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First report of the use of a Saxitoxin-Protein Conjugate to Develop a DNA Aptamer to a Small Molecule Toxin

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Abstract

Saxitoxin (STX) is a low molecular weight neurotoxin mainly produced by certain marine dinoflagellates that, along with its family of similarly related paralytic shellfish toxins, may cause the potentially fatal intoxication known as paralytic shellfish poisoning. Illness and fatality rates are low due to the effective monitoring programs that determine when toxins exceed the established regulatory action level and effectuate shellfish harvesting closures accordingly. Such monitoring programs rely on the ability to rapidly screen large volumes of samples. Many of the screening assays currently available employ antibodies or live animals. This research focused on developing an analytical recognition element that would eliminate the challenges associated with the limited availability of antibodies and the use of animals. Here we report the discovery of a DNA aptamer that targets STX. Concentration-dependent and selective binding of the aptamer to STX was determined using a surface plasmon resonance sensor. Not only does this work represent the first reported aptamer to STX, but also the first aptamer to any marine biotoxin. A novel strategy of using a toxin-protein conjugate for DNA aptamer selection was successfully implemented to overcome the challenges associated with aptamer selection to small molecules. Taking advantage of such an approach could lead to increased diversity and accessibility of aptamers to low molecular weight toxins, which could then be incorporated as analytical recognition elements in diagnostic assays for foodborne toxin detection. The selected STX aptamer sequence is provided here, making it available to any investigator for use in assay development for the detection of STX.
Keywords: aptamer, saxitoxin, marine toxin, paralytic shellfish toxins, surface plasmon resonance
1. Introduction

Saxitoxin (C_{10}H_{17}N_{7}O_{4}) is a low molecular weight toxin with a mass of 299.29 g mol\(^{-1}\). One of the most harmful marine toxins, saxitoxin (STX) is primarily produced by several dinoflagellates as well as certain cyanobacteria (Deeds et al., 2008). Along with dozens of related congeners, STX makes up a suite of neurotoxins, which is referred to as paralytic shellfish toxins (PSTs) and are responsible for the human syndrome paralytic shellfish poisoning (PSP; Etheridge, 2010). Molluscan bivalves represent the major vectors of PSTs to humans given their capacity to filter feed and accumulate high concentrations of the toxins; however, numerous non-traditional vectors such as crabs, lobster, and carnivorous snails have also been identified (Deeds et al., 2008). Despite seasonal toxic algal blooms in certain geographic regions, PSP cases are rare because of the effective biotoxin monitoring programs that have been implemented. In the United States, shellfish growing areas are closed to harvesting when toxins are found to be higher than the regulatory action level of 80 µg STX equivalents per 100 g tissue.

The longstanding regulatory method for testing overall PSP toxicity is the mouse bioassay (MBA, AOAC Official Method of Analysis [OMA] 959.08); however, it has received much criticism for ethical reasons as well as a host of other issues including poor specificity, high cost, and high variability (Etheridge, 2010). Thus, recent emphasis has been placed on developing screening assays for PSTs that could reduce animal usage and provide a means of rapid, high throughput testing. Screening assays generally rely on incorporating analyte recognition elements such as native receptors (i.e., sodium channels) and produced biorecognition molecules (i.e., antibodies). Yet numerous
challenges remain for existing analytical recognitions elements (notably antibodies), including limited availability, cost, and the continued use of animals for their production. This research sought to develop a new analytical recognition element for STX that was cost effective, did not employ the use of animals for production, and was widely available for use by the marine toxin community for assay development.

