Acyclic Identification of Aptamers for Human alpha-Thrombin Using Over-Represented Libraries and Deep Sequencing

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Abstract

Background: Aptamers are oligonucleotides that bind proteins and other targets with high affinity and selectivity. Twenty years ago elements of natural selection were adapted to in vitro selection in order to distinguish aptamers among randomized sequence libraries. The primary bottleneck in traditional aptamer discovery is multiple cycles of in vitro evolution.

Methodology/Principal Findings: We show that over-representation of sequences in aptamer libraries and deep sequencing enables acyclic identification of aptamers. We demonstrated this by isolating a known family of aptamers for human α-thrombin. Aptamers were found within a library containing an average of 56,000 copies of each possible randomized 15mer sequence. The high affinity sequences were counted many times above the background in 2–6 million reads. Clustering analysis of sequences with more than 10 counts distinguished two sequence motifs with candidates at high abundance. Motif I contained the previously observed consensus 15mer, Thb1 (46,000 counts), and related variants with mostly G/T substitutions; secondary analysis showed that affinity for thrombin correlated with abundance ($K_d = 12 \text{ nM}$ for Thb1). The signal-to-noise ratio for this experiment was roughly 10,000:1 for Thb1. Motif II was unrelated to Thb1 with the leading candidate (29,000 counts) being a novel aptamer against hexose sugars in the storage and elution buffers for Concanavalin A ($K_d = 0.5 \mu \text{M}$ for α-methyl-manno-side); ConA was used to immobilize α-thrombin.

Conclusions/Significance: Over-representation together with deep sequencing can dramatically shorten the discovery process, distinguish aptamers having a wide range of affinity for the target, allow an exhaustive search of the sequence space within a simplified library, reduce the quantity of the target required, eliminate cycling artifacts, and should allow multiplexing of sequencing experiments and targets.

Introduction

DNA and RNA aptamers [1,2,3,4] have affinities for their targets similar to antibodies [5] and are useful in biosensors [6,7,8,9], diagnostics [5,10,11] and therapeutics [12,13,14]. Unlike antibodies, aptamers are not limited to binding immunogenic epitopes but have been reported to specifically bind an array of small organic molecules, macromolecules, and cells [3,4,9,15]. Aptamers have also been utilized as in vitro tools to detect and influence biological interactions in proteomics and metabolomics research [16,17]. In contrast to antibodies, aptamers can be prepared by standard solid-phase synthesis at a fraction of the cost for antibodies, have shelf lives of years, and require no animal or cell lines. In addition to their diverse functionality, aptamers are stable over a wide range of pH and temperatures, are not immunogenic and have shown successful protection from degradation by chemical protection [3,4,18]. Aptamers for several hundred targets have been described in the open literature [3,4] over the past 20 years and perhaps a few hundred more have been discovered in unpublished efforts. Far more antibodies have been found in the same time period. Some of this is due to the larger number of scientists who specialize in antibodies compared to those focused on aptamers. However, much of the disparity can be ascribed to the lengthy, cyclic methods in common use for aptamer discovery.

In vitro evolution [2,19], often called SELEX [1,20,21], is the standard method for aptamer discovery. SELEX, compared to our acyclic protocol in Figure 1, typically uses five to fifteen cycles of target-partitioning and amplification to enrich aptamer candidates from a pool containing randomized segments of length, $m$, where $m \geq 30$ is typical (Figure 1b).
Simpler and faster methods for aptamer discovery have been sought to shorten the rather complex cycle of enrichment of naïve pools with molecules that have high affinity for their target. Improvements include a robotic SELEX workstation to perform multiple selection cycles [22,23,24], reducing tedious sample manipulations. Efficient separation technologies have been applied, including capillary electrophoresis [25,26,27,28], monoLEX [15], which joins a selection step with column fractionation, and a microfluidic device [29]. These methods have been successful in reducing the discovery time from weeks to days. Other methods include photoSELEX, a method that covalently links high affinity binding sequences to their target of interest via a UV radiation-assisted photoreaction [30]. The work described in this report [31,32] and other recent papers [29,33] have applied deep sequencing to further reduce the cycling requirement in aptamer discovery.

Typical starting libraries for SELEX have oligomers with central randomized regions (30–70 nt) flanked by fixed regions needed for amplification and cloning (overall length often ≥ 90 nt). This is illustrated in Figure 1b for human α-thrombin, where the aptamer core sequence is only 15 bases, as originally deduced by Bock et al. [34]. They used five SELEX cycles, starting with 100 pmol of a 96mer DNA library with a 60 nt randomized region. Most possible sequences are not represented in naïve (unpartitioned) SELEX pools, e.g., in 100 pmol of a pool with m = 60, the probability is only 5 × 10^23 that a particular 60mer occurs at each randomized position. The use of such structured libraries can also avoid the necessity for truncating aptamer candidates to discover minimal core binding sequences.

We chose thrombin as a target to validate acyclic identification, and found the canonical DNA aptamer and lower affinity relatives produced.

