the one hand, phages may have lower than average replication time in bacterial hosts, illustrated by the HAIYPRH example. The impact of these propagation advantages increases with every round of selection and amplification. As an example, HAIYPRH was found 41,257 times after one round of selection and amplification (0.26% of all sequenced peptides) and 237,535 times after three rounds of selection (1.8%) (Fig. 4A). These false positives can be identified by sequencing of the naïve library after one round of amplification in bacteria. On the other hand, peptides may bind to non-target substrates and propagate during subsequent selection rounds without having any affinity to the intended target. These are less easy to identify, since their enrichment pattern may be similar to that of targetspecific peptides (Fig. 4A). A notorious example is the GETRAPL peptide, identified in many independent experiments [7]. Most likely, the peptide binds to plastics in general, since it was found at a frequency of 35% in phage display selection for polystyrene binding peptides and demonstrated considerable affinity for polystyrene [18]. GETRAPL was the most abundant in 6/9 of our libraries selected on KS483 cells and amongst the top ranked peptides in the other libraries. Libraries selected against *in vivo* targets did not have an overabundance of GETRAPL (data not shown), consistent with this peptide's presumed affinity for polystyrene plastics not present in *in vivo* screens. Two databases, PepBank [7] and SAROTUP [8], are very helpful to identify non-target specific peptides. Figure 4B shows that subsequent rounds of biopanning resulted in gradual enrichment for known target-unrelated peptides, amounting to ~10% of all sequences found after round 4. Through comparisons of many deep-sequenced libraries, it is possible to filter for commonly found parasitic peptides.

Table 2: Selected peptide sequences

| Group | Peptide sequence | Peptide number (PP) | Counts in naïve library | Counts in Day 4 - surface | Counts in Day 4 -internal- ized | Counts in Day 18 - surface | Counts in Day 18 - inter- nalized | SAROTUP | Pepbank |
|--|---------------------|---------------------------|-------------------------------|---------------------------------|--|----------------------------------|--|---------|---------|
| 0 - Control peptides | LPLTPLP | 98 | 16 | 13,006 | 29,175 | 20,988 | 20,003 | Yes | Yes |
| | GETRAPL | 99 | 15 | 1,279,913 | 397,669 | 3,599,322 | 427,153 | Yes | Yes |
| | RHEPPLA | 100 | 0 | 6 | 0 | 0 | 543,337 | No | No (*) |
| 1 - Abundant in all selected libraries | AMSSRSL | 101 | 0 | 15,301 | 54,629 | 11,517 | 134,645 | No | No |
| | YRAPWPP | 102 | 1 | 34,250 | 23,765 | 56,436 | 55,732 | No | No |
| | ASSSHRS | 103 | 0 | 73,075 | 5,033 | 57,476 | 24,902 | No | No |
| 2 - Surface specific | DLKIPLR | 104 | 0 | 37,802 | 0 | 55,342 | 59 | No | No |
| | IEFSPLM | 105 | 1 | 4,765 | 0 | 43,697 | 0 | No | No |
| 3 - Day 4 specific (Internalized) | NPWTTRP | 106 | 0 | 0 | 50,155 | 0 | 0 | No | No |
| 4 - Day 18 specific (Internalized) | ALPQIVR | 107 | 0 | 0 | 210 | 0 | 34,034 | No | No |
| | DERHQHY | 108 | 0 | 0 | 0 | 0 | 19,404 | No | No |
| | WQSVPTI | 109 | 0 | 0 | 0 | 0 | 20,390 | No | No |

Note: The following selection criteria were applied: Control peptides had more than two counts in the naïve library and were included in Pepbank or SAROTUP. Peptides in group 1 had more than 1000 counts in all round three libraries. Peptides in group 2 had more than 1000 counts in all surface but not in internalized libraries. The peptide in group 3 had more than 1000 counts, but only in the internalizing phages at day 4 of differentiation. Peptides in group 4 had more than 1000 counts in internalizing phages at day 18 only. (*) mentioned as a polystyrene binder in [18]

