A Cancer Cell-Activatable Aptamer-Reporter System for One-Step Assay of Circulating Tumor Cells

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The current antibody-mediated enumeration assays of circulating tumor cells (CTCs) require multiple steps and are time-consuming. To overcome these technical limitations, a cancer cell-activatable aptamer-reporter was formulated by conjugating a biomarker-specific aptamer sequence with paired fluorochrome-quencher molecules. In contrast to the antibody probes, the intact aptamer-reporter was optically silent in the absence of cells of interest. However, when used in an assay, the aptamer selectively targeted cancer cells through interaction with a specific surface biomarker, which triggered internalization of the aptamer-reporter and, subsequently, into cell lysosomes. Rapid lysosomal degradation of the aptamer-reporter resulted in separation of the paired fluorochrome-quencher molecules. The released fluorochrome emitted bright fluorescent signals exclusively within the targeted cancer cells, with no background noise in the assay. Thus, the assays could be completed in a single step within minutes. By using this one-step assay, CTCs in whole blood and marrow aspirate samples of patients with lymphoma tumors were selectively highlighted and rapidly detected with no off-target signals from background blood cells. The development of the cancer cell-activatable aptamer-reporter system allows for the possibility of a simple and robust point-of-care test for CTC detection, which is currently unavailable.

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Introduction

As a small molecule probe, aptamers are composed of short, single-stranded oligonucleotides (RNA or ssDNA) ranging from 30 to 60 bases.1–4 Similar to protein antibodies on the basis of their three-dimensional structures, aptamers specifically recognize and bind to their targets with high sensitivity and specificity.5–9 These targets can include small molecules, macromolecules, proteins, viruses, cells, and tissues.10–14 However, in contrast to antibodies, oligonucleotide aptamers are easily generated through chemical synthesis, and can be conveniently modified with a variety of functional molecules for different purposes. Therefore, aptamers have been widely studied for potential biomedical applications as a “chemical antibody”15–17 and, more importantly, for developing new clinical applications that cannot be performed with current protein antibody technology.18,19

Metastasis or dissemination of primary tumor is the major cause of mortality in cancer patients. Recent studies have demonstrated that circulating tumor cells (CTCs) in the bloodstream are key factors in the establishment of metastatic tumors.20–22 In addition, the number of CTCs present in whole blood is directly associated with cancer progression, recurrence, survival rate, and prognosis.23–25 Therefore, accurate detection of CTCs will provide critical information for the proper management of cancer patients. Antibody technology has been widely used for a variety of biomedical applications since it was first developed. The current CTC assays use antibodies as a specific probe to target tumor cells. These antibodies are preconjugated with fluorochrome as a reporter for imaging cell detection. However, the conjugated fluorochrome is constantly emitting a fluorescent signal, regardless of the status of whether the antibody is free or binding to cells, leading to a high level of background noise and potential off-target signals from normal blood cells. Therefore, the current antibody-mediated assays require multiple steps to isolate CTCs from normal blood cells and eliminate potential off-target signals. Repeated washes to remove excess free fluorochrome-conjugated antibody from the assays are also required. Notably, these multi-step assays are time-consuming and, more importantly, lead to loss of the blood sample and damage to CTCs, which adversely impacts assay accuracy. To overcome these technical obstacles in antibody-mediated assays, a new reporter system that can selectively highlight only tumor cells without generation of background noise is urgently needed.

Results

Design of the tumor cell-activatable aptamer-reporter

For point-of-care purposes, an “ideal” CTC assay should be (i) simple (one-step reaction), (ii) high throughput (complete in minutes), and (iii) accurate (no loss of blood sample and no damage of tumor cells through the assay process). However, no such assay could be developed using current antibody-mediated technology as described above. To this end,
we designed a unique aptamer-reporter, which is composed of an oligonucleotide sequence specific for a tumor cell biomarker conjugated with a reporter of paired fluorochrome-quencher molecules. The underlying hypothesis is that the intact aptamer-reporter is optically silent in the absence of tumor cells (Figure 1a), but can be activated by tumor cells to emit fluorescence. As showed in Figure 1b, in assays containing patient blood the oligonucleotide sequence will selectively target CTCs through interaction with a specific surface biomarker, which triggers internalization of the aptamer-reporter through a natural cellular process and, subsequently, into cell lysosomes that are rich in different enzymes. This internalization mechanism has been successfully used for intracellular delivery of chemotherapeutic drugs for targeted therapy by a newly FDA-approved antibody-drug-conjugate, brentuximab.26 Rapid degradation of the aptamer-reporter by lysosomal nucleases results in separation of the paired fluorochrome-quencher molecules, the released fluorochrome becoming optically active and emitting bright fluorescent signals exclusively within the targeted tumor cells. Notably, because it is controlled by a specific aptamer-biomarker interaction and an intracellular activation switch, the aptamer-reporter will generate no background noise or off-target signals from normal blood cells and thus, the assay can be carried out in one-step, load-then-read (Figure 1c).

