Development of RNA aptamers as molecular probes for HER2+ breast cancer study using cell-SELEX

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ABSTRACT

Introduction

A variety of novel drug delivery systems have been developed to improve the selectivity of anticancer agents. Targeted drug delivery systems exploit differences between cancer and normal cells.

Breast cancer is one of the most common female cancers in the world and is the most common cause of cancer death in women aged 45–55 years (1, 2).

Human epidermal growth factor receptor 2 (HER2) belongs to a family of four transmembrane receptors involved in signal transduction pathways that regulate cell growth and differentiation. HER2 has an important role in the pathogenesis of breast cancer. Overexpression of HER2 is associated with malignancy and poor prognosis of breast cancer. One strategy for targeted cancer therapy is the use of anti-HER2 targeting agents (1, 3, 4).

Aptamers are single strand nucleic acids that fold in 3D structures (5, 6) and specifically bind to their targets such as proteins, nucleic acids, phospholipids and sugars with high affinity and selectivity (7, 8). Aptamers offer advantages over antibodies, which make them useful tools for the validation of targets. Aptamers are small and nonimmunogenic molecules that can be produced by chemical synthesis without the need for biological systems. Chemical production is cost effective and can be automated. Aptamers are stable in wide ranges of pH and temperature. They can also withstand organic solvents. The characterization and modification of aptamers is easier than antibodies (5, 9, 10).

Aptamers that bind to specific targets are generated by SELEX (the Systematic Evolution of Ligands by Exponential enrichment). During this process oligonucleotides with unique sequences are selected from a random pool.

SELEX was independently developed in 1990 by Lary Gold and Jack Szostak (11-13). In the SELEX procedure an oligonucleotide library is incubated with the target. Then oligonucleotides with affinity for the target are eluted, amplified and single stranded. SELEX cycles are repeated for several rounds until proper sequences that specifically recognize the target are obtained (14, 15).
The aim of this study was to find RNA aptamers targeting HER2-overexpressing TUBO cells using the Cell-SELEX strategy. The selected aptamers showed strong affinity to TUBO cells. Specific targeting of HER2 receptor by aptamers was further confirmed by flow cytometry.

**Materials and Methods**

**Cell lines and culture condition**

TUBO, a cloned cell line that overexpresses the rHER2/neu protein, was used as the target of selection. This cell line was kindly provided by Dr. Pier-Luigi Lollini (Department of Clinical and Biological Sciences, University of Turin, Orbassano, Italy) and was cultured in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 20% fetal bovine serum (FBS).

A murine colon carcinoma cell line, CT26, was purchased from the Pasteur Institute of Iran, and cultured in RPMI-1640 medium supplemented with 10% FBS. CT26 cells (rHER2/neu negative) were used as a negative control to remove sequences that bind to both cell lines (16, 17). Cells were regularly subcultured to maintain exponential growth.

Other cell lines were used in this project, all purchased from the Pasteur Institute of Tehran, Iran, and maintained in cell culture medium under the recommended conditions.

**Primers and library**

The random library (5’ACC GAG TCC AGA AGC TTG TAG TAC T-N35-GCC TAG ATG GAG TGG AAT TCT CCC TAT AGT GAG TCG TAT TAC-3’) was synthesized by DNA synthesizer (PolyGen) and purified by gel purification (18-20). This library was amplified using forward primer (5’-GTAATAGCCTCACATAGGGAATCTCAACTGGCCTCA-CTA-3’) and reverse primer (5’-ACCGAGTCCAGAA-GCTTGTAGT-3’). RNA library was transcribed from the PCR product using DuraScribe T7 transcription kit (Epigen Technologies). RNase A resistant product was accomplished by replacing CTP and UTP with 2’-Fluorine-dCTP (2’-F-dCTP) and 2’-Fluorine-dUTP (2’-F-dUTP) in the DuraScribe in vitro transcription reaction. After purification, the RNA library was added to 300 µl binding buffer containing HEPES-NaOH (20 mM, pH 7.4), NaCl (150 mM), CaCl2 (1.5 mM), MgCl2 (0.5 mM), and 10% yeast tRNA (Sigma). To retain correct configurations, the RNA library was denatured at 90 °C for 1 min and snap-cooled on ice.

**Cell SELEX**

TUBO cells were dislodged from the flask after a short period of incubation with trypsin and then counted. The cells’ viability was assessed by Trypan blue assay. 5-10 million cells were centrifuged, washed 3 times with washing buffer (20 mM HEPES-NaOH, pH 7.4, 150 mM NaCl, 1.5 mM CaCl2, 0.5 mM MgCl2) and resuspended in the binding buffer (washing buffer plus 10% yeast tRNA (Sigma)).