Selection of candidate peptide sequences

We were interested to identify peptides that specifically bind to and/or are internalized by KS483 cells. We applied the filter criteria described below to identify candidate peptide sequences from the different libraries, and searched for peptides with different apparent specificities. To exclude any putative false positives, we discarded all peptides with a frequency of 2 or higher in the naïve library and hits in Pepbank and SAROTUP. From the remaining list of peptides, we selected 9 putative target-related peptides (PP101-109) with apparent differences in specificities (Table 2), PP101-103 were most highly enriched in all libraries, while PP104-109 showed at least 100-fold differences in abundance between libraries and were selected for their apparent specificities. PP104-105 were selected as potential binders to the surface of cells at day 4 and day 18 of differentiated cells. We included peptides PP107-109 with apparent specificity towards fully differentiated (day 18) cells. These peptides were gradually enriched during subsequent rounds of selection (Fig. 4A), and were consistently ranked in the top 120. No other phages in the top 120 displayed similar degrees of specificity for internalization in cells at day 18 of differentiation. As negative controls, we selected three non-target related peptides (PP98-100): LPLTPLP (PP98) and GETRAPL (PP99) demonstrated a frequency of 2 or higher in the naïve library and had a hit in Pepbank. RHEPPLA (PP100) had been found to bind to polystyrene in an earlier study [18], but was unexpectedly only found in the libraries selected for internalization at day 18, where it was the most highly enriched peptide (Table 1), and not in any of the other libraries. The presumed non-target related peptides demonstrated similar enrichment profiles as PP107-109 (Fig. 4A). Unfortunately, solubility of PP98 and PP105 was too limited to allow further analysis.

Fluorescent imaging of peptide binding and uptake by KS483 cells

Selected peptides, equipped with a fluorescent FAM-label, were tested for binding to and uptake by KS483 cells at day 4 and day 18 of differentiation. The experiments with cells at day 4 of differentiation were performed three times with duplicated wells in each experiment, using cells with high and low passage numbers and incubation times of 24 and 48 hours, all generating similar results. Day 4 cells were analyzed with confocal microscopy to be able to discriminate between surface binding and internalizing peptides. Confocal microscopy was not possible for day 18 cells since the cells grow on top of each other and deposit thick matrix structures. Therefore, day 18 cells were analyzed by live cell imaging. Representative fluorescent images obtained with the two different technologies are shown in Figure 5 and Supplementary Figure 3. As expected, cells incubated with the PP99 control peptide, which has been found in many other phage display selection experiments according to SAROTUP and Pepbank, do not show any fluorescent signal. Similarly,

PP100, which was found to bind to polystyrene before [18], demonstrated only very weak cellular staining. PP102, which was highly abundant in all selected libraries, indeed displayed strong binding and uptake in day 4 and day 18 cells. Spectrophotometric analysis of the cell lysates confirmed that the majority of the presented peptides were internalized (data not shown). The intracellular distribution of PP102 differed between day 4 and day 18 cells, with more prominent staining of the nucleus on day 4 and more prominent staining of the extracellular matrix on day 18. Surprisingly, PP101 and PP103, also abundant in all selected libraries, displayed very weak or no signal. In line with the selection of PP104 as a surface binding peptide, PP104 demonstrated strong binding to the extracellular matrix. A similar staining pattern was observed for PP106, which was unanticipated since this peptide sequence was discovered in the pool of internalized phages. Peptides PP107 and PP108 were identified in day 18 cells only, and indeed appeared to give stronger staining in day 18 cells than in day 4 cells. Data for PP109 were inconclusive since the peptide was highly aggregation prone.

4. Discussion

In the studies described in the current paper, we have gained significant insight into the phage display selection process. This was achieved by sequencing millions of independent phages after different selection rounds using the latest next generation sequencing technology. We show that the enrichment factor of positive clones gradually increases during subsequent rounds of selection. We have not observed differences in the enrichment kinetics between target-specific and target-unrelated peptides. This is concordant with the observed absence of a correlation between peptide affinity and abundance in other reported phage display experiments [19].