Table 1. Representation of sequences in naïve pools.

| m | U | R (0.1) | R (25) |
|---|---|---------|--------|
| 5 | 1.0 × 10^3 | 5.9 × 10^10 | 1.5 × 10^13 |
| 10 | 1.0 × 10^6 | 5.7 × 10^7 | 1.4 × 10^13 |
| 15 | 5.0 × 10^10 | 2.2 × 10^6 | 5.0 × 10^10 |
| 18 | 6.9 × 10^9 | 5.0 × 10^8 | 7.0 × 10^8 |
| 19 | 2.7 × 10^11 | 2.2 × 10^9 | 2.7 × 10^9 |
| 20 | 1.1 × 10^12 | 5.5 × 10^9 | 1.1 × 10^9 |
| 21 | 4.4 × 10^12 | 5.0 × 10^9 | 4.4 × 10^9 |
| 22 | 1.8 × 10^13 | 3.4 × 10^9 | 1.8 × 10^9 |
| 23 | 7.0 × 10^13 | 8.6 × 10^9 | 7.0 × 10^9 |
| 24 | 2.8 × 10^14 | 2.1 × 10^9 | 2.8 × 10^9 |
| 25 | 1.1 × 10^15 | 5.3 × 10^9 | 1.1 × 10^9 |
| 26 | 4.5 × 10^15 | 1.3 × 10^9 | 4.5 × 10^9 |
| 30 | 1.2 × 10^18 | 5.2 × 10^8 | 1.2 × 10^8 |
| 40 | 1.2 × 10^19 | 5.0 × 10^7 | 1.2 × 10^7 |
| 60 | 1.2 × 10^20 | 4.5 × 10^6 | 1.2 × 10^6 |

a. See Table S1 for a more extensive list and details of calculations.
b. m = length of randomized sequence.
c. U = 4^m, the number of possible unique sequences.
d. R = the average representation of each unique molecule in a pool, for the indicated (0.1 or 25) nmol of total strands.
e. Beyond diversity limit; some possible sequences are absent.
f. Beyond synthesizer limit for 1 µmol scale synthesis; some sequences are not produced.
within a naive library that substantially over-represents each possible sequence. This library was also adapted to the sequencing platform we used (Illumina) with short read lengths. A natural result of the investigation was that all possible thrombin binding sequences within the sequence space encompassed by the m = 15 library were evaluated [31,35]. The acyclic protocol reduced the sample manipulations by a factor of six compared to [34], and required only standard methods in molecular biology coupled with deep sequencing that is now widely available.

**Results**

**Library design and optimization**

We designed a 39mer DNA library with a constant stem to present the m = 15 variable segment in the context of a hairpin loop (Figure 1a). A fixed-sequence stem was chosen to confer reasonable stability in partitioning (T_m = 70–75°C over the range of 0.1–0.3M NaCl for an unstructured 15mer hairpin loop), while maintaining efficient ligation and PCR. The four-base non-complementary tails shown in the figure were tested to ensure efficient ligation into a sequencing cassette adapted for the Illumina Genome Analyzer platform (GA, data not shown). The sequencer was capable of generating reads up to 36 bases (the read length increased after this work was completed).

Prior to partitioning with a protein target, we tested for bias in the distribution of the four bases in the randomized positions due to the details of library synthesis. A set of m = 6 stem-loops, as in Figure 1a, was “machine-mixed” using what should have been identical molar amounts of each amidite for each randomized loop site; GA-sequencing demonstrated considerable bias as evidenced by the proportion of homo-oligomers, G6:A6:T6:G6 = 0.63: 0.37: 0.0025: 0.0006, which is far from the desired mole fraction of 0.25 for each of these 6mers. This strong bias was corrected in a “hand-mixed” library, where the four phosphoramidites were mixed prior to synthesis and added from one port of the synthesizer (IDT, Coralville, IA). The resultant proportion still showed a slight bias against C-containing sequences, G6:A6:T6:G6 = 0.27: 0.24: 0.29: 0.20, but was gauged to be satisfactory for our purposes.

The possibility that PCR-artifacts might skew the distribution of sequences obtained from Illumina sequencing was also assessed. The data just presented for m = 6 also shows that there is little PCR-bias for or against loop sequences containing homo-oligomer runs of six. Another test for PCR-bias within the context of the loops in Figure 1a was conducted with four different specified m = 15 loop sequences mixed in the molar proportion: 1.00: 0.10: 0.010: 0.0010 with no partitioning against a target. The counts of 3.2 million sequenced clusters were proportional to the dose, 1.00: 0.11: 0.012: 0.0010, accurately representing the input population for these loops over three orders of magnitude. While bias in PCR and deep sequencing is known to occur, these experiments reduced concerns that adapter ligation, PCR, and bridge amplification might introduce a large bias in the sequencing results. However, PCR-artifacts associated with skipping parts of the loop sequence have been detected, as described later.

**Aptamer Selection and Identification**

Partitioning conditions used in this study were exactly as described by Bock et al. [34]. Early rounds of SELEX aim to prevent the accidental loss of high affinity aptamer candidates by avoiding competition between candidates for binding sites (typical libraries begin with single copies of any unique full-length sequence; see Tables 1 and S1). Thus, a starting ratio of 60:1 thrombin:DNA was used by Bock et al., and was used in the current study in order to preserve a direct comparison between SELEX and the acyclic approach introduced here.