Aptamers are nucleic acids or peptides that are selected in vitro to bind to a target with high affinity and specificity due to the tertiary structure formed (Ellington and Szostak, 1990; Tuerk and Gold, 1990; Bunka and Stockley, 2006). These short ribonucleic acid (RNA) or single-stranded deoxyribonucleic acid (ssDNA) analytical recognition elements are discovered from a starting random library of $10^{12} - 10^{15}$ oligonucleotide sequences via an iterative process known as systematic evolution of ligands by exponential enrichment (SELEX; Ellington and Szostak, 1990; Tuerk and Gold, 1990). Separation of target bound and unbound oligonucleotides has been performed using strategies such as affinity chromatography (Liu and Stormo, 2005), filtration (Schneider et al., 1993), SPR sensors (Misono and Kumar, 2005), and magnetic bead immobilization (Bruno and Kiel, 2002; Stoltenburg et al., 2005). Selected aptamers have been demonstrated for use in assays and platforms that have traditionally employed antibodies as analytical recognition elements. Aptamers display several important advantages over antibodies, including stability, lower cost, ease of production, and batch-to-batch consistency (Jayasena, 1999; Tombelli et al., 2005; Song et al., 2012). Further, and perhaps most important, aptamers do not depend on animal usage (live animals, cell lines, or in vivo conditions), either for development or production, making
them an ethically favorable choice over antibodies. Another feature of aptamers is that they can be easily modified via simple chemical modifications without compromising their potential for binding to the target, thereby allowing for a wide range of applications and manipulation for use with a variety of platforms (Jayasena, 1999; Song et al., 2012).

Since their discovery, aptamers have been applied to different classes of targets: small molecules (including organic dyes, metals, drugs, carbohydrates, amino acids, nucleotides, and peptides), proteins (including enzymes, antibodies, gene regulatory factors, and lectins), as well as viruses and pathogenic bacteria (see reviews by Jayasena, 1999; Stoltenburg et al., 2007). Regardless of the class and size of the desired target, there are a number of molecular features that determine the ease with which aptamers with high specificity can be generated. The target should be present in high concentrations and with high purity. Ideal target candidates include aromatic compounds and hydrogen bond donors and acceptors (Stoltenburg et al., 2007).

Furthermore, the molecule must be amenable to the method used to separate target bound/unbound oligonucleotides. In this study, a hapten-carrier complex employed for STX antibody production was used for aptamer selection. This novel strategy of using a toxin-protein conjugate for DNA aptamer selection was implemented to overcome the challenges associated with discovering aptamers to low molecular weight targets, analogous to the use of hapten-carrier complexes for enhancing immunogenicity for antibody production against small molecules that alone are generally non-immunogenic. As such, we were able to take advantage of known chemistries for coupling the hapten-
carrier conjugate to magnetic beads for SELEX and subsequent protein digestion strategies for assessing protocol performance (Figure 1). Counter-selection was conducted to remove any ssDNA that bound to the protein carrier, leaving only those sequences specific to the STX target. To the best of our knowledge, this is the first example of a SELEX method employing a hapten-carrier conjugate to discover an aptamer.

The goal of this research was to use STX conjugated to a protein carrier (keyhole limpet hemocyanin [KLH]) to develop an ssDNA aptamer that would bind to STX. Here we demonstrate (1) discovery of such an analytical recognition element using a hapten-carrier conjugate, (2) concentration-dependent binding of the aptamer to surface-bound STX, (3) aptamer binding to free STX, and (4) selectivity of binding of STX to the discovered aptamer, compared to other aptamers and random ssDNA sequences. This novel approach to aptamer selection targeted for small molecule toxins offers a promising strategy that may be employed to generate a more extensive library of aptamers to marine toxins that can then be used for developing improved diagnostic assays.

2. Experimental

2.1 Development of toxin-protein conjugate

STX was conjugated to the carrier protein, keyhole limpet haemocyanin (KLH) via a modification of the Mannich reaction incorporating 2,2’-(ethylenedioxy)bis(ethylamine), or Jeffamine, as a spacer compound in a similar manner
to that reported previously for bovine serum albumin (Campbell et al., 2007). In brief, Jeffamine was conjugated initially to the KLH via amine coupling. KLH (10 mg) was dissolved in 2-(N-morpholino) ethanesulfonic acid (MES) buffer (0.5 mL, 0.05 M MES, 0.5M NaCl, pH 5). An aliquot (250 mL) of 1-ethyl-3-[3-dimethylaminopropyl]carbodiimide (EDC, 20 mg) and 8 mg of N-hydroxysuccinimide (NHS) dissolved in MES buffer was added to the KLH and mixed for 5 min at room temperature. Then Jeffamine (50 mL, 1M) was added and the mixture allowed to react for 3 h at 20°C. The Jeffamine-KLH conjugate was purified using a PD-10 column (GE Healthcare) versus PBS (pH 7.2). Jeffamine-KLH (1 mg), STX (200 mg) and 2.5% formaldehyde (80 mL) were reacted, followed by dialysis over 24 h in 3 x 4 L of 0.15 M saline solution.