**Tumor cell-activation and intracellular signaling of the aptamer-reporter**

For the initial proof-of-concept studies, the aptamer-reporter was formulated by conjugating a 39-mer RNA sequence specific for the CD30 biomarker27–29 with a fluorochrome Cy3 compound and a BHQ2 at the 5'- and 3'-ends, respectively.

![Figure 1](image_url) **Figure 1** Schema of the tumor cell-activatable aptamer-reporter for one-step assay of circulating tumor cells (CTCs) in a whole blood sample. (a) A biomarker-specific and tumor cell-activatable aptamer-reporter, which is composed of two functional units: an oligonucleotide aptamer sequence specific for the biomarker of interest, and a tumor cell-activatable reporter composed of paired fluorochrome-quencher molecules at the 5'- and 3'-ends of the same sequence, respectively. Due to covalent chemical conjugation, the intact aptamer is able to sufficiently hold the quencher molecule close to the paired fluorochrome, rendering the fluorochrome optically "inactive" in the absence of tumor cells. (b) Tumor cell-triggered intracellular activation of the aptamer-reporter. In assays containing tumor cells, the aptamer-reporter will selectively target and specifically bind to surface biomarkers, resulting in internalization of the aptamer-reporter into the targeted tumor cells and subsequently into cell lysosomes via a natural cellular process. The degradation of the aptamer-reporter by lysosomal nucleases will lead to separation of the paired fluorochrome-quencher molecules and release of the fluorochrome, emitting bright fluorescent signals exclusively within the tumor cells. (c) Proposed one-step assay for rapid detection of CTCs. Simply load patient whole blood into the assay that contains the optically silent aptamer-reporter, and CTCs are selectively highlighted and then detected within minutes.
(Figure 1a and Figure 2a, left). Due to covalent chemical conjugation, aptamer sequence is able to hold fluorochrome and quencher molecules close to each other and, thus, the BHQ2 molecule will quench fluorescent signals emitted from the Cy3 compound on the same aptamer. A control aptamer was also synthesized by conjugating the same sequence with the fluorochrome Cy3 compound alone (Figure 2a, right).

First, to validate the optical-activatable property, the aptamer-reporter was incubated with fresh lysates of cultured Karpas 299 tumor cells of human ALCL at room temperature as a cellular enzyme source. Changes in fluorescence were kinetically quantified. Incubation with cell lysates activated the aptamer-reporter and triggered a rapid increase in fluorescent signals in a time-dependent fashion (Figure 2b). In