High affinity candidates were partitioned, isolated and prepared for sequencing by ligation of Illumina sequencing adapters and PCR-amplification (Figure S1). Following sequencing, the variable regions of output reads were aligned with respect to the invariant stem and tail regions using a Perl script (Table S2). Typically 80–90% of the sequences matched closely enough to the invariant sequences and length of the variable region to merit designation as qualified reads. The base calls for the invariant regions in qualified reads exhibited >95% accuracy for each base position with regard to substitution, deletion, and insertion (Table S3). Output sequences were ranked by the number of times each was counted. The high efficiency of our acyclic procedure is evident in Table 2, which displays the summary statistics for experiment T1. 88% of the total reads passed qualification by the Perl script. Most sequences that are counted only once arise from molecules that have little affinity for the target but are accidentally carried forward in the partitioning step. 92% of the sequences in experiment T1 were counted 1 time only and only 8 out of the 1,728,220 unique sequences were counted more than 500 times. While the total number of counts and the number of sequences in each motif vary between experiments, the trends remain similar (Table S4).

Sequence alignment within the variable region was made using ClustalX and motif (phylogeny) diagrams were made with Drawtree. Analysis of all sequences from experiment T1 with counts ≥ 10 revealed two major sequence motifs (Figure 2a).

Table 3 collects data from this experiment on several members of each motif. Motif I contained the consensus G-quadruplex 15mer from Bock et al. (Thb1), which ranked highest occurring 46,444 times in 4.7×10^6 qualified reads; this motif also includes variants having mainly G, T substitutions, designated as motif Ia and low abundance variants having many A, C substitutions, designated as motif Ib. Motif II was unrelated to motif I, containing novel carbohydrate binding aptamer sequences related to Carb1 (counted 29,405 times). Three thrombin partitioning and sequencing experiments were conducted, each showing a similar set of sequences from the sequence motifs (see Table S4).

| Table 2. Summary of sequencing results for experiment T1. |
|--------------------------|--------------------------|
| Total reads              | 2,142,146                |
| Total qualified reads    | 1,959,748                |
| Unique qualified reads   | 1,728,220                |
| Count = 1                | 1,598,788                |
| Count = 2                | 117,721                  |
| Count = 3                | 10,909                   |
| Count = 4–10             | 1,194                    |
| Count = 11–25            | 43                       |
| Count = 26–50            | 18                       |
| Count = 51–100           | 22                       |
| Count = 101–250          | 13                       |
| Count = 251–500          | 3                        |
| Count>500               | 8                        |

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Acyclic Identification of Aptamers

Figure 2. Analysis of selected aptamer candidate sequences. A. Motif (phylogeny) tree of sequences with ≥10 counts (108 sequences), in the experiment summarized in Table 2. Motif I at the upper right (blue) binds thrombin, motif II at the lower right (red) binds carbohydrates, while “jump” sequences (black) at the left are PCR-artifacts. B. SPR analysis of top contenders from each motif. Thb1 is the top candidate in motif la (close relatives of Thb1) and binds strongly to thrombin (upper SPR trace, blue). The top candidate in motif lb (distant relatives of Thb1, see Table 3) was counted ~1,000 times less than Thb1 and binds weakly (green trace). Motif II also binds weakly to thrombin (red trace). The SPR signal, ΔR%, is the change in reflectivity of the chip surface in response to analyte.

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Validation of aptamer candidates

Affinity measurements for thrombin are reported from SPR (Surface Plasmon Resonance imaging) in Table 3 for selected aptamer candidates. SPR traces are shown in Figure 2b for thrombin complexes with the top candidate within each motif immobilized on the SPR chip. Within motif Ia a correlation existed between deep sequencing counts and affinity for thrombin (Figure 3a,b), suggesting that counts may be a useful proxy for affinity within an aptamer family. The estimated Kd for the α-thrombin-Thb1 complex (12 nM) was in the range of previous measurements [4]. Motifs Ib and II made weaker complexes with thrombin (Figure 2b).

Sugar binding aptamer candidate (Carb1)

The weak SPR response to thrombin by the leading sequence from motif II, even though it had the second highest count (29,405), prompted us to investigate the likelihood that the sequence was a ligation artifact or a PCR-champion. Comparative studies of ligation efficiencies and semi-quantitative real-time PCR with Thb1 suggested that Carb1 was not a PCR-champion (Figure S2). A quantitative comparison of ligation efficiency for Thb1 and Carb1 also showed no significant difference (data not shown). However, we noted that the ratio of counts for Carb1/Thb1 increased with increasing concentrations of hexose sugars present in the partitioning step (see Table S4). While Carb1 bound α-thrombin weakly, its SPR-response was high for the two hexose sugars in the storage and elution buffers for Concanavalin A (ConA) agarose, which was used to immobilize thrombin. This novel sequence bound both α-methyl mannoside (AMM) and glucose (Figure 4); Kd ~500 nM for AMM and ~1.4 μM for glucose), with the Carb1-AMM affinity ranking in the top third of aptamer-small molecule complexes [4]. Thb1 also has affinity for both sugars consistent with reports that G-rich sequences can be carbohydrate aptamers [36,37]. Low affinity of Carb1 for thrombin was confirmed by electrophoretic mobility shift assays (Figure 4c,d). The shifted band diminished in the presence of ConA, which competes for the glycosylated residues of thrombin, and disappeared on addition of glucose or AMM. Agarose is composed of galactopyranose units, which might also contribute to the selection of motif II, although negative selection was performed by passing the library through agarose cells were not determined.