2.2 STX-Jeffamine-KLH magnetic bead coupling and verification

Following the Dynabeads Co-Immunoprecipitation Kit (Life Technologies, Grand Island, NY) instructions, 30 µg of the STX-Jeffamine-KLH conjugate were covalently bound to 2 x 10^8 (3 mg) Dynabeads® M-270 Epoxy. After the coupling and washing steps, the coated beads were suspended at 6.7 x 10^5 beads/µl, or 10 µg/µl, in PBS-T (10 mM phosphate buffer, 2.7 mM KCl, 140 mM NaCl, 0.05% Tween-20, pH 7.4; Sigma, St. Louis, MO). A second batch of Dynabeads was coated with Jeffamine-KLH, as above, and was used for counter-selection.

To assess the success of the coupling procedure, forty microliters of ligand-coupled Dynabeads (10 µg/µl) were transferred to a microcentrifuge vial and washed 3×
with 100 µl of 50 mM ammonium bicarbonate (Sigma Aldrich, St. Louis, MO). The beads were resuspended in 50 µl of 9% acetonitrile in 50 mM ammonium bicarbonate. After the addition of 1 µg of the endoprotease trypsin, the reaction was allowed to proceed for 4 hours at 60 °C before being quenched with acetic acid (1% final concentration). Five micrograms of both standard KLH (Sigma) and STX-Jeffamine-KLH, as well as 400 µg of uncoupled Dynabeads were analyzed in the same fashion as positive and negative controls, respectively. The resultant tryptic peptides were analyzed by LC-MS/MS. Briefly, 10 µl of the peptide mixture was loaded onto a 0.15 x 100 mm nanoAcquity column (Water, Milford, MA) and washed with 90% mobile phase A (0.1 M acetic acid) at a flow rate of 1 µl/min. The bound peptides were eluted with the following gradient: 0 – 1 min, 20% mobile phase B (0.1 M acetic acid in acetonitrile); 1 – 10 min, 40% mobile phase B; 10 – 11 min, 70% mobile phase B. The eluate was analyzed by a linear ion trap (LTQ) mass spectrometer (Thermo Fisher, Waltham, MA) with a data dependent acquisition of 3 MS/MS scans following a survey scan.

2.3 Aptamer development (SELEX)

The method used by DeGrasse (2012), as adapted from Murphy et al. (2003), was followed to develop the STX aptamer. The library consisted of a central string of 40 randomized nucleotides that were flanked by defined primer binding regions necessary for polymerase chain reaction (PCR) amplification. The library template was synthesized (1 µmole scale) with machine mixing for the bases within the center random sequence domain, and purified using polyacrylamide gel electrophoresis (PAGE) by Integrated
DNA Technologies (IDT, Coralville, IA). Forward and reverse primers were also generated by IDT (Table 1), as were reverse primers with a 5’ biotin label.

Ten rounds of aptamer selection were conducted starting with 5 nmoles (~3 x 10^{15} different ssDNA sequences) of the library and 50 µl of STX-Jeffamine-KLH coupled beads (3.35 x 10^{7}, Table 2) and following the SELEX protocol of DeGrasse (2012). This included running two PCRs per round using the same primers. Counter-selection rounds were included using Jeffamine-KLH coupled beads (no STX) for 6 of the 10 rounds. Otherwise, with the exception of performing 10 rounds of selection instead of the 14 rounds conducted by DeGrasse (2012), all other details of the protocol were the same, including reagents, instruments, PCR conditions, and sequencing of final products (see below). All secondary PCR products were visualized, at the end of each round and again all together upon completion of the entire SELEX process, on a 4% E-Gel® (Life Technologies) with an E-Gel 25bp DNA ladder (Life Technologies).

