RNA recognition by a human antibody against brain cytoplasmic 200 RNA

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ABSTRACT
Diverse functional RNAs participate in a wide range of cellular processes. The RNA structure is critical for function, either on its own or as a complex form with proteins and other ligands. Therefore, analysis of the RNA conformation in cells is essential for understanding their functional mechanisms. However, no appropriate methods have been established as yet. Here, we developed an efficient strategy for panning and affinity maturation of anti-RNA human monoclonal antibodies from a naïve antigen binding fragment (Fab) combinatorial phage library. Brain cytoplasmic 200 (BC200) RNA, which is also highly expressed in some tumors, was used as an RNA antigen. We identified MabBC200-A3 as the optimal binding antibody. Mutagenesis and SELEX experiments showed that the antibody recognized a domain of BC200 in a structure- and sequence-dependent manner. Various breast cancer cell lines were further examined for BC200 RNA expression using conventional hybridization and immunoanalysis with MabBC200-A3 to see whether the antibody specifically recognizes BC200 RNA among the total purified RNAs. The amounts of antibody-recognizable BC200 RNA were consistent with hybridization signals among the cell lines. Furthermore, the antibody was able to discriminate BC200 RNA from other RNAs, supporting the utility of this antibody as a specific RNA structure-recognizing probe. Intriguingly, however, when permeabilized cells were subjected to immunoanalysis instead of purified total RNA, the amount of antibody-recognizable RNA was not correlated with the cellular level of BC200 RNA, indicating that BC200 RNA exists as two distinct forms (antibody-recognizable and nonrecognizable) in breast cancer cells and that their distribution depends on the cell type. Our results clearly demonstrate that anti-RNA antibodies provide an effective novel tool for detecting and analyzing RNA conformation.

Keywords: RNA recognition; noncoding RNA; BC200 RNA; human monoclonal antibody; breast cancer cells

INTRODUCTION
Analysis of the human genome led to the surprising revelation that only 2% of the total genomic sequence comprises protein-coding regions (International Human Genome Sequencing Consortium 2004). Unexpectedly, transcription is prevalent throughout the mammalian genome, yielding complex pools of transcripts, including those with no protein-coding capacity (Carninci et al. 2005). Recent studies have identified several cellular RNAs that function as a class of gene regulators, a role previously assumed to be reserved mainly for proteins (Witusz et al. 2009; Kugel and Goodrich 2012). In many cases, the biological roles performed by RNAs in cells rely on their three-dimensional structures, although specific sequences have been shown to be essential for function (Chowdhury et al. 2006; Wan et al. 2011, 2012; Breaker 2012). However, effective tools for recognizing the conformations of structured RNA are rare. In general, RNAs are detected via hybridization with complementary nucleic acid probes. However, direct probing of structured RNAs with the classical hybridization method is difficult, since the hybridization procedure requires partial denaturation conditions to secure single-stranded regions for base-pairing between RNA and the probe, which could cause conformational changes in RNA. One possible method of effectively probing structured RNAs is the use of specific antibodies. While antibodies against specific proteins can be easily generated, it is considered impossible to produce antibodies recognizing the RNA structure through immunization due to the intrinsic instability of RNA, which leads to rapid degradation upon injection into animals. Furthermore,
nucleic acids such as RNA or DNA are not normally immunogenic, owing to recognition by immune cells as self-antigens (Pokkuluri et al. 1994), although Stollar (1980) reported that antibodies to single-stranded DNA can be induced by linking them to proteins or polypeptides, followed by injecting to animals. On the other hand, anti-RNA antibodies can be obtained through panning and affinity maturation from an antibody library because previously Piccirilli’s group reported the selection of specific antigen binding fragments (Fabs) against a domain derived from the Tetrahymena group I intron using a synthetic phage-display library (Ye et al. 2008; Koldobskaya et al. 2011).

BC200 RNA (brain cytoplasmic 200 RNA) is a small noncoding RNA (Fig. 1) that operates as a translational modulator in human cells (Cao et al. 2006). BC200 RNA is implicated in the inhibition of local synaptodendritic protein synthesis in neurons and is not detected in somatic cells other than neurons (Tiedge et al. 1993). A number of tumors (carcinomas of breast, cervix, esophagus, lung, ovary, parotid, and tongue) are reported to express BC200 RNA (Chen et al. 1997). Moreover, this noncoding RNA appears to be expressed at higher levels in invasive carcinomas than in benign tumors of the breast, suggestive of a role in tumorigenesis (Iacoangeli et al. 2004). However, the biological relevance of high BC200 RNA expression in tumor tissues is yet to be clarified.