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Table 3. Aptamer sequences and Kd values (in μM).

| Rankb | Sequence | Countc | Tbd | Glcd | AMMd |
|-------|----------|--------|-----|------|-------|
| Motif Ia (top sequence is Thb1) | | | | | |
| 1 | GGTGGTGCGTGGG | 46,444 | 0.012 | 2.5 | 0.78 |
| 3 | GGTGGTGCTGGTGG | 2,451 | 0.17 | | |
| 9 | GGTGGTGCGTGGG | 419 | 0.65 | | |
| 25 | GGTGGTGCGTGGG | 96 | 2.2 | | |
| Motif Ib (more distant Thb1 relatives) | | | | | |
| 51 | GGTGGTGCGTGGG | 53 | | | |
| 54 | GGTGGTGCGTGGG | 48 | | | |
| 56 | GGTGGTGCGTGGG | 42 | | | |
| 121 | GGTGGTGCGTGGG | 10 | | | |
| Motif II (top sequence is Carb1) | | | | | |
| 2 | GCCTGCTGCGCGCG | 29,405 | 1.4 | 0.52 | |
| 4 | GCCTGCTGCGCGCG | 1,040 | | | |
| 10 | GCCTGCTGCGCGCG | 354 | | | |
| 34 | GCCTGCTGCGCGCG | 80 | | | |

a. See Table 2 for counting statistics. Four sequences from each motif are shown: underlined bases in motifs I and II differ from the top candidate; bold fonts and capitals are used to highlight motif similarities.
b. Order of abundance in the deep sequencing results.
c. Counts for indicated sequence.
d. Kd values (in μM) measured by SPR for Tb = Thrombin, Glc = glucose, AMM = α-methyl mannoside (accuracy estimated at ±30%). Kd values for blank cells were not determined.

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Variation in the Thb1 core binding sequence

The consensus thrombin binding aptamer (Thb1) discovered by Bock et al. forms two stacked G-quartets connected by three loops; the minimal form that retains high activity has fifteen residues [34,38,39] (Figure 5). We aligned the first 54,140 counts of motif I (Table S5). The shifted band diminished in the presence of ConA, which competes for the glycosylated residues of thrombin, and disappeared on addition of glucose or AMM. Agarose is composed of galactopyranose units, which might also contribute to the selection of motif II, although negative selection was performed by passing the library through ConA-agarose without thrombin. Future work will distinguish the preference of Carb1 for various sugars.

Functional assay

As expected Thb1 strongly inhibited thrombin’s enzymatic activity [34], while variants had reduced performance in the order
of their counts (Figure 6; see Tables 3 and S4 for sequences of the 6 oligonucleotides used in the clotting assay). Structural data has established that the Thb1 G-quadruplex binds α-thrombin’s anion-binding exosite I, effectively inhibiting its protease activity [38,39,41]. Carb1 and its related motif II sequences had no observable effect on thrombin’s activity suggesting that its (weaker) anion-binding exosite I, effectively inhibiting its protease activity [38,39,41].

Estimation of the noise threshold

Clustal was also used to estimate the contribution of background noise to the sequence counts. Figure S3 shows tree diagrams at thresholds of count = 10 (used for Fig. 2a) down to count = 4 (at a count threshold of four, there were more than 1000 sequences, the upper limit for ClustalX, so not all sequences were included). As the threshold was reduced, more and more sequences were included in Motifs I and II, and a progressively larger number of sequences were uncorrelated to either Motif I or II. At a threshold of five, these uncorrelated “noise” sequences were roughly equal in population to the “signal” sequences in Motifs I and II. Going from a threshold of five to four, new signal sequences appeared, but the noise sequences became the overwhelming majority.

Discussion

Representation

There is the potential for confusion with regard to the word, sampling, which is often used to describe the occurrence of candidate sequences in an aptamer library, with the degree of sampling afforded by deep sequencing. We prefer to use sampling for the latter, i.e., to take a sample for sequencing. The term used here to describe the occurrence of unique sequences within an aptamer library is the average representation, R. The number of possible unique sequences, \( U \), in a pool with \( m \) randomized nucleotides is \( 4^m \) when four nucleotides occur at each randomized position in equal population.

For the \( 6 \times 10^3 \) molecules in 100 pmol of a library with \( m = 15 \), \( U = 1.1 \) billion unique molecules; \( R = 56,000 \) for each of these, making this an over-represented library (see Tables 1 and S1). For a library with \( m = 22 \) and \( R = 3.4 \), \( R \) is the average taken over a distribution of discrete objects so the largest number of sequences are present in three copies and nearly as many occur in four copies. Such a random distribution of 18 trillion unique molecules in a total of 60 trillion will have some sequences present in two and five copies, as well as other multiples. Poisson statistics predict that there will be some sequences present in zero copies—absent from the distribution. We refer to \( m = 22 \) as being beyond the diversity limit for 100 pmol of library; full representation of all sequences is unlikely. By contrast, Poisson statistics predict for \( m = 21 \) (\( R = 14 \)) that the overwhelming majority of possible sequences are present. As noted in the Introduction, \( R = 5 \times 10^{-23} \) when \( m = 60 \); thus, the original pool of Bock et al. [34] was very sparsely represented. In such a pool (whenever \( R < 1 \)) it is extremely unlikely for any sequence to occur in more than one copy.

At a pool size of 25 nmol, \( m = 26 \) is beyond the diversity limit (last column of Table 1). It is common to perform library synthesis on the 1,000 nmol scale so acyclic aptamer discovery with \( m = 25 \) should be practical if a sufficient amount of the target is available. However, the synthesizer limit is passed at \( m = 30 \) for a 1,000 nmol synthesis. That is, \( U = 1.2 \times 10^{18} \) and fewer molecules will be produced from synthesis at this scale (\( \sim 6 \times 10^{13} \)). While it is possible to increase the synthesis scale, it may become prohibitively expensive to supply enough target for acyclic selection and identification.