After round 10, the PCR product was cloned using the One Shot® Top10 E. coli with the TOPO® TA cloning vector (Life Technologies) according to manufacturer instructions. The E. coli (50 µl) was plated and grown overnight on pre-warmed (37 °C) LB agar plates containing 100 µg/ml ampicillin. The plates with a few hundred colonies were sent to GENEWIZ (South Plainfield, NJ) where 50 colonies were randomly selected for Sanger sequencing using the T3 sequencing primer that was present in the TOPO TA vector. The software program GeneiousPro (Drummond et al., 2010) was used to remove the primer and plasmid sequences from the 42 successful sequences returned
from GENEWIZ. The trimmed 40 nt sequences were aligned in GeneiousPro and forced into zero gaps to directly compare each potential aptamer sequence nucleotide by nucleotide. Another alignment was generated in GeneiousPro using the ClustalW (Drummond et al., 2010) algorithm that allowed for gaps to find potential motifs among the sequences. The settings in GeneiousPro were: cost matrix set to IUB (a scoring matrix), a gap open penalty of 9, and a gap extend cost of 3.

The structures of the resulting sequences were compared using the program mfold (Zuker, 2003) with the following settings: linear DNA fold, folding temperature set to 24 °C, ionic conditions for Na⁺ and Mg²⁺ were 140 and 0 mM, respectively (Peyret, 2000), suboptimality was 5%, the window parameter was set to 2, and there was no set limit on the maximum distance between paired bases. If more than one structure was returned, the lowest ΔG in kcal/mol was selected for the comparison (SantaLucia, 1998).

2.4 STX aptamer (APT⁶CTX) binding evaluation with an SPR sensor

Standard laboratory materials for the SPR assays (e.g., sensor chips, amine coupling kit, and running buffers) were procured from Biacore (GE Healthcare, Piscataway, NJ). Nuclease free water (IDT) was used in the preparation of all reagents for SPR experiments. SPR running buffers (a. 10 mM HEPES, 150 mM NaCl, 3 mM EDTA, and 0.05% [v/v] surfactant P20 buffer [HBS-EP+] and b. 10 mM HEPES, 150 mM NaCl [HBS-N]) were prepared from 10× concentrates.

The STX dihydrochloride used for chip formation and inhibition experiments was the FDA reference standard, currently available from the National Institute of Standards
and Technology (81.0 μg/mL free base in 20% ethanol/80% water). APT\textsuperscript{STX1}, APT\textsuperscript{SEB1}, APT\textsuperscript{RAND1}, and APT\textsuperscript{RAND3} were synthesized by IDT (See Table 1 for sequences) for use in the SPR assay. The sequences APT\textsuperscript{RAND1} and APT\textsuperscript{RAND3} were produced using a random sequence generator (http://www.bioinformatics.org/sms/index.html).

2.4.1 STX aptamer (APT\textsuperscript{STX1}) binding to surface-bound STX

APT\textsuperscript{STX1} binding to STX was evaluated with a Biacore T200 surface plasmon resonance instrument (GE Healthcare). The STX sensor chip development and instrumentation parameters used herein were detailed in Yakes et al. (2011). Briefly, a CMS sensor chip was prepared with three active channels via EDC/NHS activation, Jeffamine coupling, ethanolamine blocking, and STX coupling as well as one reference channel via EDC/NHS activation and ethanolamine blocking. Sensor chip activity was verified with a standard STX inhibition assay (Yakes et al., 2011). The chip performed as expected in the STX inhibition assay with 8 μg/mL burro anti-STX yielding 80 resonance units (RU) and a decreasing response with a corresponding increase in STX concentration.