Multiple rounds of selection will be helpful to reduce the background of non-binders and is therefore essential when sequencing a limited number of clones (*cf.* our simulation study in Fig. 3D). In contrast, significant enrichment can already be observed after the first round of selection when sequencing millions of clones. The high correlation between abundances after the first and subsequent rounds suggests that further rounds of enrichment may not have additional value. The large effects of small differences in propagation rates on the final abundances, nicely illustrated in ref. [19], may even result in exclusion or down-weighting of interesting phages from the pool. Instead of performing multiple rounds of selection, it may be preferred to perform the first round of selection in duplicate or triplicate, to account for variability in the stringency of binding conditions.

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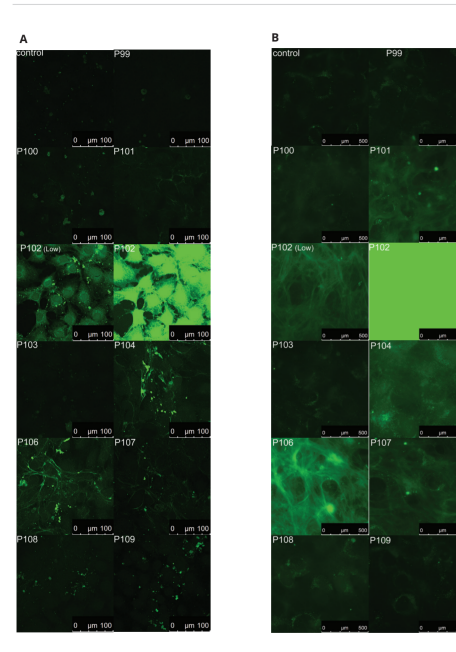


Figure 5: Uptake of fluorescently labeled peptides by KS483 cells. KS483 cells at day 4 (Panel A) or day 18 (Panel B) of differentiation were incubated with 2.25 μ M of FAM-labeled peptide in serum free medium and analyzed by confocal microscope (Panel A) or by live cell imaging with an inverted microscope (Panel B). Since the incubation with PP102 resulted in high uptake and very bright fluorescent, images obtained with two different gains (PanelA: 528 (also used for the other peptides) and 424 (labeled with LOW)) or exposure times (Panel B: 220 ms (also used for the other peptides) and 25 ms (labeled with LOW)) are shown.

Thus, our results question the validity of the traditional phage display screening approach: the high number of selection rounds required to arrive at peptides observed multiple times in tens to a few hundred of sequenced clones will come with the risk of an over-reduction in the library complexity and the loss of potentially interesting phages. Moreover, NGS-based characterization of phage display libraries is more cost effective than traditional clone picking because of the lower number of selection rounds required. It is also less laborious than methods previously developed to mitigate amplification biases employing phage amplification in isolated compartments such as monodisperse droplets [20].

Although full characterization of the first round library would require sequencing more than 10 million reads, identification of the most abundant phages requires substantially lower sequencing depth. The required sequencing depth depends on the complexity of the library and the enrichment factor. Generally speaking, sequencing of 1 million reads should be sufficient with relatively low enrichment factors obtained after one round of biopanning, since they would be represented >50 times according to simulated random draws of 1 million sequences from our round one library. This would allow for a high level of multiplexing of different phage display libraries in a single lane of the Illumina sequencer, and thus further reduce sequencing costs.

Previous characterization of phage display libraries were done with Roche 454 sequencing. Dias-Neto et al. sequenced around 70,000 phages isolated from human tissues after phage infusion [11]. In their experiment, the number of unique phages recovered was much lower than in our experiment - probably due to the distribution of the phages over different organs and the small biopsies taken - and sequencing of around 40,000 phages was sufficient to fully characterize the library. In an independent study, a cDNA phage display library selected for binding to transglutaminase was characterized by Roche 454 sequencing [21]. After sequencing of 120,000 phages, no decrease in diversity compared to the naïve library was observed. This may have been caused by a lack of high affinity binders in the phage pool and /or an insufficient sequencing depth. Illumina sequencing technology, like SOLiD and Helicos technologies, provides millions of reads. This may be an important reason why we observed a clear decrease in library complexity already after the first round of sequencing.

