Abstract. It is significant to develop a probe with sensitivity and specificity for the detection of cancer cells. The present study aimed to develop an ‘activatable’ aptamer-based fluorescence probe (AAFP) to detect cancer cells and frozen cancer tissue. This AAFP consisted of two fragments: aptamer TLS11a that targets HepG2 cells, and two short extending complementary DNA sequences with a 5'- and 3'-terminus that make the aptamer in hairpin structure a capable quencher to fluorophore. The ability of the AAFP to bind specifically to cancer cells was assessed using flow cytometry, fluorescence spectroscopy and fluorescence microscopy. Its ability to bind to frozen cancer tissue was assessed using fluorescence microscopy. As a result, in the absence of cancer cells, AAFP showed minimal fluorescence, reflecting auto-quenching. In the presence of cancer cells, however, AAFP showed a strong fluorescent signal. Therefore, this AAFP may be a promising tool for sensitive and specific detection of cancer.

Introduction

The growing incidence of cancer and associated mortality worldwide highlights the importance of timely and effective diagnostic methods (1). Fluorescent probes that specifically recognize molecular targets have proven useful in many areas of biology and medicine, and they show potential for sensitive cancer diagnosis (2-4). Such probes are usually created by conjugating a reporter fluorophore to an affinity reagent, such as a monoclonal antibody, peptide or aptamer, which binds specifically to the target of interest (5-8). To ensure specificity and sensitivity, the fluorophore must emit a strong signal and be photostable. It may emit minimal fluorescence in the absence of a target, however, in the presence of one the affinity reagent may bind to the target with high specificity and affinity (9). A key challenge in designing fluorescence probes is minimizing background signals in the absence of a target.

Nucleic acid aptamers, which are short, single-stranded RNA or DNA oligonucleotides, show substantial promise as affinity agents (10-13). They have several advantages over other affinity reagents, including high affinity and specificity, facile synthesis and modification and a structure-controlled design. Compared to other affinity reagents, aptamers may be taken up more quickly by tissue, penetrate more deeply into tissue, accumulate more in target tissue and persist less in non-target organs and circulation (14,15). These characteristics make aptamers well suited for cancer detection.

In the present study, we described the development of an activatable aptamer-based fluorescence probe (AAFP) to detect cancer cells and frozen cancer tissue (Fig. 1). This probe was ‘activatable’ since it was designed to emit minimal signals in the absence of a target, which may lead to decreased background interference than ‘always on’ fluorescence probes (16). Our AAFP was a short, single-stranded DNA oligonucleotide: we used the DNA aptamer TLS11a, which binds with high affinity to cancer cells (17), and added one short DNA sequence (C-strand) to each end. The two C-strands were complementary to each other, and the 5' C-strand was conjugated to the fluorophore FAM, while the 3' C-strand was conjugated to the quencher Eclipse. In the absence of a target, the two C-strands were complementary to each other, and the 5' C-strand was conjugated to the fluorophore FAM, while the 3' C-strand was conjugated to the quencher Eclipse. In the absence of a target, the two C-strands were complementary to each other, and the 5' C-strand was conjugated to the fluorophore FAM, while the 3' C-strand was conjugated to the quencher Eclipse. Upon target binding, the two C-strands separated, strongly increasing FAM fluorescence signals.

Materials and methods

Materials. All DNA oligonucleotides were synthesized by Shanghai Sangon Biological Engineering Technology & Services (Shanghai, China). These oligonucleotides included...
Cells and animals. Cell lines were obtained from the National Center for International Research of Biological Targeting Diagnosis and Therapy of Guangxi Medical University. Human hepatocellular carcinoma HepG2, and human normal liver L02 cells were cultured at 37°C in Dulbecco's modified Eagle's medium (DMEM) supplemented with 10% fetal bovine serum (FBS; HyClone, Logan, UT, USA), and 100 U/ml penicillin-streptomycin in a 5% CO2 atmosphere. Six-week-old female BALB/c nude mice from the Guangxi Laboratory Animal Center (Guangxi, China) were raised in sterile conditions in a laminar flow hood. All experiments were carried out according to the guidelines of the Federation of European Laboratory Animal Science Associations, and all protocols were approved by the Institutional Animal Care and Use Committee of Guangxi Medical University.

Flow cytometry. Cells were cultured at a density of 5.0x10^5 cells/ml, collected by centrifugation and resuspended in 0.5 ml of phosphate-buffered saline (PBS). In order to investigate the ability of AAFP to bind to cancer cells, cells were incubated with FAM-TLS11a, AAFP or AAFP-mis (250 nM) at 4°C in the dark for 30 min. Binding reactions were analyzed by fluorescence spectroscopy as described above.