Lyophilized APT\textsuperscript{STX1} was reconstituted in HBS-EP+ to a concentration of 100 μM, and then diluted to 240 μg/mL, heated at 95 °C for 5 min, and cooled on ice for 10 min prior to use. This stock was serially diluted in HBS-EP+ to yield 120, 60, 30, 15, 7.5, 3.75, 1.88, 0.94, and 0.47 μg/mL of aptamer. A dilution of APT\textsuperscript{STX1} was injected over the sensor chip surface at 12 μL/min for 90 sec followed by a 30 sec dissociation period.

After regeneration was performed with 50 mM NaOH for 60 sec, the next analysis cycle
was performed. Binding analysis was evaluated using the reference channel subtracted data to account for bulk refractive index variations. Within the sensorgram, a 5 sec window, 15 sec post-sample injection stop, was used to quantify the amount of APT\textsuperscript{STX1} binding. Studies to determine specificity of the created sensor surface for the selected aptamer versus other ssDNA (APT\textsuperscript{SEB1}, APT\textsuperscript{RAND1}, and APT\textsuperscript{RAND3} each assayed at 10 µM) followed the same assay procedures. In addition, a modified domoic acid (DA) sensor chip (Traynor et al., 2006) was used to evaluate APT\textsuperscript{STX1} binding to a similar, small molecule marine toxin. Accurate chip performance was first ensured by evaluating the response with a DA antibody, and the DA substrate performed as expected in a DA inhibition assay with 1:200 anti-DA yielding 400 RU and a decreasing response with a corresponding increase in DA concentration. The STX aptamer (240 µg/mL) was exposed to the DA sensor chip for 90 sec at 12 µL/min. The curves were fit in GraphPad Prism (v. 5.02, La Jolla, CA) with a one-site, total binding model.

2.4.2 STX aptamer (APT\textsuperscript{STX1}) binding to free STX

To perform studies on the aptamer binding to free STX, an antibody sensor substrate in conjugation with a direct toxin assay was performed. To create the substrate, a CM7 sensor chip was activated with EDC/NHS, polyclonal anti-STX (a generous gift from Beacon Analytical Systems [Saco, ME]; 75 µg/mL in pH 4.5, 10 mM sodium acetate buffer) was coupled, and ethanolamine was used to block remaining active sites. In addition, one reference channel was incorporated via EDC/NHS activation and ethanolamine blocking.
After determining assay conditions (HBS-N buffer, 50 µL/min flow rate, 180 sec contact time, and regeneration with 10 mM glycine HCl, pH 3.0 for 30 sec at 30 µL/min), direct toxin assays were performed. First STX was injected over the sensor surface and then STX mixed with 50% of 240 µg/mL APT\textsuperscript{STX}, prepared as for previous SPR assays, HBS-N buffer, was flowed over the antibody substrate. Final concentrations of STX in solution were 10,000, 5,000, 2,500, and 1,250 ng/mL. Curve fits were performed in GraphPad Prism using the log(agonist) vs. response (three parameters) function.

3. Results and Discussion

3.1 STX-Jeffamine-KLH magnetic bead coupling and verification

The coupling of KLH and STX-Jeffamine-KLH to the M-270 epoxy coated Dynabeads was successful, as demonstrated by proteomic analysis. During the LC-MS/MS analysis of the standard KLH, an MS/MS spectrum was observed that resulted from the fragmentation of a peptide (811 m/z) that eluted from the column at 8.5 min (Figure 2A). This signature spectrum was also observed during the analysis of the Jeffamine-KLH-coupled and the STX-Jeffamine-KLH-coupled beads (Figure 2B and 2C, respectively). This spectrum was not observed in the negative control (uncoupled beads).