Sampling

A primary advantage of using deep sequencing in aptamer discovery is that many more sequences are sampled in the partitioned pool than when Sanger sequencing is utilized. We define the Sampling Ratio, \( SR \), as the ratio of total qualified reads to the total number of sequences in the pool. This is the fraction that was physically sampled; in the work presented here \( SR \) is taken as the fraction of the partitioned pool that was sampled.

Bock et al. [34] reported 0.01% recovery of DNA from the 100 pmol applied to thrombin immobilized on ConA-agarose beads in the first cycle of SELEX (determined using \( ^3 \)P-labeled DNA). Thus, about \( 6 \times 10^9 \) sequences survived partitioning from the initial pool of \( 6 \times 10^{13} \). As described above, the overwhelming majority of \( m = 60 \) sequences in the initial pool were present as single copies. If Bock et al. had sequenced 32 clones following cycle 1 (as they did after cycle 5) \( SR \) would have been \( 32/6 \times 10^9 = 5 \times 10^{-5} \), sampling one molecule per 200 million.

By contrast, the experiments reported in this paper have a much higher value for \( SR \). Assuming that partitioning has the same efficiency as for Bock et al., and for 2 million qualified GA reads for experiment T1 (Table 2), \( SR = 2 \times 10^9/6 \times 10^9 = 5 \times 10^{-4} \), about 1 molecule was sampled per 200,000 recovered after partitioning.

Signal to Noise

The counts for authentic aptamers are regarded as the signal in our experiment. We have shown that the signal is strongly
correlated with binding affinity (Figure 3). If the partitioning step is efficient, the signal for Thb1 should be proportional to the number of Thb1 sequences in the naïve pool. Relatives of Thb1 will have signals reduced by their less efficient retention by the Thb protein in partitioning. The same should hold for Carb1 and its relatives. On the other hand, random noise should arise primarily from sequences that are accidentally carried forward in the partitioning step. Most signal sequences will be correlated to Thb1 or Carb1,

Figure 4. Motif II sequences bind carbohydrate moieties. SPR analysis of (Carb1, red), Thb1 (blue) and an Aβ control (green) with A, α-methylmannoside (AMM) and B, glucose (glc; note the expanded vertical scale). C, D, Electrophoretic mobility shift analysis: The affinity of Carb1 for α-thrombin (αT) diminished on addition of Con-A, and disappeared on addition of glucose. DNA hairpins consistently had two bands in EMSAs, the upper (minor) band presumably due to a low concentration of dimerized hairpins.

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Figure 5. Folding of the canonical thrombin binding sequence. In Thb1 the 15mer hairpin loop folds into the G-quadruplex structure on the stem and tails shown in Figure 1.

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Figure 6. Effect of selected motif I aptamers on clotting activity of α-thrombin. Thrombin activity was measured in the presence of varying concentrations of selected sequences with the corresponding counts shown in the legend (the sequences are listed in Tables 3 and S4). Motif II sequences had no effect on the clotting times.

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while noise sequences should be uncorrelated to Thb1, Carbl, and themselves. The noise threshold is due to multiples of non-binding sequences that must be expected for molecules that are sampled randomly from the partitioned pool. As discussed under Sampling, above, there should be about $6 \times 10^9$ total sequences in this pool, not all unique because $U = 1.1 \times 10^8$ (Table 1). This distribution is skewed toward sequences having some affinity for the targets. We will show next that the skewness is small-only $\sim 2\%$ of the sample contains signal sequences. At $SR = 3 \times 10^{-4}$, the $2 \times 10^6$ qualified reads must contain some multiples of noise sequences. (In this analysis, both signal and noise sequences are assumed to be amplified equally by PCR prior to clustering on the GA flow cell. Although PCR may be biased against particular sequences or structural motifs, it is unlikely to strongly alter the distribution of millions of non-binding noise sequences that pass through a selection process by random chance. Neither is it likely to distort the distribution of signal sequences for the Thb1 family, as the counts have been shown to correlate with affinity for thrombin-see Figure 3.)

It is possible to make an empirical estimate of the noise threshold from the Clustal analysis. Most sequences with 9, 8, and 7 counts in experiment T1 are included within the Thb or Carb motif (Figure S3). More outliers become evident as the count is reduced. At a count of five, roughly half of the sequences are no longer classified within the Thb or Carb families, yet new signal sequences continue to accumulate. Uncorrelated sequences dominate at four counts and fewer. Thus, a count of five represents an upper bound on the noise threshold, and four counts is a lower bound.

We conclude that the signal/noise ratio in the thrombin-selection experiment, T1, was very high, specifically, $S/N \equiv 10,000$ (46,444 divided by 4 or 5). There were 89,761 total counts for all unique sequences with five counts or more; this group, constituting $1.9\%$ of all of the qualified reads, is clearly dominated by signal sequences. It is likely that some weaker-binding signal sequences occur at 4 counts and less, and that there are sequences that are uncorrelated to either Thb1 or Carbl that have low, but nonzero affinity. Thus, there may be more than 100,000 total signal sequences in the data set. Most of the unique sequences (1.6 million), were counted only once and most appear to be uncorrelated to Thb1 or Carbl. At 0.01% recovery [34], as discussed above, $6 \times 10^6$ sequences would have survived partitioning. If 100,000 signal sequences were present among the GA counts, correcting for SR gives $3 \times 10^5$ signal sequences after partitioning; this still leaves $5.7 \times 10^4$ uncorrelated sequences out of $6 \times 10^5$ total. It is not surprising that the majority of unique reads occur one time only.