Cells were incubated with a solution of library in the binding buffer at 4 °C for 45 min. After 2 washes, sequences that bound to TUBO cells were recovered by denaturation of RNA sequences as well as surface proteins at 95 °C for 5 min. Cells were precipitated and RNA sequences were retrieved by ethanol precipitation of the supernatant. The obtained RNA sequences were reverse-transcribed using the Cloned AMV first-strand cDNA synthesis kit (Invitrogen), and PCR-amplified. The purified PCR products were transcribed in vitro using the Dura Scribe T7 transcription kit (Epigen Technologies). After third round of selection, counter selection was performed to subtract sequences with affinity for both the control and target cells. For negative selection, the RNA sequences eluted from TUBO cells were incubated with the CT26 cell line, and unbounded sequences were ethanol precipitated (19, 21, 22).

**Cloning, sequencing, and structure analysis of selected aptamers**

After 12 rounds of selection, the PCR amplified dsDNAs were cloned in to Escherichia coli DH5-α using the TOPO TA cloning kit (Invitrogen K4500-40). Individual white colonies were picked and cultured in a liquid LB (Luria-Bertani) medium. After a brief centrifugation, plasmids were purified using Gene lute TM HP Fine-Minute Plasmid Miniprep (Sigma Aldrich) and sequenced by Bioneer Company (23).

Sequences were aligned using the sequence alignment program Clustal Omega (https://www.ebi.ac.uk/Tools/msa/clustalo). A phylogenetic tree was constructed using the DNAMAN version 6 software (Lynnun Corporation) and the sequences were grouped. Representative sequences from different groups were selected as candidate aptamers for further characterization (21, 24).

**Flow cytometry binding assay**

To monitor the enrichment of the library from aptamers with affinity for TUBO cells, eluted aptamers after 1, 3, 5, 8, and 12 SELEX cycles were labeled by Cy-5. Aminohexyl-ATP was incorporated into the RNA structure during the Dura Scribe transcription reaction followed by post transcriptional labeling with Cy5-NHS ester.

Then the Cy-5 labeled RNA sequences were incubated with about 5×10⁵ target and negative cells for 30 min on ice in the dark in 500 µl binding buffer in the presence of 10% FBS. After incubation, the cells were washed two times and suspended in binding buffer and analyzed by flow cytometry (BD FACSCalibur cytometer).
The flow cytometry data was analyzed using FCS Express 4 Flow Cytometry (De Novo Software, Los Angeles, CA).

**Determination of aptamer selectivity**
The selectivity of aptamers for TUBO cells was tested by incubation of aptamers with the following cells: human prostatic carcinoma cell line (PC3), transformed mouse embryonic fibroblast cell line (NIH3T3), human breast cancer cell line (SK-BR-3), human Burkitt’s lymphoma cell line (Raji), and murine colon adenocarcinoma cell line (C26).

**Determination of secondary structure of aptamers**
The secondary structure of selected aptamers was analyzed by free-energy minimization using the algorithm according to the method of Zuker in mfold web based software (http://mfold.rna.albany.edu/?q=mfold) (25).

**Determination of apparent dissociation constant of aptamers**
The range of Cy5 labeled aptamers and control library were incubated with a constant number of TUBO cells. The mean fluorescent intensities of aptamer and library at each concentration were determined. All binding assays were performed in triplicate. The mean fluorescence intensity of the unselected library was subtracted from that of the corresponding aptamer with target cells. Then, apparent dissociation constants (Kd) for the aptamer-cell interaction were calculated by fitting the dependence of fluorescence intensity (Y) and the concentration of aptamers (X) into the one-site saturation equation $Y = B_{\text{max}} X/(K_d + X)$ using Prism version 5 (GraphPad Software, San Diego, CA). In this equation $B_{\text{max}}$ is the maximum specific binding with the same unit as Y (21).

**Effect of trypsin treatment on binding of aptamers to TUBO cells**
TUBO cells were incubated with trypsin for 8 min at 37 °C. To stop trypsin activity, cells were mixed with ice cold culture medium containing FBS. Then, the cells were quickly washed and centrifuged. After resuspension in the binding buffer, cells were incubated with aptamers. Binding of aptamer to treated cells was assessed by flow cytometry and compared with untreated TUBO cells as positive control (26, 27).