3.2 Aptamer development (SELEX)

Using the hapten-carrier conjugate as a target for SELEX also allowed us to apply an established protocol for discovering an aptamer to a protein target (DeGrasse, 2012).
to a small molecule target. This led to the efficient discovery of $\text{APT}^{\text{STX1}}$ in 10 rounds of selection. To remove candidate sequences that bound either directly to Jeffamine-KLH or to the solid support, counter-selection was aggressively employed. Before proceeding with the next round of selection, it was imperative to check the product. After each of the ten rounds of SELEX, the quality and size of the resultant enriched library (78 bases) was confirmed by running the secondary PCR product on a 4% agarose gel. The presence of a band with the proper molecular weight (~78 bases) was the cue to proceed with the next round. Figure 3 illustrates the secondary products from all rounds. The success of the SELEX protocol was measured by the enrichment of a few sequences from a vast library of $\sim 3 \times 10^{15}$ different ssDNA sequences, in this case 31 out of 42 sequences were identical as is discussed below.

Product from the final round of aptamer selection was sequenced to reveal the selected STX-binding aptamers. Of the fifty positive bacterial colonies selected for sequencing, 42 sequences were returned (84%). The sequences were trimmed to remove known plasmid and primer regions and were assessed for quality (i.e., proper length and sequence reaction confidence). The trimmed 40 nt sequences were aligned both with no gaps (Figure 4A) and with adjusted parameters that allowed for gaps using ClustalW (Figure 4B). Out of the 42 sequences, 31 sequences, labeled $\text{APT}^{\text{STX1}}$, were identical (73.8%) (Table 1 and Figure 4). The remaining 11 sequences were not significantly homologous to each other with a percent identity of 2.5% (Figure 4A). In fact, the only common base was at the first position; however, in the clustal alignment
(Figure 4B) a few common motifs were noted, but there were no more than 3 base runs and a small increase in percent identity to 7.7%.

When synthesizing APT$^{STX1}$ to evaluate binding, the priming sites were included, not only to enable PCR amplification, but also because the primers may play a role in structural stability or conformation. The sequence of APT$^{STX1}$ was reported as 5’- GGT ATT GAG GGT CGC ATC CCG TGG AAA CAT GTT CAT TGG GCG CAC TCC GCT TTC TGT AGA TGG CTC TAA CTC TCC TCT (40 nt consensus in bold). The lowest free energy secondary structure of APT$^{STX1}$ was predicted based on conditions during aptamer development (Figure 5). The estimated secondary structure for APT$^{STX1}$ revealed four stem and loop segments, two involving the primers. The secondary structures of the other 11 sequences were also predicted. Again, the sequences were not significantly homologous to each other or APT$^{STX1}$, but many (8) of the structures as predicted by mfold did include a stem and loop structure that could possibly play a role in aptamer function (data not shown).

3.3 Binding evaluation of APT$^{STX1}$ using SPR sensors

To demonstrate binding, varying concentrations of APT$^{STX1}$ were exposed to a well-established STX sensor chip (Yakes et al., 2011; Yakes et al., 2012). While this chip has been optimized for PST antibody evaluation, binding of APT$^{STX1}$ to the STX substrate was seen. APT$^{STX1}$ bound to the STX surface, with the change in RU directly correlating with the amount of APT$^{STX1}$ in solution (Figure 6). This increase in signal with an increase in solution APT$^{STX1}$ concentration was reproducible, with low error and no dependence
on sample order or injection. As a negative control, the APT\textsuperscript{STX1} was exposed to the reference channel, and no change in response was observed (data not shown); as such, non-specific binding of APT\textsuperscript{STX1} to the chip was ruled out.

To determine whether other ssDNA molecules would non-specifically bind to the sensor chip surface, several ssDNA sequences of the same length were exposed to the STX chip. APT\textsuperscript{SEB1}, APT\textsuperscript{RAND1}, and APT\textsuperscript{RAND3} did not exhibit binding to the STX sensor surface (data not shown), indicating that the STX sensor surface selectively bound to APT\textsuperscript{STX1}. A modified DA sensor chip was used to evaluate specificity of APT\textsuperscript{STX1} binding relative to another small molecule marine toxin that often co-occurs in regions where STX may be present. After accurate chip performance was verified using an anti-DA/DA inhibition assay, APT\textsuperscript{STX1} was exposed to the DA sensor surface. Binding to the substrate was not observed (i.e., approx. 1 RU Fc2-1 response), thus indicating that APT\textsuperscript{STX1} did not cross-react with DA (data not shown) and further supporting that the selected aptamer was specific to STX.