In this study, we have developed an efficient strategy for panning and affinity maturation of human monoclonal antibodies binding to RNA from a naïve Fab combinatorial phage library, using BC200 RNA as the antigen. We identified MabBC200-A3 as the optimal binder, which interacted with BC200 RNA at a dissociation constant of ∼7 nM. Mutagenesis and SELEX experiments showed that the antibody recognizes BC200 RNA in a sequence- and structure-dependent manner. Expression of BC200 RNA in various breast cancer cell lines was further examined using conventional hybridization and immunoanalysis with MabBC200-A3.

RESULTS

Fabs selection against BC200 RNA

We used a human naïve Fab combinatorial phage library (I Park and HJ Hong, in prep.) to select Fabs recognizing BC200 RNA (Fig. 1). In principle, we adapted the procedures used for the selection of RNA-binding proteins and synthetic antibodies from the groups of Belasco (Laird-Offringa and Belasco 1996) and Piccirilli (Ye et al. 2008; Koldobskaya et al. 2011), respectively. We used a streptavidin-coated immunotube and biotin-tagged BC200 RNA for RNA immobilization, which is known to facilitate more effective RNA binding and clearer separation of buffers than streptavidin-coated beads. The specificity of clones against RNA was additionally enhanced by eluting RNA-binding phages from the immunotube with yeast total RNA as competitors. We selected two unique Fab phage clones, FabBC200-A and FabBC200-B, after four rounds of panning (Fig. 2A).

Binding of the selected clones to BC200 RNA was confirmed using dot-blot analysis, where an individual phage immobilized on a membrane as a dot was incubated with labeled BC200 RNA. FabBC200-A and FabBC200-B were used to generate soluble whole-antibody forms, MabBC200-A and MabBC200-B, respectively. Determination of their binding affinities with the filter binding assay revealed $K_d$ values of ∼36 nM for MabBC200-A and ∼76 nM for MabBC200-B (Fig. 2B,C). Interestingly, sequence analysis disclosed the same hexapeptide sequence “ARGSPR” on the early CDR3 region of the heavy-chain (CDR-H3) in both clones. Considering that the library originates from human naïve antibodies and rarely has bias toward any specific class of residues, this hexapeptide match appears significant, implying a critical role of the peptide sequence in binding to BC200 RNA.

Affinity maturation of the anti-RNA antibody

Next, we performed affinity maturation to enhance the affinity of the BC200 RNA-binding antibody, starting with the higher affinity clone, FabBC200-A. For this purpose, a novel
affinity maturation method was designed (Supplemental Fig. 1). The essential features of this method were sib selection strategy (McCormick 1987) combined with PCR mutagenesis using randomized codons of the nucleotide sequence NNS (N = A, G, C, or T; S = G or C). The previous affinity maturation study of anti-RNA antibodies screened from a synthetic library disclosed that mutations were observed in the CDR-L3 regions of higher-affinity antibodies at a relatively high frequency, rather than in other CDR regions (Koldobskaya et al. 2011). Accordingly, CDR-L3 of FabBC200-A was chosen as an initial target CDR region for affinity maturation. Nine mutant sets were generated, each containing clones with random mutations in one of nine residues in CDR-L3. The mutant set with the highest affinity against antigen was selected. Individual clones were separated from the selected mutant set and their binding affinities assayed. We simply examined a pool of 96 independent clones from the selected set, which included all 20 amino acids with a probability of >90%. The mutant set of residue S107 showed the most improved binding affinity, relative to parental FabBC200-A. Further analysis of 96 clones randomly selected from the S107 mutant pool revealed four clones with higher affinity than the parental clone. Sequence analysis further disclosed S107G, S107A, S107C, and S107V mutations in CDR-L3 (Fig. 3A). Mutant clones were designated FabBC200-A1, FabBC200-A2, FabBC200-A3, and FabBC200-A4, respectively. Their whole-antibody forms (MabBC200-As) were generated and tested for binding affinity to BC200 RNA, revealing two- to fivefold enhanced $K_d$ values of 7–18 nM, compared with the parental clone (Fig. 3).