Figure 2 Tumor cell activation and intracellular signaling of the aptamer-reporter. (a) As a demonstration model, a tumor cell-activatable aptamer-reporter was formulated by conjugating a CD30 biomarker-specific aptamer sequence with the fluorochrome Cy3 compound and quencher molecule BHQ2 at the 5'- and 3'-ends, respectively (left). A control aptamer was also synthesized by conjugating the same aptamer sequence with fluorochrome Cy3 compound alone (right). (b) To test activation potential, the formulated aptamer-reporter was incubated with fresh lysates of Karpas 299 tumor cells as a cellular enzyme source. Resultant changes of fluorescent signals in reactions containing the aptamer-reporter or the control aptamer were kinetically quantified and compared in graph. (c) To rule out structure change effect, the aptamer-reporter and the control aptamer were treated in different pH conditions from pH 4.0 to pH 7.4, and changes in fluorescent signals were kinetically monitored and graphed. (d) To test biostability, the aptamer-reporter and control aptamer were incubated in human serum alone or in the presence of fresh cell lysates. The changes in fluorescence were kinetically monitored and graphed. (e) To study tumor cell-induced activation, the aptamer-reporter was incubated with fresh culture of Karpas 299 cells (a human anaplastic large cell lymphoma expressing surface CD30 biomarkers) at room temperature, and cells were directly examined by fluorescent microscope at different time points as indicated. Quantified fluorescent signals of cells were showed in graph. (f) To confirm intracellular signaling, cells were treated with the aptamer-reporter and cell membranes were then stained with Alexa 488. Fluorescent signals derived from the aptamer-reporters (red) and cell membranes (green) were detected by confocal microscope. (g) Similarly, cells treated with the aptamer-reporter and cell lysosomes were then stained with Lyso-ID. Co-location of fluorescent signals emitted from the aptamer-reporters (red) and stained lysosomes (green) were detected by confocal microscope.
contrast, cell lysates had no effect on the control aptamer which constantly emitted high levels of fluorescent signals. To rule out a potential effect due to conformation change, the aptamer-reporter was treated under different pH conditions, from pH 4.0 to physiological pH 7.4, at room temperature for 30 minutes (Figure 2c). Notably, cell lysosomes have a low pH environment (pH 4–5) which can cause denaturing of the aptamer sequence. Change in pH had no effect on fluorescent emission of the aptamer-reporter or the control aptamer. These findings indicated that the formed aptamer-reporter can be optically activated by cell lysates, but not affected by the low pH-induced structure change. To further strengthen these observations, the control and aptamer-reporter were also incubated in human serum alone or in serum containing fresh cell lysates under. Kinetic analysis of fluorescence revealed that the aptamer-reporter was stable up to 90 minutes in human serum with minimal background, and was rapidly activated by the addition of cell lysates (Figure 2d).

To study tumor cell-induced activation, the aptamer-reporter was simply added into a culture of Karpas 299 cells, which express high levels of the CD30 biomarker on the cell surfaces. Cells were then directly examined under fluorescent microscope. As shown in Figure 2e (upper panel), the aptamer-reporter was rapidly optically activated by the tumor cells and, thus, highlighted the tumor cells. The fluorescence was initially observed at 10 minutes postincubation and lasted up to 120 minutes, with no background noise. Quantified fluorescent signals of tumor cells at different time points were calculated and summarized in graph in Figure 2e.

Subsequently, to validate intracellular optical activation, cells were treated with the aptamer-reporter at room temperature for 30 minutes and cell membranes were then stained with the green fluorescent dye, Alexa 488. Cells were directly examined by a confocal fluorescent microscope (Figure 2f). Fluorescent signals derived from the aptamer-reporters (red) and prestained cell membranes (green) were recorded.

Figure 3 Specific and selective highlighting of tumor cells by the aptamer-reporter. (a) To validate specificity, tumor cells with or without CD30-biomarker expression were used as indicated. The aptamer-reporter was selectively activated by and, thus, highlighted CD30-expressing tumor cells (left panels) and did not react to tumor cells that were negative for CD30 (right panels). Additional validation data with different tumor cells are listed in Table 1. (b) For a sensitivity test, a cell mixture was made by diluting CD30-expressing Karpas 299 tumor cells in CD30-negative U937 cells at ratios from 1:10 to 1:10,000. For identification purposes, U937 cells were pre-stained with green fluorescence. Cell mixtures were incubated with the aptamer-reporter and examined under a fluorescent microscope. The merged images revealed that the aptamer-reporter was sensitive and could selectively highlight tumor cells of interest (red) among many background off-target control cells (green).
separately. Merged images confirmed that the aptamer-reporter emitted fluorescent signals exclusively confined within the targeted cells. Moreover, to detect intracellular distribution, the cells were treated with the aptamer-reporter and cell lysosomes were then stained with the fluorescent dye, Lyso-ID. Merging of confocal microscopic images revealed the colocation of fluorescent signals from the aptamer-reporter (red) and prestained lysosomes (green) within the same cells (Figure 2g). These findings confirmed that the aptamer-reporter could be optically activated and emit fluorescent signals exclusively within the targeted cells.