Finally, to show the selected aptamer has the ability to bind to free STX, an assay that detects STX from solution was developed. The highly-coupled antibody substrate (approx. 40,000 RU of anti-STX) has the ability to bind STX from solution and has an operating range of approx. 1,000 to 25,000 ng STX/mL with responses between 10 and 50 RU. When the STX solutions were pre-mixed with APT\textsuperscript{STX1} and then injected over the antibody surface, no binding of STX to the sensor surface was observed (Figure 7). The lack of response in the presence of the aptamer indicated that the STX in solution was
bound to the aptamer and was not available to bind to the antibody surface. Together, 
these data demonstrated that the selected aptamer not only binds to surface-bound 
STX, but also to free STX in solution. Such assay versatility, along with potential 
selectivity, suggests that aptamers may be able to outcompete antibodies in small 
marine toxin diagnostics. Efforts to design detection systems with the STX aptamer as 
well as expand our aptamer library are underway.

4. Conclusions

To our knowledge, we have discovered the first aptamer to a marine toxin and 
demonstrated that it binds selectively to STX in a concentration-dependent manner. For 
discovery of APT$_{STX1}^*$, we employed a novel approach of using a hapten-carrier conjugate 
as the SELEX target. Coupling a small molecule to a solid support often poses a challenge 
to aptamer development. In particular, it can be difficult to assess the efficiency of such 
a coupling reaction. To avoid such a challenge, we took advantage of a hapten-carrier 
(STX-Jeffamine-KLH) conjugate that was produced originally for antibody production. 
The conjugate essentially converted a small molecule (~299 Da) into a protein moiety 
(~350 kDa), thus allowing conventional surface chemistries (e.g., amine/epoxy reaction) 
to be used for coupling the conjugate to a solid support. Additionally, we were able to 
verify the coupling reaction to ensure that the paramagnetic beads were coated with 
the conjugate. This assessment was crucial given that the hapten-carrier reaction 
reduces the number of free amines on the carrier available for coupling to the epoxy 
active sites on the Dynabeads.
This study represented the first step in aptamer development for marine toxins and resulted in a promising strategy for continued aptamer selections to target marine toxins. These aptamers can be developed in a very short period of time (weeks compared to the months required for antibody production), are inexpensive to produce, and have the potential for implementation into a wide range of detection assays and platforms. While antibodies are often proprietary, this work reports the sequence of the aptamer selected to STX, thereby making it available to any researcher who wishes to employ it as an analytical recognition element in their own assay development. Future research will include discovering aptamers to other relevant PSTs and incorporating these analytical recognition elements into operational assays. It is anticipated that the use of aptamers in marine toxin detection methods could reduce or eliminate animal-based models, as well as the numerous challenges associated with antibody-based methods.

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Ethical statement

The authors declare that the study reported in the manuscript has been conducted in accordance with Elsevier ethical guidelines.

Conflict of interest statement

The authors declare no competing interests.
References


Figure captions:

Fig. 1: Schematic of aptamer development. The iterative process began with the STX-Jeffamine-KLH coupled magnetic beads being incubated with the random ssDNA library. Following incubation, bound versus unbound ssDNA were separated and then the bound ssDNA were eluted from the STX-Jeffamine-KLH coupled magnetic beads, replicated by PCR, and used as an enriched library for the next round of selection. During rounds 4-10 an additional step was included for counter-selection. Magnetic beads coupled to Jeffamine-KLH were incubated with the PCR product from the previous round. Only the ssDNA that did not bind to the beads coupled with Jeffamine-KLH were used for the next round of selection. This process consisted of a total of 10 rounds.