Specificity of the antibody to BC200 RNA

The structural mapping showed that the region of BC200 RNA protected by the antibody MabBC200-A3 is localized to a structural domain spanning residues 60–110. To examine how this region is involved in the antibody binding, we generated an RNA fragment of 73 nucleotides (nt), named RNA(51-120), which contained BC200 RNA sequences of residues +51 to +120 and two extra nucleotides at the 5' end.
end and one extra nucleotide at the 3′ end. The binding assay revealed that RNA(51-120) bound to MabBC200-A3 with a similar affinity to that for BC200 RNA (Fig. 5), suggesting that this truncated version is sufficient for binding to the antibody. To examine which bases in the protected region are important for the RNA–antibody interaction, we performed a SELEX (systematic evolution of ligands by exponential enrichment) (Supplemental Fig. 3; Table 1). First, we chose 11 bases residing on loops in the secondary structure model among the total 19 protected bases. Then we substituted randomized base N in place of each chosen base for generation of a partially randomized RNA pool derived from RNA(51-120). This RNA pool of $4^{11} \times 4.2 \times 10^6$ diversity was subjected to SELEX. RNA species binding to the antibody were selected by immunoprecipitation. After eight rounds of selection, 50 independent clones were isolated and sequenced (Fig. 6; Table 1). Surprisingly, 46 clones had the exactly same sequence of RNA(51-120). The other four clones differed in sequence one another: They had five to 10 mismatches among the 11 bases as compared with RNA(51-120), and they all showed their $K_d$ values of $>1$ mM for MabBC200-A3. Since their lack of binding was comparable to that of BC200 RNA to the control antibody, we concluded that they had no binding affinity to MabBC200-A3 and that RNA(51-120) is the only one binder out of the partially randomized RNA pool.

Therefore, the SELEX results suggest that all the 11 bases are important for determining specificity of the antibody. Next we tested whether recognition by the antibody would require a specific structure. We performed mutagenesis on RNA(51-120) to disrupt a stem formed by base-pairing between residues +71 to +77 and residues +91 to +97 (Fig. 5), which is essential for RNA secondary structure formation. To do this, we substituted 3 nt of +73 to +75 (CCA to GGU) in the middle 3 bp within the stem because the 3 bp does not belong to bases protected by the antibody. While the resulting mutant RNA, RNA(51-120)-M1, did not bind to the antibody, the binding ability of RNA(51-120)-M12 generated by a compensatory mutation in the other strand was restored to the same level as that of RNA(51-120) (Fig. 5), suggesting that the RNA structure is also crucial for the antibody–RNA interaction.

RNA discrimination by the anti-RNA antibody

Experiments to ascertain whether MabBC200-A3 effectively discriminates BC200 RNA from other RNAs were performed.

![FIGURE 4](image-url)

**FIGURE 4.** Antibody binding regions of BC200 RNA. (A) Hydroxyl radical footprints of BC200 RNA with 0, 5, 10, and 50 nM MabBC200-A3. Protected regions are denoted with black bars (a and b). The regions were detected from the range of a 10 nM antibody, consistent with a $K_d$ of $\sim7$ nM, as observed using the nitrocellulose filter binding assay. Untreated and RNase T1 ladders of denatured BC200 RNA are shown in lanes C and T1, respectively. (B) Binding regions in the secondary structure of BC200 RNA are highlighted in red letters. (C) Predicted 3D structure of a region of BC200 RNA (residues 51–119) carrying the anti-RNA antibody recognition motif. The binding regions are indicated in red.

![FIGURE 5](image-url)

**FIGURE 5.** Binding affinity of truncated derivatives of BC200 RNA. (A) Possible secondary structures of RNA(51-120) and its derivatives (Zuker 2003), and the $K_d$ values of the complexes between RNAs and the MabBC200-A3 antibody. RNA(51-120) of 73 nt contains BC200 RNA sequences of residues +51 to +120, two extra nucleotides at the 5′ end, and one extra nucleotide at the 3′ end. Protected regions by the MabBC200-A3 antibody are highlighted in red letters. The mutated sequences in RNA(51-120)-M1 and RNA(51-120)-M12 are boxed. (B) Binding assays for determination of $K_d$ values of the antibody–RNA complexes were carried out as for Figure 2B.
using total cellular RNA pools from several breast cancer cell lines. First, BC200 RNA expression in various breast cancer cell lines was examined with Northern blot analysis (Fig. 7A). We observed significant differences in expression levels among the cell lines. BC200 RNA expression increased in the order: MDA-MB-231 = SKBR3 < T47D < Hs578T < MDA-MB-435 < MCF7. MDA-MB-231 and SKBR3 cells showed little expression, similar to the normal breast cell line, MCF10A. Next, the immunoprecipitation assay was performed with total cellular RNAs prepared from these cells (Fig. 7B). Notably, BC200 RNA was immunoprecipitated from BC200 RNA-expressing cells with MabBC200-A3, but not the control antibody. 7SL RNA, a component of the signal recognition particle, is a predecessor of Alu, the most abundant transposable elements in the human genome (Walter and Blobel 1982; Ullu and Tschudi 1984). BC200 RNA is composed of a 5′ Alu domain, which shares high sequence similarity with that of 7SL RNA (Bovia and Strub 1996). While the region of BC200 RNA protected by the MabBC200-A3 antibody was within the Alu domain, 7SL RNA was not precipitated from total RNAs of cells with the antibody, suggesting that MabBC200-A3 specifically recognizes BC200 RNA.