**Effect of temperature and culture medium on the binding of aptamers to cells**
All aptamers were isolated at 4 °C in binding buffer. To verify the ability of target recognition of aptamers at 37 °C, binding assay was performed at 37°C and in culture medium as described in the binding assay section (26, 28).

**Verification of aptamer binding to extracellular domain of HER2**
TUBO cells were prepared as described in the flow cytometry section. TUBO cells were incubated with anti-HER2 Affibody® molecules (Affibody AB). After two washes, cells were incubated with Cy5 labeled aptamers. TUBO cells were also incubated with Cy5 labeled aptamers as positive control.

Mean fluorescence intensity of aptamers binding to Affibody pretreated cells was determined and compared with aptamers binding in positive control.

**Results**

**SELEX and aptamer enrichment monitoring**
SELEX was used to screen aptamers specifically binding to the breast cancer cell line TUBO, with CT26 as counter selection. Following every two continuous rounds, binding assay was performed. The enrichment of aptamer pool was monitored by flow cytometry. As we progressed with SELEX, the fluorescence intensity gradually increased, while there was no significant change in the same experiment with the counter cells. Selection cycles were stopped when significant difference between the mean fluorescence intensities of the unselected library and the selected RNA pool was observed (Figure 1).

**Cloning, sequencing and structure analysis of selected aptamers**
After 12 rounds of selection the RNA pool was reverse transcribed into cDNA and amplified by PCR. The highly enriched pools were cloned into pTZ57R/T vector, and after culturing, positively inserted clones were picked, and plasmids were purified and sequenced using M13F (-20) and M13R (-40). Sequences were aligned and clustered based on their sequence homology (Figure 2). From the sequencing and alignment results TSA6, TSA7, TSA10, TSA12, and TSA14 were synthesized as potential aptamer candidates.

**Determination of aptamer selectivity**
We investigated the binding affinity of selected aptamers for transformed mouse embryonic fibroblast cell line (NIH3T3), SKBR3 (hHER2+) breast cancer cell line, human prostate cancer cell line (PC3), Burkitt’s lymphoma cell line (Raji), and C26 murine colon carcinoma cell line (Table 1).

Although some levels of recognition were observed with these cell lines, affinity of aptamer candidates to TUBO cell line was much higher. There were no sequences that recognize TUBO cells exclusively. All selected aptamers had fair affinity to SKBR3 cell lines. As shown in Table 1 interaction of aptamers and normal cell line was like control cells. Affinity of aptamers for C26 cell line was measured, because it was closely related to control cell line.
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TSA14 showed clearly stronger signal with TUBO cells compared to other sequences. Mean fluorescence intensity of TSA14 was about 2.5 times higher than other aptamers. TSA12 aptamer did not recognize CT-26, and PC3 cell lines and had minimum affinity for cell lines other than TUBO cells. According to this data, TSA14 and TSA12 were selected for further studies.

Determination of the secondary structure of aptamers

The secondary structure of RNA aptamers was predicted with the mfold web based software by calculation of minimum free energy (http://mfold.rna.albany.edu/?q=mfold). Figure 3, A and Figure 3, B show the predicted secondary structure of TSA12 and TSA14 aptamers. The free energy (ΔG) of the most stable structures of TSA12 and TSA14 was -8.29 and -9.68, respectively. Free energy was more negative for TSA14.

The predicted structure of TSA14 was more complicated, having 3 loops.

Determination of aptamer binding affinity

Dissociation constant (Kd) of aptamer-TUBO cell interaction was determined. Kd of TSA12 and TSA14 were 191.9±21.77 nM and 133.9±12.78 nM, respectively (Figure 3, C and Figure 3, D). TSA14 had higher affinity for target cells.

Table 1. Specificity determination of TSA6, TSA7, TSA10, TSA12, and TSA14 aptamers

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Disease</th>
<th>TSA6</th>
<th>TSA7</th>
<th>TSA10</th>
<th>TSA12</th>
<th>TSA14</th>
</tr>
</thead>
<tbody>
<tr>
<td>TUBO</td>
<td>mouse breast cancer</td>
<td>**</td>
<td>***</td>
<td>**</td>
<td>-</td>
<td>****</td>
</tr>
<tr>
<td>CT26</td>
<td>mouse colon carcinoma</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>-</td>
<td>*</td>
</tr>
<tr>
<td>PC3</td>
<td>human prostate cancer</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>-</td>
<td>*</td>
</tr>
<tr>
<td>NIH3T3</td>
<td>transformed mouse embryonic fibroblast cell line</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>SKBR3</td>
<td>human breast cancer</td>
<td>*</td>
<td>**</td>
<td>**</td>
<td>**</td>
<td>**</td>
</tr>
<tr>
<td>Raji</td>
<td>human Burkitt’s lymphoma</td>
<td>*</td>
<td>*</td>
<td>*</td>
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<td>*</td>
</tr>
<tr>
<td>C26</td>
<td>mouse colon carcinoma</td>
<td>*</td>
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<td>*</td>
</tr>
</tbody>
</table>