**Over-representation limit**

It is interesting to estimate the limit in diversity beyond which aptamers can no longer be distinguished by simple counting. We have shown here that acyclic identification is robust for 100 pmol of over-represented libraries with $m = 15$ ($R = 56,000$). Due to Poisson noise, the $S/N$ must tend toward one before $R$ is reduced to one. Distinguishing aptamers by counting after acyclic identification cannot succeed when $S/N \approx 1$, and must instead rely on motif recognition and/or cyclic enrichment. Preliminary experiments where oligonucleotides were spiked at known concentrations suggested that acyclic identification can be achieved at $m = 18$ [42] and perhaps longer ($R = 880$ at $m = 18$ and 220 at $m = 19$ see Table 1). It should also be possible to find aptamers in larger pools, e.g., 25 nmol, where $m = 22$ to 23 have $R = 860$ and 210 (Table 1). Using these pools should reduce the signal 50 to 300-fold for an aptamer requiring the full diversity of $m$ sites, giving $S/N \sim 40$ if the experiment scales according to our thrombin results. The signal may increase for increasing values of $m$ for aptamers that do not require the full diversity of $m$ sites. For example, when $m = 16$, there are eight ways to fit the canonical 15mer Thb1 into a 16-base randomized region (four with any base at the 3'-side and four at the 3'-side), and four ways to increase each of the three loops by one base (see Figure 5). Thus, there should be twenty new ways to represent high affinity versions of Thb1 at $m = 16$, with only a four-fold reduction in all unique sequences in going from $m = 15$ to 16. This corresponds to a five-fold increase in high-affinity signal, and does not begin to consider all of the new lower affinity versions corresponding to low-count sequences in Figure S3. There is also reason to expect a lower noise threshold from more diverse libraries ($m > 18$). That is because the Poisson distribution predicts that fewer multiples of sequences will occur in these larger libraries, so fewer multiples will be randomly carried forward by partitioning. Therefore, the background could be lower than the estimate of four to five counts determined above. Experiments in progress aim to define practical over-representation limits for acyclic identification.

**Variants of Thb1**

The sequencing results were used to explore variability within the Thb motif (see Table S3). The first 6 bases, GGTTGG (Figure 5), were $> 99\%$ conserved. The largest variability occurred at G8, G10, G14 and G15. The G8 to T8 substitution caused a huge decrease in counts, so the canonical Thb1 is clearly optimal among 15mers. Variability at G14 and/or G15 may be compensated by using the 3'-neighboring stem base (also a G) to complete the G-quartet structure. This would come at the expense of one base-pair in the stem.

Other high-affinity DNA aptamers are known for thrombin that involve quadruplexes. These aptamers adhere to the same general motif outlined in Figure 5; most have longer loops, which are not encompassed within the present libraries.

**Jump sequences**

There are systematic artifacts resulting from deletion of a portion of the library, probably due to PCR. Figure S4 shows the most abundant artifact sequence that was a result of the deletion of parts of the library construct for the DNA $m = 15$ experiment. These jumps escape filtering by PERL scripts as adapter 2 has short stretches nearly identical to the authentic 3'fixed region (see Figure S4). Perl scripts discriminate against PCR-artifacts, which are a small proportion of the 2–5 million qualified reads. Other PCR-artifacts may exist that do not fit this pattern, so examination of sequence motifs within candidates and assaying for affinity and function are important.

**Comparison with SELEX**

The acronym, SELEX, is derived from the term, Systematic Evolution of Ligands by Exponential enrichment. Prominent is the word, evolution. We have described an acyclic procedure for distinguishing DNA sequences with high affinity for proteins or small molecules that does not rely on cyclic evolution.

The variable region within a library of $10^{14}$ molecules with a large value of $m$ will contain all possible sub-sequences below a certain threshold length ($\sim 22$ nt for 100 pmol of library-see Table 1). This has been used to argue that complete representation is unnecessary [19]. However, when each sequence is present only once, as in sparse representation, cyclic enrichment is mandatory before an aptamer can be distinguished by counting. It would be difficult to find a single copy of a high-affinity sequence among the
billions of sequences that are accidentally carried forward through the partitioning step.

By contrast, acyclic selection and identification allows the distinction of aptamers simply by counting. We have described how this approach easily located a known thrombin aptamer and lower-affinity variants. In other work, our lab has shown that this approach is compatible with RNA libraries and RNA/DNA chimeric libraries (Chen, L, JEC, MPM, and FNB, unpublished).

It can be difficult to distinguish the key binding elements in long aptamers because contact sites are dispersed in a folded structure that is not known in advance. This may require extensive post-SELEX optimization [20,43,44,45]. Furthermore, diversity in aptamers with long variable regions is often consumed in presenting the combining domain in a favorable context. This suggests that libraries with pre-defined secondary structures can simplify the process of determining the active binding form. It is easy to multiplex such libraries using deep sequencing, either before the partitioning step or in the sequencing step. Winning sequences can be distinguished by matching against the library templates, which have fixed bases and randomized regions of defined length.