Next, to determine whether MabBC200-A3 recognizes BC200 RNA present within the cell, flow cytometric analysis was performed using MabBC200-A3 as primary antibody (Fig. 8A). For this purpose, breast tumor cells were suspended following fixation in formaldehyde and permeabilization with Tween-20 to allow antibody penetration into the cells. The fluorescence intensities determined using flow cytometry were inconsistent

| Table 1. Sequences of selected RNAs after eight rounds of selection |
|---------------------------------------------------------------|
| RNA (5′→3′) | Frequency | Sequence |
|-------------|-----------|----------|
| RNA(51-120) | -         | GGAGGCCGGGAGGAGAUAUGUGAGCCCAAGAGAUUCGAGGACC |
| Randomized RNA pool | -         | GGAGGCCGGGAGGAGAUAUGUGAGCCCAAGAGNNNNNACC |
| S1 | 46 | GGAGGGCGGGGAGGAGAUAUGUGAGCCCAAGAGJUCGAGGACC |
| S7 | 1 | GGAGGGCGGGGAGGAGAUAUGUGAGCCCAAGAGJUCGAGGACC |
| S14 | 1 | GGAGGGCGGGGAGGAGAUAUGUGAGCCCAAGAGJUCGAGGACC |
| S40 | 1 | GGAGGGCGGGGAGGAGAUAUGUGAGCCCAAGAGJUCGAGGACC |
| S50 | 1 | GGAGGGCGGGGAGGAGAUAUGUGAGCCCAAGAGJUCGAGGACC |

After eight rounds of SELEX, selected RNAs were subcloned and sequenced. Protected regions by the MabBC200-A3 antibody are highlighted in bold. The sequences of the randomized regions are boxed, and the bases that deviated from the BC200 RNA sequences are indicated by shaded letters.
with Northern signals of BC200 RNA in the cells (Fig. 8B). Northern signals representing the cellular content of BC200 RNA increased in the order: T47D < Hs578T < MDA-MB-435 < MCF7, while the order of fluorescence signals was MDA-MB-435 < T47D < MCF7 < Hs578T. To confirm that the flow cytometry signals result from BC200 RNA–antibody complexes, the complexes were immunoprecipitated directly from cell lysates and the BC200 RNA content analyzed using Northern blotting (Fig. 9A) and RT-PCR (Supplemental Fig. S4). Comparative data revealed that BC200 RNA levels immunoprecipitating from different breast cancer cells are consistent with flow cytometry signals (Fig. 9B). These results suggest that BC200 RNA exists as two forms in the cell: one that is recognized by MabBC200-A3 and another that is not recognized by the antibody. Interestingly, while the two estrogen receptor (ER)–negative cell lines, MDA-MB-435 and Hs578T (Neve et al. 2006), expressed comparable amounts of BC200 RNA, the amounts of BC200 RNA recognized by MabBC200-A3 were distinct between the two cell lines. The majority of antigen was not recognized by MabBC200-A3 in MDA-MB-435, while Hs578T cells contained the highest amount of antibody-binding BC200 RNA.

FIGURE 7. RNA recognition by MabBC200-A3. (A) Total cellular RNAs purified from breast cancer cell lines (MCF7, SKBR3, MDA-MB-231, MDA-MB-435, Hs578T, and T47D) and a normal breast cell line, MCF10A, were analyzed for BC200 RNA content via Northern blot. The membranes were hybridized with the BC200 RNA probe (left) or 7SL RNA probe (right). (B) Total cellular RNAs were immunoprecipitated with MabBC200-A3. The BC200 RNA (left) or 7SL RNA (right) content in immunoprecipitates was analyzed using Northern blot, with MabN as a negative control antibody.