Immunofluorescence imaging of cells. Cells were seeded into 6-well plates and cultured for 24 h. Cells were fixed in 4% paraformaldehyde (Sigma-Aldrich, St. Louis, MO, USA) for 20 min, washed with PBS, and then incubated with FAM-TLS11a, AAFP or AAFP-mis in binding buffer at 4°C in the dark for 30 min. Cells were then washed again with PBS and stained with 4’,6-diamidino-2-phenylindole dihydrochloride (DAPI; Life Technologies, Foster City, CA, USA) for 5 min in the dark. Finally, cells were washed and examined by fluorescence microscopy (Nikon DS-Ri1; Nikon, Tokyo, Japan).

Cancer model and immunofluorescence imaging of frozen cancer sections. BALB/c nude mice received a subcutaneous injection of 5x10^6 HepG2 cells on their backside. Tumors were allowed to grow for 15-20 days until reaching a diameter of 0.5-1.5 cm. Mice were sacrificed, the cancer tissue was excised, and frozen sections 6-8 mm of thickness were prepared immediately. As a control, normal liver tissue was obtained from the mice that did not receive HepG2 injections.

Frozen sections were fixed with 4% paraformaldehyde (Sigma-Aldrich) for 10 min, washed with PBS, stained with DAPI for 5 min, washed again, and were then incubated with FAM-TLS11a, AAFP or AAFP-mis (250 nM) at 4°C in the dark for 30 min. Finally, the sections were washed again with PBS and observed using fluorescence microscopy.

Statistical analyses. Each experiment was carried out in triplicate. Data were expressed as the mean ± SD or as the median (range). All statistical analyses were performed using GraphPad Prism 6.02 (GraphPad Software, San Diego, CA, USA). The threshold of significance in all analyses was P<0.05.

Results

Activation of AAFP fluorescence by target cancer cells. Flow cytometry was used to investigate the ability of target cancer cells to activate AAFP fluorescence. Both FAM-TLS11a and AAFP bound to HepG2 cells, but not to L02 cells, while AAFP-mis did not bind to either cell line (Fig. 2A and B). FAM-TLS11a and AAFP showed similar rates of binding to HepG2 cells, which were significantly higher than the rate of AAFP-mis binding.

Fluorescence spectroscopy was used to analyze the increase in FAM signals due to AAFP binding to HepG2 cells (Fig. 2C and D). Negligible fluorescence was detected in the cell suspensions without the probe or in a solution of AAFP (250 nM) without cells. Incubating HepG2 cells with AAFP led to a strong fluorescence signal at 518 nm, which was much weaker when L02 cells were incubated with AAFP. These results indicate that AAFP fluorescence increased specifically in response to HepG2 cancer cells.

Optimization of C-strand length on AAFP. We tested various lengths of the C-strand on AAFP, from 3 to 5 bases, in an effort to minimize background fluorescence while maintaining strong affinity for HepG2 cells. The highest signal-to-background ratio was obtained when the two C-strands contained...
4 bases, corresponding to 4 base pairs in the hairpin structure that forms in the absence of a target (Fig. 3A). This ratio gradually decreased as the number of bases was increased above 4. We speculated that C-strands with fewer than 4 bases are incapable of keeping the fluorophore and quencher close together, whereas C-strands longer than 4 bases decrease the affinity of the aptamer for target cells.

**Optimization of the detection conditions.** We investigated the optimal conditions for AAFP binding to HepG2 cells. Cells
were incubated with AAFP at 250 nM in the dark for 30 min at 4, 25 or 37°C, and then activation of AAFP fluorescence was assessed using fluorescence spectroscopy (Fig. 3B). As the temperature increased, fluorescence activation decreased, although it remained well above the background at all temperatures tested. These results indicate that although higher temperatures, including physiological temperatures, slightly decrease the binding of AAFP to target tissues in vitro, the fluorescence signal remains strong relative to the background.

Next, we incubated HepG2 cells with varying AAFP concentrations up to 375 nM at 4°C for 30 min, and examined the fluorescence signal (Fig. 3C). The signal increased progressively up to an AAFP concentration of 250 nM, after which it almost plateaued, suggesting saturation.

Finally, we incubated HepG2 cells with AAFP at 250 nM for different lengths of time at 4°C (Fig. 3D). The fluorescence signal increased gradually from 0 to 30 min, after which it plateaued, suggesting complete binding.

**Cancer cell detection by AAFP.** To assess the sensitivity of AAFP in the detection of HepG2 cells, we incubated the probe (250 nM) with HepG2 cells at concentrations of up to 10^6 cells/ml in 0.5 ml of binding buffer, and then assessed the fluorescence signal spectroscopically. The signal increased over the entire
range of cell concentrations assessed (Fig. 4). Regression analysis of the plot of the signal intensity and logarithm of the cell concentration (fluorescence = 214.4 x log C - 6.981) revealed that under these experimental conditions, the probe was capable of detecting cancer cells at concentrations as low as ~100 cells/ml.