Note: The mean fluorescence intensity (MFI) of TSA14 aptamer with TUBO cells was chosen as maximum intensity (****). The MFIs with other cells and other aptamers were therefore compared and accordingly assigned approximate binding intensity. A “*” indicates that the MFI is similar to MFI of unselected library with cell line.
Figure 2. (A) Sample data of sequence alignment using Clustal Omega showing homologous families and nonhomologous sequences of potential aptamer sequences. (B) Phylogenetic tree of selected sequence.

**Effect of trypsin treatment on binding of aptamer to TUBO cells**

Figure 4, A shows that trypsin pretreatment decreased the fluorescence intensities compared to untreated cells. Trypsin treatment assay indicated that TSA14 aptamer bound to surface proteins that were digested by trypsin.

**Effect of temperature and culture medium on the binding of aptamers**

As shown in Figure 4, B and Figure 4, C the binding of aptamers changed slightly at 37°C.

Moreover, TSA12 and TSA14 showed reduced, but still significant binding to TUBO cells in culture medium (Figure 4, D and Figure 4, E).

**Verification of aptamer binding to the extracellular domain of HER2**

In order to confirm that aptamers bind to HER2 on the surface of TUBO cells, the external domain of HER2 was masked with anti-HER2 affibodies. Flow cytometry data showed that binding of Cy5-labeled aptamers decreased considerably after the incubation of TUBO cells with affibodies (Figure 5).
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Figure 3. Software simulated secondary structure of TSA12 (A) and TSA14 (B) aptamers, binding curve of TSA12 aptamer (C) and TSA14 aptamer (D) with TUBO cells. Cells were incubated with varying concentrations of Cy5-labeled aptamer and unselected library in triplicate. Mean fluorescence intensity of each concentration was determined. The mean fluorescence intensity of the unselected library (background binding) was subtracted from the mean fluorescence intensity of corresponding aptamer. Using Prism, the apparent dissociation constant (Kd) of aptamer-cell interaction was obtained by fitting the dependence of fluorescence intensity of specific binding on the concentration of aptamers to the one-site saturation equation Y= Bmax X/(Kd + X)

Discussion

We successfully performed cell SELEX to isolate RNA aptamers that bind with high affinity and selectivity to TUBO cell line (rHER2/neu +). Human epidermal growth factor receptor 2 (HER2/neu), a 185-kDa tyrosine kinase receptor, is a proto-oncogene, and activating mutations of this gene can result in constitutive tyrosine kinase activity and uncontrolled proliferation (29-31). Her2/neu is over-expressed in 20-30% of breast cancers, typically as a result of gene amplification, and is associated with aggressive disease and a poor prognosis (32). TUBO cells are a murine cell line that overexpresses rHER2/neu protein (17).

Some efforts have previously been made to find aptamer molecules that specifically bind to HER2/neu (24, 33-35). Previous studies have used human cell lines to perform cell SELEX. Dastjerdi et al employed cell-SELEX procedure to generate an enriched pool DNA aptamers by using HER2-positive cells as the target for aptamer selection (36). Kim and Jeong developed an RNA aptamer against HER2 protein, and proposed that the selected aptamer could potentially be utilized as imaging agents for HER2-positive cancers (33). Giangrande et al used “cell-internalization SELEX” to select a series of RNA aptamers against HER2 that specifically recognized and were efficiently internalized by HER2-positive breast cancer cells (37). Here, we picked TUBO cells to proceed with the SELEX process. With no or limited access to nude mice facilities, the development of a proper targeting ligand to a rodent cell line, like TUBO cells, may facilitate subsequent in vivo studies in BALB/c mice. Since rHER2/neu and human HER2/neu are highly homologous (38), the SELEX products may be used as targeting ligand in hHER2/neu overexpressing cell lines.