DISCUSSION

In the present study, we developed an efficient strategy for panning and affinity maturation of human monoclonal antibodies binding to RNA from a naïve Fab combinatorial phage library. Especially, sib selection combined with PCR mutagenesis using randomized NNS codons was successfully utilized for affinity maturation. We expect that our affinity maturation method should be faster than the existing protocols using libraries that are newly constructed through mutagenesis or chain shuffling across antibodies (Gram et al. 1992; Schier et al. 1996). While the diversity of the clones examined with our maturation method is relatively low compared with that obtained with the new library construction methods using mutagenesis or shuffling, the selected set has a greater probability of containing clones having higher affinity.

We identified antibodies that recognize structural motifs of BC200 RNA, implying that naïve B cells have clones capable of RNA recognition. Footprinting experiments performed with the best binder, MabBC200-A3, revealed that the antibody exclusively binds to a specific region of BC200 RNA. The SELEX and mutagenesis data further demonstrated that MabBC200-A3 specifically recognizes BC200 RNA by binding this domain in a sequence- and structure-dependent manner. When MabBC200-A3 was tested with total RNAs purified from various breast cancer cell lines, immunoanalytical signals for BC200 RNA coincided with conventional hybridization signals. Furthermore, the antibody was able to discriminate BC200 RNA from other RNAs in the purified total RNA, supporting its usefulness as a specific RNA structure-recognizing probe for the RNA. Intriguingly, however, when permeabilized cells were used instead of purified total cellular RNA,
the amounts of antibody-recognizable BC200 RNA were different, indicating that BC200 RNA exists as two distinct forms (antibody-recognizable and nonrecognizable) in the cell and that their distribution depends on the cell type. For BC200 RNA recognition by the antibody, its structure should maintain the integrity of the antigenic motif, and antibody binding to this motif should not be hindered by other BC200 RNA-binding proteins (Fig. 10). The functional differences between BC200 RNA recognized by the antibody and that lacking antibody-binding affinity remain to be established.

RNA function depends on both its conformation and sequence. Furthermore, RNA is a molecule displaying complexity and dynamic behavior that exists in various conformations and as different ribonucleoproteins in cells. Therefore, detecting the presence or absence of free RNA motifs in cells is essential for understanding the cellular functions of RNA. In this respect, RNA structure-recognizing antibodies can potentially overcome the limitations of complementarity-based RNA detection through hybridization, which prevents retention of structure. In addition, all existing methods of antibody technology can be applied for enhancing utilization of RNA structure-recognizing antibodies in basic and applied research. For instance, given that RNA usually folds into extensive structures, an anti-RNA antibody may provide an innovative tool for dissecting RNA motifs with specific conformations. The specific motifs recognized by the anti-RNA antibody may act as potent biomarkers for diagnosis. Moreover, antibody delivery systems into living cells are currently under development (Kuo et al. 1998; Fang et al. 2007), and cells expressing disease-related RNAs may thus be targeted by anti-RNA antibodies more effectively than antisense nucleic acids.

**MATERIALS AND METHODS**

**In vitro preparation of RNA**

A DNA template for BC200 RNA was obtained from human genomic DNA (Roche) via PCR using a primer pair of BC200-T7-up (5'-GAATTCATACTAACAAGCTC/TATA GGGAGGGCGGGGTTG)/BC200-dn (5'-CCC AAGCTTTTAAATGGGCGGGGTG TTG) and cloned into the EcoRI and HindIII sites of pUC19 (New England Biolabs). The resulting plasmid DNA was cleaved with Dral and used as a template for in vitro transcription with T7 RNA polymerase (Promega) to generate in vitro transcripts having the same 5' and 3' ends as the natural BC200 RNA. Truncated BC200 RNA derivatives, RNA(51-120), RNA(51-120)-M1, and RNA(51-120)-M12 were generated similarly. Briefly, the DNA template for RNA(51-120) was amplified via PCR using a primer pair,

RNA(51-120)-T7-up (5'-CCGGAATTCTAATACGACTCACTATA GGGAGGGCGGGGTTG)/RNA(51-120)-dn (5'-TCCCCCGG

![FIGURE 8. Flow cytometric analysis. (A) Fixed and permeabilized cells were treated with MabBC200-A3, followed by FITC-conjugated anti-human IgG, and subjected to flow cytometric analysis. Plots for cells treated with MabBC200-A3 and MabN are indicated by shaded and white areas, respectively. (B) Comparison between the flow cytometric signals of A and the Northern blot data of Figure 7A. Levels were normalized with MabN and 5S for flow cytometric and Northern blot analyses, respectively. The relative signals are presented in arbitrary units for each cell.](image)