There are advantages in combining results from sparse representation and over-representation. For instance, ordinary SELEX can guide construction of over-represented libraries to refine aptamer candidates by acyclic identification. A related item is that SELEX has found aptamers in long randomized regions; such aptamers may allow more widely dispersed contacts with proteins that could be important in some applications. Such long randomized regions are not amenable to acyclic identification as described in this paper. Techniques inspired by SELEX [15,46,47] that focus on highly efficient separations can be coupled to deep sequencing in either over-represented or sparsely-represented libraries. Further, over-represented and sparsely represented libraries can be mixed prior to partitioning. Acyclic identification is also compatible with many modified RNA nucleotides that are refractory to enzymatic replication of an evolving pool, but can be copied into DNA for sequencing, e.g., by reverse transcription.

Acyclic identification should afford new opportunities for multiplexing targets. In the absence of cyclic evolution, aptamers with high affinity do not compete against those with moderate or low affinity. Thus, it should be possible to mix multiple targets prior to partitioning against a library, and the aptamers matched to their individual targets in secondary screens. Multiplexing the targets was demonstrated here in the co-isolation of aptamers for a-thrombin and carbohydrates. The protein was consistently free of self-cleavage products. The primary carbohydrate binding sequence has Kd = 500 nM for a-thrombin and carbohydrates. The canonical thrombin aptamer was distinguished at a signal/noise ratio of ~10,000:1. More than 1,000 aptamer candidates were found with counts ≥4, and sequences similar to the canonical thrombin aptamers. Counts for thrombin aptamers correlated with affinity over at least a factor of 200 in Kd. The primary carbohydrate binding sequence has Kd = 500 nM for a-methyl mannoside. This affinity ranks it in the top third of aptamers discovered to date that bind small molecules.

Methods

Thrombin analysis by MALDI TOF mass spectrometry

Human a-thrombin was purchased from Haematologic Technologies (Essex Junction, VT) in 50% glycerol stocks. Before spotting it on the MALDI plate, the thrombin was dialyzed in 1 liter of PBS buffer twice at 4°C over 24 hours. MALDI TOF mass spectrometry was performed on a Bruker AutoFlex mass spectrometer operated in reflectron mode. Final spectra were the average of 50 shots/position at 10 different positions (Figure S5).

Aptamer selection

Partitioning, elution of thrombin-DNA complexes, and DNA extraction exactly followed Bock et al. [34]. The m = 15 DNA library was purchased from IDT (Coralville, IA) hand mixed to provide nearly equimolar amounts of the four bases at each degenerate position. 100 pmol of the m = 15 library in 1 mL of partitioning buffer (20 mM Tris-HCl, 140 mM NaCl, 5 mM KCl, 1 mM CaCl2 and 1 mM MgCl2 at pH 7.4) was subjected to negative selection in a 1 mL slurry of pre-equilibrated concanavalin-A beads (Pierce Biotechnology), recovering ~93% of the DNA. The DNA was then applied to 6 nmol of human a-thrombin (Haematologic Technologies) immobilized on 1 mL of Con-A beads equilibrated in partitioning buffer. The column was washed with partitioning buffer, and complexes were eluted with 0.1 M AMM. DNA was recovered by phenol extraction and ethanol precipitation. It is important to check the DNA recovery after negative selection on Con-A beads as some lots of beads were observed to bind more than 90% of the DNA. Even with high recovery after negative selection, some bead lots also exhibited lower recovery of Thb1 and related sequences. This lot-to-lot inconsistency was found with beads from GE Life Sciences as well as those from Pierce.

Preparation for deep sequencing

Refer to Figure S1 for an overview of preparation for sequencing, and the end of this section for the list of DNA sequences used.
The 3’ strands of the Illumina GA adapters were modified to include sequences complementary to the invariant regions of the m = 15 library. Adapters and complementary splints were added at 50 μM to the partitioned library, heated for 3 min at 90°C, ligated with T4 DNA ligase (New England Biolabs) at 25°C for 30 min, and the reaction terminated by heat denaturing the enzyme at 70°C for 3 min. The reaction was cooled to room temperature, the ligated library (which has two 45 bp double-stranded segments, see Figure S1) was extracted using a QIAquick PCR-purification kit, purified by 2% agarose gel electrophoresis, and stained with ethidium bromide (Figure S6a); visualization of a DNA ladder and the excess adapters helps to choose the region from which to excise the ~120 nt ligation product from the gel (Figure S6a). The DNA was extracted using a QIAGEN MiniElute Gel Extraction Kit. The library was then PCR-amplified using Pfu Turbo DNA polymerase (Stratagene) to extend the flanking regions for clustering on an Illumina flowcell. PCR-conditions were: (a) 2 min at 94°C, (b) 18 cycles of (1 min at 94°C, 1 min at 61°C, and 1 min at 72°C), (c) 10 min at 72°C. The PCR-product was purified using the QIAquick PCR-purification kit and size checked on a 2% agarose gel (Figure S6b) before sequencing by the Illumina GA.

Sequencing Data Analysis
A PERL script was used to identify sequence strings that closely matched the 5’- and 3’-fixed regions flanking the degenerate bases (F5-m-F3). Table S2 specifies the match regions within F5 and F3, which are adjacent to the central region of m bases. The match criteria were used to generate a file of Qualified reads for sequences with the desired length of central m-bases. An Nmer count file was generated to give the number and rank for each unique sequence.

Sequence Alignments
All sequences with a count ≥10 were aligned using ClustalX, and plotted using DRAWTREE.