One-step assay of CTCs in whole blood sample

To validate assay specificity, multiple cultured cells were incubated with the aptamer-reporter at room temperature for 30 minutes, and directly examined under fluorescent microscope. As shown in Figure 3a, the aptamer-reporter was activated by the tumor cells that express the CD30 biomarker and, thus, highlighted the tumor cells of SUDHL-1, HDLM2, and KMH2 cells, but remained optically silent in control cultures of Maver-1, U937, and CA46 cells that do not express CD30, with no background noise. Additional studies with carcinoma tumor cells and normal peripheral cells are summarized in Table 1. To assess sensitivity, a cell mixture was made by diluting Karpas 299 tumor cells at ratios from 1/10 to 1/10,000 in U937 control cells, which were prestained with green fluorescence (Figure 3b). The cell mixtures were incubated with the aptamer-reporter, and fluorescent signals derived from the aptamer-reporters (red) and prestained control cells (green) were directly recorded. Merged images revealed that the aptamer-reporters were selectively activated and highlighted by CD30-expressing tumor cells, but did not react to control cells in the cell mixtures. Taken together, these findings indicate that the aptamer-reporter was both specific and sensitive for detection of tumor cells.

For preclinical evaluation, peripheral blood and marrow aspirate were collected from patients with ALCL tumors under a protocol approved by the IRB for the Houston Methodist Research Institute. Diagnosis of ALCL was made by tissue biopsy, and CD30 expression on lymphoma cells was confirmed by immunohistochemical staining of tumor tissues with anti-CD30 antibody (Figure 4a). To detect CTCs of ALCL, whole blood samples were added into a 12-well plate (40 million total blood cells/well), which was preloaded with the aptamer-reporter (5 nmol/l final concentration). After incubation for 30 minutes at room temperature, plates were directly examined under a microscope. Blood cells were well preserved for morphologic identification, including intact RBC, white blood cells, and possibly rare tumor cells that could not be distinguished (Figure 4b). However, the CTCs in whole blood samples were selectively highlighted by the aptamer-reporter with bright fluorescence and without background noise or off-target signals from normal blood (Figure 4b, right panels). Similarly, patient marrow aspirate was also added into the assays, and CTCs were selectively highlighted by the aptamer-reporter as detected by fluorescent microscopy (Figure 4c).

Currently, antibodies have been used as the "gold standard" probe for tumor cell detection in the clinical setting. Double-staining of CTCs were performed to confirm specificity of the aptamer-reporter. Patient marrow cells were initially probed by the FITC-conjugated anti-CD30 antibody. After removal of free antibody from the reaction, the cells were then treated with the aptamer-reporter and directly examined under a fluorescent microscope. Cell signals derived from antibody (green) and the aptamer-reporters (red) were recorded separately (Figure 4d). Merged images revealed that the CTCs in patient marrow were double-stained by both antibody and the aptamer-reporters, confirming specificity of the one-step assay. Notably, the aptamer and antibody did not compete for the same binding site(s) on the same tumor cells.23 Additional blood specimens from different patients with ALCL tumor were also analyzed, and CTCs were specifically highlighted and detected with a high signal-background noise ratio (Figure 4e, upper panel). To validate assay capacity, a high-concentration blood sample was loaded into a 12-well plate (120 million blood cells/well) to form multiple layers of cells (Figure 4e, lower panel). Interestingly, cellular fluorescent signals were easily detected, although the highlighted CTCs were overlapped by normal blood cells and could not be observed under a light microscope. As a normal control, whole blood samples from healthy donors were tested under the same assay and showed no off-target signals (Figure 4f).

Discussion

It is generally believed that there are only a scant number of detectable CTCs in patient blood, although the actual number ranges are not yet fully known.20–22 A recent study demonstrated that the average number of CTCs ranges from a mean of 79 to 196 cells per ml in whole blood specimens of patients with different types of cancers, which is much higher than the cut-off cell number (5 cells/ml) detected by current