Using AAFP for fluorescence imaging of cancer cells. To visualize directly the specificity of AAFP binding at the cellular level, and thereby complement the flow cytometry experiments as described in Materials and methods, we incubated HepG2 and L02 cells with FAM-TLS11a, AAFP or AAFP-mis and analyzed them using fluorescence microscopy. Consistent with the flow cytometry experiments, AAFP and FAM-TLS11a were observed to bind to HepG2 cells, but not to L02 cells, while AAFP-mis did not bind to either cell type (Fig. 5).

Using AAFP for fluorescence imaging of frozen cancer tissue. We examined whether the specific AAFP binding observed with HepG2 cells in suspension may also apply to frozen sections of solid HepG2 tumors from a xenograft mouse model. As a control, we also examined AAFP binding to frozen sections of normal liver tissue from the same type of mice. Both AAFP and FAM-TLS11a were observed to bind to sections of cancerous tissue, but not to sections of normal tissue (Fig. 6). AAFP-mis did not bind to either type of tissue.

Discussion

We designed an ‘activatable’ aptamer-based fluorescence probe (AAFP) that sensitively and specifically detects cancer cells. The probe emitted negligible fluorescence in the absence of target cancer cells since the fluorophore and quencher, conjugated to C-strands on either side of a cancer-specific TLS11a aptamer, lie close together. Then, in the presence of target cancer cells, the fluorophore and quencher separated, giving rise to a strong FAM signal. This AAFP may be a valuable tool for the early detection of cancer, which can
facilitate timely initiation of treatment. Using an AAFP avoids the need to conjugate a fluorophore to biological tissue, in contrast to other types of fluorescent probes, such as quantum dots (18,19), carbon nanodots (20) and fluorescent dyes (21,22). Under optimal incubation conditions, our AAFP was able to detect cancer cells at concentrations as low as ~100 cells/ml.

The TLS11a aptamer, although originally selected for its binding to mouse hepatoma BNL 1ME A.7R.1 (MEAR) cells, also binds with high affinity to human liver cancer cells. TLS11a is likely to be a membrane protein that can be internalized into cells (23), which may explain this dual binding specificity. We found that TLS11a on its own or as part of our AAFP bound efficiently to HepG2 cells, consistent with a previous study (24). In contrast, neither TLS11a nor AAFP bound to normal human L02 liver cells, confirming the cancer specificity of AAFP binding. Further confirmation came when we found that the mismatch-containing AAFP-mis failed to bind to either HepG2 or L02 cells. These results using flow cytometry and fluorescence spectroscopy to examine cells in suspension were confirmed using fluorescence microscopy to examine cells in culture.

We further confirmed the specificity of AAFP binding using frozen sections of HepG2 tumor xenografts and normal liver tissue. Frozen sections are often used for histopathological examination, in part since many proteins retain their activity as in fresh tissue sections, and since they are easier and faster to prepare than paraffin-embedded sections (25).

‘Activatable’ probes can be superior to ‘always on’ probes since they emit much less fluorescence in the absence of a target, thereby increasing the signal-to-background ratio (26,27). Our experiments with fluorescence spectroscopy revealed that AAFP emitted minimal fluorescence in the absence of HepG2 cells, likely reflecting fluorescence resonance energy transfer between the fluorophore and quencher due to aptamer binding to targets within the cancer tissue. The high signal-to-background ratio observed for AAFP, even at physiological temperatures, suggests its potential for highly sensitive cancer cell detection.

Optimization studies showed that AAFP binding efficiency depended on the length of the C-strands, the incubation temperature and incubation time. This suggests that studies of AAFP or similar probes may need to screen these parameters carefully to maximize sensitivity and specificity, and in particular to avoid false-negative results. Our observation of maximal binding efficiency at an AAFP concentration of 250 nM is consistent with a previous study (28). The fact that our AAFP bound noticeably less well at 37°C than at 4°C reflects the fact that the TLS11a aptamer was selected at low temperatures (17). This suggests that optimization of the TLS11a aptamer sequence may improve target binding at physiological temperatures. It also highlights the need to design AAFP-like probes using affinity sequences selected near *in vivo* temperatures.

The present study further demonstrated the power of aptamers in the detection and imaging of tumor tissues (29). The small size of aptamers may give them an advantage over antibodies for imaging intracellular targets (30), however, they may be rapidly degraded by nucleases or cleared from tissues. Chemically modifying aptamers can render them nuclease-resistant and increase their stability (31,32). Our results open the door to further investigations of AAFP with different chemical modifications.

In summary, we developed a simple, highly sensitive, specific AAFP for the detection of cancer cells and frozen cancer tissue. Our AAFP not only holds great potential in clinical early diagnosis, which can facilitate timely initiation of treatment, but also suggests the possibility of replacing the TLS11a aptamer with other sequences to target other tissue types in cancer or even other diseases.

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