SPR analysis
Aptamer affinities were measured using a GWC SPRimager®II (GWC Technologies, Inc.) and 16 and 25 SpotReady™ chips at 25°C. SPR data was acquired with V++ imaging software and analyzed in Microsoft Excel. Functionalizing a chip involved its immersion into a 1 nM solution of 8-amino-octanethiol (Dojindo Molecular Technologies, Inc.) in absolute ethanol at room temperature overnight. Rinsing was done with absolute ethanol and drying under nitrogen and incubation in 1 mM 4-(N-maleimidomethyl) cyclohexane-1-carboxylic 3-sulfo-n-hydroxy-succinimide ester (SSMCC) (Pierce Biotechnology) for an hour. Reduced 3’-thiolated oligonucleotides (2 mM) were then spotted in 5 replicates per sequence onto the SSMCC treated chip and allowed to react overnight, then washed with nuclease-free water and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen. The chip was blocked overnight with 4 mM mPEG-thiol (MW 1000) (Nanocs) to cap unreacted and dried under nitrogen.

Electrophoretic mobility shift assays
Aptamer candidates were heated at 95°C for 3 min and snap cooled on ice for 10 min before pre-incubation for 30 minutes with respective components as explained in each corresponding gel legend, in selection buffer. Rapide cooling was used for these hairpin sequences to limit dimerization involving the self-complementary stems. Samples were analyzed on native polyacrylamide gels (14% [w/v]) in 1X Tris-glycine running buffer at 100 V for 1 hour at 4°C. Immediately after electrophoresis, gels were stained with SYBR Gold nuclear acid stain (Invitrogen) for 1 hour, imaged and subsequently stained with Coomassie Brilliant Blue protein stain.

Thrombin functional assay
Clotting times were measured in duplicate using a mechanical fibrometer, Oatoclot 2 (Helena Laboratories). Normal human plasma and varying concentrations of DNA aptamer candidates (0.1 nM–700 nM) were incubated for 4 min at 37°C before adding α-thrombin diluted in selection buffer to a final thrombin concentration of 7.5 nM. The extent of thrombin inhibition was calculated from a standard curve generated by measuring the plasma clotting time versus thrombin concentration.

DNA sequences
All sequences are listed in 5’ to 3’ direction.
Note: Adapter 1 & 2 complements possess overhangs complementary to the constant stem and tail regions of the library from each direction. The forward PCR-primer also introduced a 5’-overhang complementary to an oligomer planted on the Illumina flowcell to anneal the amplified library for sequencing.

Supporting Information
Figure S1 Detailed experimental outline. A. Alpha thrombin was immobilized on concanavalin-A-agarose beads; a library of DNA hairpin loops was applied after negative selection.
against conA. After several wash steps, high affinity binding sequences were co-eluted with the alpha thrombin. The high affinity binding sequences were extracted by phenol and chloroform extraction and concentrated by ethanol precipitation. B. The 15mer library used against alpha thrombin was a hairpin library with the 15mer degenerate library region indicated as region m in red. C. After extraction the high affinity binding sequences had adapter constructs ligated as required by the Illumina sequencing platform. Split strands ensured proper ligation. D–F. The ligated, partitioned library was PCR-amplified to introduce a 5’ overhang that annealed the sequences to the complement immobilized on the Illumina flow cell. This was followed by bridge amplification on an Illumina cluster station prior to the sequencing by synthesis process. G. The first one or two base reads and reads after base 36 are less accurately determined than the rest; all experiments generated 2 to 5 million, 36mer reads.

Figure S2 Investigation of potential amplification bias in aptamer candidates by semi-quantitative real time PCR. A, B Analysis of PCR-amplification rates by gel electrophoresis and absorbance measurements at 260 nm. In panel A are six pairs of PCR time points for Thb1 (motif I) and Carb1 (motif II), respectively, in a 2% agarose gel (cycle numbers: 10, 14, 18, 22, 26, 30) and in B, are the 260 nm readings of a duplicate experiment.

Figure S3 Background determination in thrombin aptamer identification. Following deep sequencing, the occurrence of each sequence was determined and ranked. High occurrence sequences were aligned to determine conserved motifs. The panels show phylogenetic trees generated from sequences counted 10 times or higher, 9+/8+, 7+/6+, 5+ and about one thousand of 4+, respectively. Replicate experiments consistently had large numbers of uncorrelated sequences, starting at a count of 4 to 6.

Figure S4 Systematic artifacts in surviving sequences. The full adapter-ligated construct prior to PCR-amplification is shown in A. Highlighted in yellow are the stem and tails of the hairpin library designed for the 15mer thrombin experiment, while highlighted in green, the letters, N, signify the 15mer library loop region. The red bases are the flanking regions used by the Perl script to find qualifying reads. In the case of the most abundant systematic artifact, the Perl script recognized the 5’ flanking region but substituted the second flanking region with the adapter sequence underlined above, thereby presenting the first 15 bases of adapter 2 as the library region. This could only occur with the jump shown in dotted line and from the sequence shown in B.

Figure S5 Thrombin analysis by MALDI TOF mass spectrometry. Purity of α-thrombin was verified by MALDI TOF mass spectrometry prior to use in selection experiments and was consistently >90% pure with minimal degradation.

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Author Contributions
Conceived and designed the experiments: PNB MPM GVK JEC. Performed the experiments: GVK JEC. MPM. Analyzed the data: GVK JEC MPM PNB. Contributed reagents/materials/analysis tools: GVK JEC MPM PNB. Wrote the paper: GVK PNB.
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