| Culture tumor cells | CD30-expressing status | One-step assay by aptamer-reporter |
|---------------------|------------------------|-----------------------------------|
| SU-DHL-1 (ALCL)     | +                      | +                                 |
| Karpas 299 (ALCL)   | +                      | +                                 |
| HDLM2 (Hodgkin lymphoma) | +                | +                                 |
| L-428 (Hodgkin lymphoma) | +                | +                                 |
| KMH2 (Hodgkin lymphoma) | +                | +                                 |
| L-540 (Hodgkin lymphoma) | +                | +                                 |
| CA-46 (Burkitt lymphoma) | –                 | –                                 |
| Jeko-1 (B cell lymphoma)  | –                   | –                                 |
| SUP-T1 (T cell lymphoma) | –                 | –                                 |
| U937 (Histiocytic lymphoma) | –                 | –                                 |
| Jurkat (T cell leukemia)  | –                   | –                                 |
| CCRF-CEM (T cell leukemia) | –                 | –                                 |
| SK-BR-3 (Breast carcinoma) | –                 | –                                 |
| T47D (Breast carcinoma)  | –                   | –                                 |
| LNCaP (Prostate carcinoma) | –                 | –                                 |
| VCap (Prostate carcinoma) | –                 | –                                 |
| PC-3 (Prostate carcinoma)  | –                   | –                                 |
| A-431 (Epidermoid carcinoma) | –                 | –                                 |
| HT-29 (Colon adenocarcinoma) | –                 | –                                 |
| Normal blood mononuclear cells | –               | –                                 |
| ALCL, anaplastic large cell lymphoma. | –               | –                                 |
commercial tests. In addition, employing the Cell Search Profile Kit could detect more than 30-fold of the median CTC counts that are identified by the clinically used Cell Search Epithelial Kit in the same blood samples of patients with breast or lung cancers. These findings indicate the presence of much more CTCs in patient blood than previously thought, suggesting the possibility that, with a sensitive detection approach, the current standard enumeration of CTCs can simply be justified by employing different-sized plates as clinically indicated with 96-, 24-, 12-, 6-, and single-well(s) to meet specific clinical needs. In addition, the load-then-read technology provides the possibility of performing high-throughput screening assays for rapid detection of CTCs. Moreover, good preservation of intact CTCs in this one-step assay will allow us to collect viable CTCs for further analysis of cellular protein and/or gene expression profiles at a single-cell level.

To eliminate background noise in whole-blood samples and enable intracellular highlighting of tumor cells, the one-step assay was designed by taking advantage of tumor cell internalization of the aptamer-reporter through a natural biological cell process, which is triggered by specific binding of aptamer sequences to surface CD30 receptors on tumor cells (Figure 1b). Interestingly, a similar internalization mechanism is employed for intracellular delivery of chemotherapeutic drug by an antibody-drug-conjugate, brentuximab, a newly FDA-approved, targeted therapeutic specific for CD30-expressing lymphomas. Thus, it is considerable that, in addition to the potential for diagnostic applications, aptamers can also be used as a chemical antibody for targeted therapy.

In summary, our study introduced a unique tumor cell-activatable aptamer-reporter technology and demonstrated a simple, one-step assay for detection of CTCs in patient whole blood. Since this technology improves upon the existing antibody-driven methods, we believe it holds a high potential in early identification of patients who are at a higher risk for
developing tumor metastases due to their elevated levels of systemic CTCs.

Materials and methods

Reagents and cells. In the proof of principal studies, a 39-mer RNA-based aptamer sequence, previously shown to specifically bind CD30-expressing tumor cells, was used.27-29 A novel tumor cell-activatable aptamer-reporter was then chemically synthesized (Bio-Synthesis, Lewisville, TX) by conjugating a fluorochrome Cy3 together with a Black Hole Quencher 2 (BHQ2) molecule at the opposing ends of the aptamer sequence as follows: Cy3-5′-rGrArUrCrArUrArGrUrGrUrGrUrCrArGrGrGrArGrGrGrArUrGrArUrCrG-3′-BHQ2. A control aptamer probe with the same sequence was conjugated with a fluorochrome Cy3 compound at the 5′ end. Alexa Fluor 488 conjugates for cell membrane staining were purchased from Invitrogen (Grand Island, NY) and Lyso-ID Green Detection Kit for lysosome membrane staining were purchased from Enzo Life Sciences (Farmingdale, NY). The FITC-conjugated anti-human CD30 antibody was purchased from BD Biosciences (Franklin Lakes, NJ).