![FIGURE 9. Immunoprecipitation of BC200 RNA. (A) Lysates from cells treated with MabBC200-A3 were subjected to immunoprecipitation. The BC200 RNA content in precipitates was analyzed via Northern blot using the BC200 RNA probe. It was confirmed via Western blot with lysates of antibody-treated cells that all the cell lines were permeable to antibodies to similar extents. After electrophoresis on a 5% SDS-PAGE gel, the lysates were blotted with MabBC200-A3 or anti-tubulin antibody (α-tubulin). NB and WB indicate Northern blot and Western blot, respectively. (B) Comparison between the NB data after cellular immunoprecipitation (Cellular IP) of A and the flow cytometric signals of Figure 8A.](image)
GAGAACGGGGTGTCGC). The DNA template for RNA(51-120)-M1 was amplified using a primer pair, RNA(51-120)-M1-up (5’-GGATACGCTTGACCGTGAGTTGCTGAGACCTGCC) and RNA(51-120)-M1-dn (5’-GGCAAGTCTCGAACCTCACGCTCAAGCATTCC). The DNA template for RNA(51-120)-M12 was amplified via recombinant PCR using two primer pairs, RNA(51-120)-M1-up/RNA(51-120)-M1-dn and RNA(51-120)-M12-PCR2-up (5’-GGAGTTCCGAGACCTGCCGACCGGAAATAGGGAAGGAGACGACCCAAGG)/RNA(51-120)-dn. The amplified DNA fragments were cloned into the EcoRI and Smal sites of pUC19. The resulting plasmid DNAs were cleaved with Smal and used as templates for in vitro transcription with T7 RNA polymerase.

The DNA template for an RNA(51-120)-derived library for SELEX was obtained via recombinant PCR using two primer pairs, RNA(51-120)-T7-up/RNA(51-120)-SELEX-dn (5’-GCCAGGCAAGAACCCACCGCTCAAGCTATCCTCGCC/TTATAGTGGCTGATTAATGCGG) and RNA(51-120)-SELEX-up (5’-GATAATTCTGACCGGAGGNNNNNNACCTGCCGTCGCGGCG/TAATACGACTCACTATAGG) and RNA(51-120)-SELEX-dn and amplified via PCR using a primer pair, RNA(51-120)-T7-up and RNA(51-120)-dn. The amplified DNA was cleaved with Smal and subjected to in vitro transcription with T7 RNA polymerase to generate the RNA(51-120)-derived library. After selection, cDNAs were generated via RT-PCR using a primer pair, RNA(51-120)-T7-up and RNA(51-120)-dn, and were again cleaved with Smal and subjected to in vitro transcription for additional rounds of selection.

All the in vitro transcripts were purified via gel elution. Before use, RNA was renatured by heating for 5 min at 65°C, chilling on ice for 10 min, and slowly returning to room temperature.

To prepare a biotinylated BC200 RNA for panning experiments, a DNA fragment was amplified by PCR with a primer pair of BC200-T7-up (5’-CTCCGGATCCCGGATCCCGGGCTTCTTCGACGCAACAGCTTAC) and BC200B-dn (5’-CTTCGGATCCCGGATCCCGGGCTTCTTCGACGCAACAGCTTAC) and cloned into pUC19. The resulting plasmid DNA was cleaved with Stul and used as the DNA template for in vitro transcription. The in vitro transcript and a 5’ biotinylated oligonucleotide (5’-AGGATCCGACCGGATCCCGGGCTTCTTCGACGCAACAGCTTAC) were mixed in the ratio of 1:2.5 and incubated for 5 min at 37°C, chilled on ice for 10 min, and slowly returned to room temperature to anneal through the extra adaptor sequence of RNA at the 3’ end as well as for RNA renaturation.