It is not yet feasible to sequence the complete naïve library, since it is claimed to contain 2 billion different phages. We sequenced a fraction of about 0.3%. Still, the sequencing of the naïve library has proven extremely useful for two reasons. First, it revealed non-random occurrences of the different amino acids at the different positions. We confirm the findings by Krumpe *et al.* obtained by random sequencing of 100-400 phages from

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similarly constructed CX7C and X12 M13 phage display libraries [17], and demonstrate that proline residues are overrepresented at all but the first position, and that cysteine residues are significantly underrepresented. Based on the fact that this was already apparent in the naïve library, which went through one round of amplification in bacteria, we tend to explain this by differences in availability of the amino acids, non-random incorporation probabilities during translation and suboptimal codon usage in bacteria. The non-random representation of amino acids reduces the chance to identify the highest affinity binders and therefore some optimization of the identified peptide sequences may still be useful. Second, sequences that were already found at higher frequencies than expected by chance are likely to have a selective or growth advantages. Indeed the GETRAPL peptide (PP99), which was found already 15 times in the naïve library, was not binding to KS483 cells, nor effective as a viral targeting peptide [22]. The lack of signal in our plastic dishes also suggests that GETRAPL is not a polystyrene binding peptide as previously claimed [18]. Its identification in a screening for plastic binding phages may have been caused by the same selection advantage. Moreover, the PCR amplification step applied during sample preparation may introduce biases that would be revealed by sequencing of the naïve library. We therefore strongly recommend to sequence naïve phage display libraries after one round of amplification in bacteria.

Despite our elimination of possible false positives caused by amplification or PCR biases or non-specific binding, other factors appear to contribute to the high false positive rate in peptide phage display screenings and were also observed in our experiment. A likely explanation for this is that the binding of synthetic 7-mer peptides may be largely different from the binding of a phage displaying a 7-mer peptide sequence as part of the phage plll protein, giving rise to substantial differences in secondary and tertiary structures.

We have identified one peptide (PP102) with exceptionally strong binding and uptake by KS483 cells and two peptides with strong binding to the extracellular matrix. Further research should demonstrate the potential of these peptides to target bioactive agents or drugs to osteoblasts and/or shuttle these across the plasma membrane. PP102 was ranked 14 after the first round of selection and remained in the top-20 during subsequent rounds of selection. Interestingly, there were two independent phage clones coding for the PP102 amino acid sequence: TATAGGGCTCCTTGGCCGCCT was the most prominent one followed by TATCGGGCTCCTTGGCCGCCT, which was observed at frequencies of approximately 10% of the former. The other selected peptides, which displayed less strong binding to KS483 cells, were not represented by multiple independent phages, since DNA variants were present at frequencies <1%. This suggests that the identification of independent phages displaying the same peptide has positive predictive

value for the affinity of the peptide. Clearly, more research evaluating larger sets of peptides needs to be done to confirm this.

The 1% threshold is our current estimate of the total sequencing error rate for a 21-mer sequence. This estimate is based on the steep increase of the number of DNA variants with a one mismatch difference at frequencies <=1% of the more abundant sequence. Another positive predictive feature would be the identification of related peptides with slightly different amino acid compositions. Single amino acid substitutions at frequencies lower than one percent can be explained by sequencing errors and should not be considered. However, for PP102 we found a second peptide, SRAPWPP, differing by just one amino acid, at DNA sequence-based frequencies ranging between 3 and 20% of the frequency of YRAPWPP (PP102). It proved difficult to perform a systematic search for clusters of related peptide sequences, since existing peptide sequence clustering software is not designed for the clustering of very large sets of very short peptides.

In our experiment, reads of only 21 nucleotides were required to identify unknown 7-mer peptide sequences. Other types of phage display libraries, like cDNA and antibody phage display libraries are also amenable to NGS-based characterization, aided by the much longer read length currently obtained with next generation sequencers. Likewise, NGS was recently applied to characterize the antibody variable regions of bone marrow plasma cells and this proved to be a very efficient method for monoclonal antibody selection [13]. Taken together, NGS can significantly empower phage display screenings and accelerate the finding of specific binders while restraining the number of false positive hits.

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