Cancer cells lines, including human anaplastic large cell lymphoma (ALCL) cells (Karpas 299 and SUDHL-1 cell lines from Mark Raffeld at NIH), Hodgkin lymphoma cells (HDLM2 and KMH2 cell lines from Barbara Savoldo, Baylor College of Medicine, Houston, TX), B cell lymphoma cells (Mino and Mave-1 cell lines from ATCC), and leukemia cells (U937 cell line from ATCC) were cultured in RPMI 1640 medium supplemented with 10% FBS, 100 u/ml penicillin, and 100 µg/ml streptomycin at 37 °C under the atmosphere of 5% CO2 and ≥95% humidity.

Activation of the aptamer-reporter. Fresh tumor cell lysates were prepared to confirm the cell-activatable property of the formed aptamer-reporter. Briefly, cultured Karpas 299 cells were harvested, washed twice, and suspended in PBS (5 x 10^5/ml). Cells were frozen at −80 °C, thawed at room temperature twice, and centrifuged at 12,000g for 20 minutes at 4 °C. Finally, 100 µl supernatants of cell lysates were collected and added into wells of a 96-well black wall plate, which was preloaded with the aptamer-reporter (5 nmo/l final concentration). After incubation at room temperature, fluorescence signals of the assays were quantified by a Synergy H4 microplate reader (BioTek, Winooski, VT) at different time points, as described. Each condition was tested in triplicate and experiments were repeated more than three times with similar results. The mean value of fluorescence was calculated and shown as the mean ± SD. Similarly, the control aptamer, containing the same sequence and conjugated with the fluorochrome Cy3 compound alone, was tested under the same treatment conditions and change in fluorescent signals of the control group was kinetically monitored as described above. To rule out the effects of low pH conditions, which is seen in cell lysomes and might result in denature of the aptamer sequence 3D-structure, the aptamer-reporter (5 nmo/l final concentration) was added into PBS with pHs 4.0, 5.0, 6.0, 7.0, and 7.4 in a 96-well black wall plate. After incubation at room temperature for 30 minutes, fluorescence of the assays was quantified by a Synergy H4 microplate reader. Each condition was tested in triplicate and experiments were repeated more than three times with similar results. The mean value of fluorescence was calculated and shown as the mean ± SD. Finally, the aptamer-reporter (5 nmo/l final concentration) were incubated in 100 µl human serum (Atlanta Biological, Lawrenceville, GA, USA) in a 96-well black wall plate and changes in fluorescence were quantified as described above. In control group, 100 µl of cell lysate supernatants were also added into the human serum reaction and fluorescence was monitored. Each condition was tested in triplicate and experiments were repeated more than three times with similar results. The mean value of fluorescence was calculated and shown as the mean ± SD.

One-step assay and tumor cell detection. To validate the tumor cell-induced activation, 100 µl of cultured Karpas 299 cells (5 x 10^5/ml) were preseeded in a 96-well black wall plate in PBS, and 5 µl of the aptamer-reporter was then added to each well at 5 nmo/l final concentration. Without any additional preparation steps, the plates were directly examined under an Olympus IX81 fluorescent microscope and cell fluorescent signals were kinetically imaged using a TRITC filter under 547 nm of peak excitation wavelength and 572 nm of peak emission wavelength. For control purposes, cells were also treated with the control aptamer that was conjugated with fluorochrome Cy3 compound alone under the same condition. Fluorescent images were recorded using the same exposure time (300 microseconds) at different time points post treatment as described in Figures. Meanwhile, cell morphology was confirmed by light microscopy images. Each experiment with triple wells was repeated at least three times with similar results. For quantification, intensity of fluorescent signals from individual tumor cells was analyzed by Image J software (NIH, Bethesda, MD). A total of 90 cells from triplet wells (30 cells/well) at each time point were randomly gated and recorded. To get the background value of fluorescent intensity, 10 random noncellular locations with an average area of cell sizes were gated and calculated similarly. The mean fluorescent intensity of tumor cells was derived after background signals were deducted, calculated with standard deviation, and shown in graph with ± SD error bars.