Selection of human Fab

For selection of Fab phage clones via panning, the phagemid vector, Fab1, was used to construct phage display libraries with a repertoire of human V1 or VHu. The resulting Fab libraries had 5.0 × 10^8 diversity. The libraries were panned for four rounds for biotinylated BC200 RNA, which was immobilized on a streptavidin-coated immunotube at room temperature. Briefly, phages were preincubated three times with the immunotube and recovered phages were amplified with VCS M13 helper phage at 20-fold MOI. Phages were purified through PEG precipitation, and the affinity for BC200 RNA was measured with dot-blot assays. Briefly, each clone phage was transferred to nitrocellulose membrane using MiniFold dot-blot apparatus (Whatman) with β^2P-labeled BC200 RNA as a probe. The membrane was exposed to imaging plate BAS-IP (Fuji) and analyzed on a phosphor-image analyzer, FLA-7000 (Fuji). Positive clones were sequenced.

Expression and purification of whole antibody forms

V_{H} and V_{L} of selected Fabs were fused with Ig heavy- and light-chain leader sequences (Ryu et al. 1996) using PCR, respectively. Amplified products were digested with EcoRI and ApaI for heavy-chain and HindIII and BsrXI for light-chain and subcloned into the corresponding sites of pdCMV-dhfrC-A10A3 (Lee et al. 2012). The resulting expression plasmid was transfected into human embryonic kidney (HEK) 293T cells using Lipofectamine 2000 (Invitrogen), according to the manufacturer’s instructions. Transfected cells were cultured in protein-free CD293 medium (Invitrogen), and culture supernatants were subjected to affinity chromatography on a Protein G-Agarose (Invitrogen) column. The purity of the antibody was determined using SDS-PAGE, and concentrations were measured using protein assay dye reagent (Bio-Rad), according to the manufacturer’s instructions.

Filter binding assays

RNA was renatured by heating for 5 min at 65°C, chilling on ice for 10 min, and slowly returning to room temperature. RNA–antibody complexes were formed in 100 μL PBS containing 0.6 nM RNA,
2 nM–100 nM antibodies, and 2 units/µL RNase inhibitor (Enzy-nomics) for 30 min at room temperature. The reaction mixture was applied to a dot-blot apparatus in which Hybond ECL membrane (GE Healthcare) and Hybond-XL membrane (GE Healthcare) were layered with Whatman 3 MM paper in between. In this process, RNAs that passed through the Hybond ECL membrane without binding to the antibody were immobilized on the Hybond-XL membrane. The Hybond-XL membrane was washed twice with PBS and hybridized with 5'-32P-labeled antisense oligonucleotide probe (5'-TTTGAGGGAGGTTACGCTTAT for BC200 RNA, 5'-GGGAGACGGCGGCTCAGCGG for truncated BC200 RNA derivatives, or 5'-GGGAATCCAGATGCCGACC for Escherichia coli 6S RNA as an irrelevant RNA), exposed to the imaging plate, and analyzed on the Phosphor-image analyzer. The amount of radiolabeled RNA on each filter was quantified with ImageJ software (NIH), and plots of bound fractions of RNA versus concentration of antibody were obtained with Origin software (OriginLab). Binding constants (K_d) were calculated by fitting data to the Hill equation.

### Affinity maturation

To enhance affinity for RNA, we implemented sib selection combined with PCR mutagenesis using randomized codons. Each of the nine residues of CDR-L3 of FabBC200-A was randomly mutated via PCR with primers containing the NNS codon. PCR products were subcloned into the BstXI site of Fab1. After the transformation of E. coli TG1 cells, each mutant set was transformed into phages. The resulting mutant phage sets were analyzed for affinity using dot-blot assays. A set with mutation at the third residue of CDR-L3 showed the highest affinity for RNA. Cells of the selected mutant set were grown in 2× YTA plates, and 96 individual clones were randomly obtained. The following equation shows that the 96 clones cover all 20 amino acids with a probability of >90%:

\[
P = \frac{1}{X^N} \times \left( X^C - 1 \times (x - 1)^N - x^C - 2 \times (x - 2)^N + \cdots \right),
\]

where P is probability for a pool containing all independent clones, X represents independent clones to be covered, and N is the number of clones to be tested. Four clones displaying higher affinity than the original clone were subjected to sequence analysis as well as filter binding assays to determine dissociation constants (K_d).