Fig. 2: LC-MS/MS confirmation of ligand coupling to Dynal M-270 expocy-coated beads. The MS/MS spectrum of parent ion 811 m/z was observed in (A) standard KLH, (B) KLH-coupled beads, and (C) STX-Jeffamine-KLH coupled beads. This spectrum was not observed with the negative control of uncoupled Dynabeads.

Fig. 3: Gel image of all 10 rounds of aptamer selection showing the products from the 2nd PCR. Lane 1 is the ladder (25bp-500bp), lane 2-11 are rounds 1-10 respectively, and lane 12 is a no template control.
Fig. 4: Alignments generated in GeneiousPro representing the 42 clonal sequences returned (with the primer and plasmid regions trimmed). Sequence 1 is APT\textsuperscript{STX1} and represents 31 clones with 100% matches to each other. All other recovered sequences are shown (with no identical matches). Common bases are marked with a darker square. Panel A represents a manually edited alignment where each base was forced into position with no gaps. Panel B is a ClustalW alignment allowed for gaps to explore potential motifs.

Fig. 5: Predicted folding and structure of APT\textsuperscript{STX1}, as generated by the mfold web server (Zuker, 2003). Bases that were similar in all 12 sequences are highlighted in gray.

Fig. 6: SPR calibration curves showing the concentration-dependent binding of the STX aptamer (APT\textsuperscript{STX1}). (A) Raw response (in RU) from flow cell 2 with reference subtraction (flow cell 1) for a concentration series of APT\textsuperscript{STX1} and (B) average normalized response from four replicate measurements. Curves were fit in GraphPad Prism with a one-site, total binding model.

Fig. 7: SPR calibration curves showing: (1) STX direct binding to an STX antibody sensor and (2) inhibition of STX binding when APT\textsuperscript{STX1} was added to the analyte solution prior to injection over the antibody substrate. Each data point is the
average response from two replicate measurements (standard deviation shown by the error bars) of blank subtracted data, normalized to STX 10,000 ng/mL.
Highlights:

- We present the first reported aptamer selected to a marine biotoxin (saxitoxin).
- A novel strategy of using a toxin-protein conjugate was utilized.
- Initial step towards new aptamers and their application to marine toxin assays.
Table 1: The primer and library sequences used in this study as well as the sequence of the STX aptamer (APT^{STX1}) discovered in this work. APT^{STX1} was reported in 31 out of 42 sequences. Also two random sequences (APT^{RAND1} and APT^{RAND3}) were generated and used for experiments to evaluate selectivity of discovered sequences to the target.

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CCT CAC TTA ACA GCC CGG TCA
TGC TGA GTG TTA TTA GCT CGA
TGG CTC TAA CTC TCC TCT

APT²⁵⁶
5’-GGT ATT GAG GGT CGC ATC 23
CAC TGG TCG TTG TTG TCT GTT
GTC TGT TAT GTT GTT TCG TGA
TGG CTC TAA CTC TCC TCT
Table 2: Values for the number of beads and incubation times used in each round. The values were modulated in later rounds to increase the stringency of the SELEX protocol.

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<td>$3.35 \times 10^7$</td>
<td>$6.7 \times 10^5$</td>
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10 selection rounds

Use PCR product for next round of selection

Bind DNA

Heat to elute bound DNA

Wash

Round 4-10, counter-selection bead

PCR product from previous round

Bind DNA

Remove beads with bound DNA, leftover goes into next selection round

KLH: Keyhole limpet hemocyanin (350-390 kDa)

ssDNA: STX (399Da); Jaffamine:
RT: 8.5 min
Parent Ion: 811 m/z

Standard KLH

Beads-KLH

Beads-KL H-STX

Relative Abundance

Mass (m/z)
### Table A

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**Legend:**

- **S1** to **S20**: Sequence positions from 1 to 20.
- **Clone 50**: Sequence consists of 20 nucleotides.
- **Clone 9**: Sequence consists of 20 nucleotides.
- **Clone 36**: Sequence consists of 20 nucleotides.
- **Clone 12**: Sequence consists of 20 nucleotides.
- **Clone 36**: Sequence consists of 20 nucleotides.