Cell staining. For cell membrane staining, cultured Karpas 299 cells were treated with the aptamer-reporter at room temperature for 30 minutes. After washing once with PBS, the cells were suspended in PBS at room temperature, stained with Concanaavalin A conjugated to Alexa Fluor 488 (1 mg/ml) for 2 minutes, fixed with 4% formaldehyde for 10 minutes, washed with PBS, and suspended in PBS/glycerin (weight/weight ratio of 1:1). Finally, 15 µl of cell solution was loaded onto glass slides and examined by a confocal microscope to determine intracellular localization of the aptamer-reporter (FITC filter for Alexa Fluor 488 dye, TRITC filter for Cy3-labeled aptamer-reporters). For lysosomal detection, Karpas 299 cells were treated with the aptamer-reporter as described above. After washing once with PBS, cells were stained with Lyso-ID green dye (1:1,000 dilution) at room temperature for 1 hour, fixed with 4% formaldehyde for 10 minutes, washed with PBS, and then suspended in PBS/glycerin. Finally, 15 µl
of cell solution was loaded onto glass slides and examined by a confocal microscope as described above to determine possible colocalization of aptamer-reporters with the lysosomes.

Specificity and sensitivity validation. CD30-expressing cells (Karpas 299, SUDHL-1, HDLM2, and KM2) and CD30-negative control cells (Maver-1, U937, and CA46) were tested to confirm specificity of the aptamer-reporter. The cells were incubated with the aptamer-reporter for 30 minutes at room temperature and the fluorescence was detected as described above. Additional cell lines (breast, prostate, and colon carcinomas) and peripheral blood collected from healthy donors were also tested and are listed in Table 1.

To test sensitivity of the aptamer-reporter, cell mixtures were prepared from CD30-expressing Karpas 299 cells and CD30-negative U937 control cells. For tracking purposes, U937 cells were stained with ConcanaVail A conjugated to Alexa Fluor 488 and Karpas 299 cells were added in 1:10 to 1:10,000 ratios, as indicated. Subsequently, cell mixtures were added into 96-well black wall plates (2 x 10^5 cells/well) that were preloaded with the aptamer-reporting system (5 nmol/l final concentration). After 30 minutes incubation at room temperature, cells were examined by a fluorescent microscope (FITC filter for Alexa Fluor 488 to detect U937 cells, TRITC filter for Cy3 derived from aptamer-reporters).

Detection of CTCs in patient whole blood and marrow aspirate samples in a single-step reaction. Under an approved IRB protocol, peripheral blood and bone marrow samples were collected from patients (n = 6) with stage IV ALCL tumors diagnosed by a histological evaluation of H&E-stained tumor tissue and immunohistochemical demonstration of CD30 expression on tumor cells. Ten blood and marrow samples collected from healthy donors were used as a control group. Following cell counting, whole blood and marrow aspirate specimens were added into wells of a 12-well plate that contained the aptamer-reporter (5 nmol/l final concentration) for a total of 40 million blood or marrow cells/well. After incubation at room temperature for 30 minutes, single-step labeled CTCs were directly detected by a fluorescent microscope. For a morphological evaluation, CTC images were also obtained with a light microscope within the same view fields.

To confirm specificity of the aptamer-reporter, marrow cells were double-stained with FITC-conjugated anti-human CD30 antibody and the aptamer-reporter. Briefly, bone marrow cells were diluted in 1 ml PBS and incubated with FITC-conjugated anti-human CD30 antibody (BD Biosciences, San Jose, CA) at room temperature for 30 minutes. To eliminate unbound antibody and minimize damage to CTCs, the reaction supernatant containing some red blood cells (RBCs) was gently removed by a pipette. The remaining cells were then suspended in 200 μl PBS and added into well plates that were preloaded with the aptamer-reporter at a 5 nmol/l final concentration. After 30 minutes incubation at room temperature, plates were directly examined by a fluorescent microscope and cell fluorescence (green: antibody, red: aptamer-reporter) recorded separately. Merged images showed that CTCs in patient marrow were double-stained, confirming specificity of the aptamer-reporter.

To test the effect of blood concentration, a total of 120 million blood cells from patients diagnosed with ALCL were added into 12-well plates to form multiple cell layers. After 30 minutes incubation with the aptamer-reporter, plates were examined by light and fluorescent microscopes to detect CTC fluorescence as described above. Merged images indicated that the assay was able to detect CTCs in high-concentration blood samples, even when CTCs overlapped with normal blood cells and could not be observed under a light microscope. To validate these results, whole blood samples from healthy donors were also tested in the single-step assay, which revealed no false-positive or off-target staining in the normal blood cells.

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