### RNA footprinting

Hydroxyl radical footprinting analysis was conducted using the Fe/EDTA-mediated Fenton reaction. Experiments were performed as described previously (Koldobskaya et al. 2011). Briefly, BC200 RNAs were 3’ end-labeled with [32P]dCTP using T4 RNA ligase (New England Biolabs). Labeled RNA molecules were renatured by heating for 5 min at 65°C, chilling on ice for 10 min, and slowly returning to room temperature. Next, 1 µL of 5 mg/mL heparin, 1 µg yeast tRNA, and the corresponding antibody were added to labeled RNA, yielding a total reaction volume of 7 µL. This binding reaction was carried out in 1× PBS for 30 min at room temperature. Subsequently, 1 µL of 1 mM Fe(II)(NH4)2(SO4)2, 1 µL of 1.25 mM EDTA, and 1 µL of 60 mM sodium ascorbate were added, and the reactions were incubated for 30 min at room temperature. Reactions were terminated by adding 1 µL of 100 mM thiourea and 11 µL gel loading buffer II (Ambion). Samples were denatured for 5 min at 95°C and loaded on a 5% (v/v) polyacrylamide–9 M urea sequencing gel. RNase T1 ladders were obtained according to the manufacturer’s instructions (Ambion).

### SELEX

The RNA(51-120)-derived library of 4^{11} (4.2 × 10^6) diversity was used for SELEX. In the first round of selection, RNA (0.2 ng, ~5 × 10^5 molecules) in 50 µL PBS was preincubated with 10 µL Protein G–agarose for 30 min to remove false-positive binders, followed by mixing with 2.8 nM MabBC200-A3 in 100 µL PBS and incubated for 30 min at room temperature. The RNA–antibody complex was immunoprecipitated with Protein G–agarose. The immunoprecipitates were washed five times with PBS containing 0.05% Tween-20 and were phenol-extracted. The RNA was recovered from ethanol precipitation and amplified through RT-PCR. The amplified cDNA was used to generate the RNA pool for additional rounds of selection. After the eighth round of selection, the amplified cDNA was cloned into a T-blunt vector (SeloGent). Individual cDNA clones were identified by DNA sequencing.

### Northern blot analysis

Total cellular RNA was extracted from cells with the easy-blue kit (Intron), according to the manufacturer’s instructions. For Northern blot analysis, RNA samples were electrophoresed on a 6% polyacrylamide gel containing 7 M urea and electrotransferred to Hybond-XL membrane (GE Healthcare). Hybridization was carried out with the 5’-32P-labeled BC200 RNA antisense oligonucleotide probe, as previously described (Lee et al. 2013). 7SL RNA and 5S RNA were probed with 5’-32P-labeled 7SL (5’-GAGGTCCACCATATTGATCGGAACCTTAGTG) and 5S (5’-CATCCAGTACTAACCAGGCC) antisense oligonucleotide. The membrane was analyzed as described in the filter binding section.

### Immunoprecipitation

After washing in ice-cold PBS, collected cells were resuspended in PBS containing 10% fetal bovine serum (Invitrogen). Cells were fixed in 0.1% formaldehyde for 15 min at room temperature, permeabilized in PBS containing 0.1% Tween-20, and subsequently incubated with MabBC200-A3 for 30 min at room temperature. After every step, cells were washed twice with ice-cold PBS. Collected cells were resuspended in ice-cold RIPA lysis buffer (150 mM NaCl, 1% sodium deoxycholate, 0.1% SDS, 1% Triton X-100, 2 mM EDTA, and 50 mM Tris-HCl at pH 7.5), incubated on ice for 30 min, and centrifuged for 20 min at 16,000 g. The supernatant fractions were used as cell lysates for immunoprecipitation with Protein G–agarose. Immunoprecipitates were analyzed via Northern blot for BC200 RNA and 7SL RNA content. Whole-cell lysates used as input were prepared using the same procedure, except for antibody treatment. To measure the levels of input antibodies, cell lysates were prepared without the final washing procedure and subjected to Western blot analysis. Where necessary, purified total cellular RNAs were subjected to immunoprecipitation. Briefly, purified total RNA was renatured, as described with in vitro transcripts above, and
incubated with MahBC200-A3 in 50 μL PBS for 30 min and immunoprecipitated with Protein G–Agarose. The immunoprecipitates were phenol-extracted, and the BC200 RNA content in the precipitates was analyzed using Northern blot.

Flow cytometry

Procedures for cell preparations and primary antibody treatments were the same as those for immunoprecipitation. Cells were subsequently treated with fluorescein isothiocyanate (FITC)–conjugated anti-human IgG (Bethyl) for 30 min at room temperature in the dark. Collected cells were filtered with Strainer capped tubes (BD) and analyzed using the LSRII flow cytometry system (BD). Plots were obtained using Flowing software (Cell Imaging Core).

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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