RNA aptamers have advantages over antibodies and DNA aptamers; they can be chemically synthesized in short time; they are small and nonimmunogenic; they are stable in harsh situations. Chemical modifications (e.g. 2’deoxy, 2’F, 2’NH3, 2’OMe) improve RNA aptamer
stability against RNase in biological fluids like blood. Because of its single stranded nature, RNA can fold into many different three-dimensional shapes, allowing for tighter and more specific binding to target molecules. Moreover, RNA aptamers have smaller size compared to DNA aptamers of the same size, and thus pass cell membranes much easier than DNA aptamers (39, 40).

In this project we performed positive selection by retrieving sequences that bind to TUBO cells. After 3 rounds of selection, counter selection was performed using CT26 cell line (rHER2/neu negative)(16, 17). As both target and control cell lines were murine cancerous cells, it was assumed that they have common surface cancerous ligands with at least one exception for HER2 molecules on TUBO cells (17).
In each round, cell suspension was prepared by short incubation, about 1 min, of cell culture with trypsin. This treatment did not have a significant effect on the binding of aptamers. We noticed that enzyme free dissociation buffers, like enzyme free dissociation solution (Millipore), increased the number of dead cells in flow cytometry assay. To generate aptamers that specifically bind to target cells with high affinity, we gradually increased the stringency of selection conditions during the SELEX process: the volume of binding buffer and washing buffer were increased and the number of target cells and incubation time was reduced. Starting from the fourth round of SELEX, 10% FBS was added to the binding buffer and gradually increased up to 20% (21). After 12 rounds of selections, cloned and sequenced aptamers were tested for their ability to bind TUBO cells and other cancerous or normal cell line. The ultimate aim of this study was to design aptamers that can bind to the extracellular domain of HER2 protein. Hence it was important that selected aptamers could recognize SKBR3 cell line. All selected aptamers had fair affinity for SKBR3 cell lines. It could be due to the high percent of homology (about 90%) between the extra cellular domains of human HER2/neu and mouse HER2/neu (41). We selected TSA12 that had the most affinity for TUBO cells, and TSA12 that had the most specificity to TUBO cells for further studies. Then the secondary structures of TSA12 and TSA14 were determined with the mfold program. The predicted structure of TSA14 was more complicated, having 3 loops. Three hairpin motif in the secondary structure of TSA14 makes it more stable compared to TSA12. Hairpins
Figure 5. Verification of aptamer binding to extracellular domain of HER2. Selected aptamers were incubated with TUBO cells (blue), and with TUBO cells treated with anti-HER2 affibodies. Black histogram represents binding of unselected library to TUBO cells serve as recognition motifs in RNA aptamer structures (42). Aptamers with hairpin structures have more affinity for their targets. TSA14 had higher affinity to target cells and had a more stable secondary structure. The obtained results corroborated the relationship between the structure and affinity of aptamers. It seems reasonable to assume that aptamers bind to cell surface proteins. This behavior has been reported in previous studies (26, 43, 44). Trypsin treatment assay is a preliminary test that has been used to study the binding of aptamers to cell surface proteins (26, 45-48). Trypsin pretreatment assay was performed to confirm that the interaction of designed aptamers with cell lines is through an interaction with cell surface proteins. The results showed decreased fluorescence intensities of trypsin pretreated cells compared to untreated cells. This means that TSA14 aptamer binds to surface proteins that are digested by trypsin.

In order to find aptamers that bind to the surface of target cells, Cell SELEX process was performed at 4 °C (21, 45). Since, it is necessary to test binding activity of aptamers at physiologic temperature.

We also assessed aptamer binding behavior in culture medium supplemented with FBS. The data showed that selected aptamers could be adopted in different conditions and still keep their binding ability to TUBO cells, which is important for in vivo studies.

As previously mentioned the purpose of this study is to produce aptamers that can recognize HER2 proteins on the surface of TUBO cells. In order to determine whether the target of the aptamers is a HER2 protein on the cell surface, anti-HER2 affibodies were incubated with TUBO cells and their binding of aptamers was analyzed by flow cytometry. The decrease in binding efficiency after incubation with affibodies strongly implied that aptamer targets on TUBO cells were most probably extracellular domain of HER2.

Conclusion

Whole cell SELEX was performed to find RNA aptamers against TUBO cell line. After 12 rounds of selection, a panel of aptamers was selected that had high affinity to target cells. Among them TSA14 aptamer had the most affinity for target cells. Different experiments showed that the interaction of TSA14 aptamer with TUBO cells is mediated by binding to the extracellular domain of HER2. Therefore, the TSA14 aptamer could be a candidate molecule for targeting the HER2+ tumors and merits further investigation.

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