RNA Aptamers Targeting Integrin α5β1 as Probes for Cyto- and Histofluorescence in Glioblastoma

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Nucleic acid aptamers are often referred to as chemical antibodies. Because they possess several advantages, like their smaller size, temperature stability, ease of chemical modification, lack of immunogenicity and toxicity, and lower cost of production, aptamers are promising tools for clinical applications. Aptamers against cell surface protein biomarkers are of particular interest for cancer diagnosis and targeted therapy. In this study, we identified and characterized RNA aptamers targeting cells expressing integrin α5β1. This αβ heterodimeric cell surface receptor is implicated in tumor angiogenesis and solid tumor aggressiveness. In glioblastoma, integrin α5β1 expression is associated with an aggressive phenotype and a decrease in patient survival. We used a complex and original hybrid SELEX (selective evolution of ligands by exponential enrichment) strategy combining protein-SELEX cycles on the recombinant α5β1 protein, surrounded by cell-SELEX cycles using two different cell lines. We identified aptamer H02, able to differentiate, in cyto- and histofluorescence assays, glioblastoma cell lines, and tissues from patient-derived tumor xenografts according to their α5 expression levels. Aptamer H02 is therefore an interesting tool for glioblastoma tumor characterization.

INTRODUCTION

Glioblastoma (GBM), the highest-grade glioma tumor (grade IV), is the most aggressive and the most common malignant form of astrocytoma. Standard therapy consists of surgical resection to an extent that is safely feasible, followed by radiotherapy and concomitant chemotherapy with temozolomide (Stupp protocol).1 Despite these therapies, patients with GBM rarely live longer than 2 years.2 Histological features that characterize GBM are the presence of necrosis and abnormal growth of blood vessels around the tumor. Defining molecular profiles aims to develop molecularly guided approaches for the treatment of patients. The 2016 World Health Organization (WHO) classification scheme3 integrated phenotypic and genotypic parameters for CNS tumor classification. GBMs are divided into isocitrate dehydrogenase (IDH) 1 wild-type (about 90% of cases; corresponds to the most frequent primary or de novo GBM) and IDH1-mutant GBM (about 10% of cases; corresponds to secondary GBM). Some of the GBM biomarkers that have been and are being discovered4 are cell surface protein biomarkers.5–8 Expression of cell surface protein is often remodeled in cancers. Genetic and epigenetic features altered in cancer include modification of copy number (under- or overexpression), truncations, mutations, and post-translational modifications. These modified proteins are major clinical targets for diagnosis and therapies, considering their accessibility for pharmacological compounds.

Tumor-specific tools such as aptamers can be used as recognition ligands to discriminate a tumor cell from another cell, as agonists or antagonists, or as carriers to deliver therapeutic payloads to targeted tumor cells.9–13 Aptamers are single-stranded DNA or RNA molecules that constitute an alternative class of molecules emerging as cancer-specific therapeutic, diagnostic, and theranostic tools.14,15,18 They are selected through an in vitro selection process, published for the first time in 1990 by three independent research groups.19–21 known as SELEX (selective evolution of ligands by exponential enrichment).5,18 Aptamers17 from the Latin aptäre (to fit) and from the ancient Greek meros (part) are often referred to as chemical antibodies15 because they bind to their specific targets with high affinity and specificity. Aptamers possess numerous advantages over antibodies, like smaller size, temperature stability, self-refolding, fewer side effects for immunotherapy, lack of immunogenicity and toxicity, more efficient penetration into biological compartments, chemical synthesis with high batch fidelity, and the option of site-specific and flexible introduction of linkers, reporters, functional groups.
small interfering RNA (siRNA), nanoparticles, drugs, and so forth. Apolmers toward a wide variety of targets have been identified, the most common ones remaining proteins. We recently reviewed aptamers to more than 30 different tumor cell surface protein biomarkers, a few of them being heterodimeric receptors, such as tyrosine kinase receptors and cell adhesion molecules. However, selection of aptamers to cell surface proteins remains a complex process.

Among cell surface biomarkers, integrins are heterodimeric cell surface receptors for cell migration, differentiation, and survival, composed of α and β subunits; their deregulation leads to cancer progression and therapy resistance. In mammals, 24 distinct integrins are formed by the combination of 18 α and 8 β subunits. Specific heterodimers preferentially bind to distinct extracellular matrix proteins. Integrin α5β1, the fibronectin receptor, belongs to the arginine, glycine, and aspartate (RGD)-binding integrin family. Overexpressed on tumor neoepithelia and on solid tumors, integrin α5β1 is implicated in tumor angiogenesis and solid tumor aggressiveness. We and others have shown that α5β1 integrin is a pertinent therapeutic target for GBM through its active role in tumor proliferation, migration, invasion, and resistance to chemotherapy. At the mRNA level, high α5β1 integrin expression is associated with more aggressive tumors in patients with glioma. At the protein level, to date, only a few in situ analyses of GBM tumor section have been described. Further investigation of α5β1 expression in GBM tumor cells as a potential prognostic factor and/or biomarker for diagnosis requires rapid and accurate tools.

In our study, aptamers to integrin α5β1-expressing cells were selected by an original and complex hybrid SELEX process. This SELEX combines three rounds of protein-SELEX surrounded by 15 rounds of cell-SELEX on two different cell lines genetically modified to overexpress integrin α5, the human GBM U87MG cell line, and the Chinese hamster ovary CHO-B2 cell line. Counterselection steps were performed on isogenic cell lines underexpressing α5 for U87MG or ones that do not express α5 for CHO-B2. We identified and characterized an aptamer named H02. Directly coupled to the cyanine 5 fluorophore, aptamer H02 was able to discriminate between 10 GBM cell lines expressing high and low levels of integrin α5. Aptamer H02 is internalized at 37°C. As a proof of concept, we also demonstrated that aptamer H02 is very efficient in afa-fluorescence assays to distinguish GBM tumor tissues from patient-derived tumor xenografts expressing high levels of α5 from GBM tumor tissues expressing low levels of α5.

RESULTS

Identification of Aptamers Binding to α5-Expressing Cells

Aptamer H02 was the most frequently represented over all sequences (6 times over 82 sequences, 7.3%). Aptamers H03, G10, and G11 represented 3.7% of all sequenced molecules and aptamer B03 1.2%. The predicted secondary structures of the five aptamers (H02, G11, B03, G10, and H03) are shown in Figure 2A. Fixed regions (shown in dark red in Figure 2A) were designed to display partial complementarity and pre-organize aptamers in hairpin structures. The secondary structure predictions of aptamers H02, G10, and B03 are highly similar and very different from those of G10 and H03.

Identification of α5β1-binding aptamers was performed using confocal fluorescence microscopy by incubating cyanine-5 (Cy5)-labeled aptamers at 4°C with the cells used for the cell-SELEX process. None of the five aptamers binds to U87MG α5− and CHO-B2 cells used for the counterselection steps (Figures S2A and S2B). Only aptamer H02 binds to the U87MG α5+ and CHO-B2 α5+ cells used for positive selection steps (Figures 2B and 2C). We next checked
whether, under the same methodological conditions, these aptamers were able to bind to U87MG cells at 37°C, the temperature used for the cell-SELEX process. H02, G11, B03, G10, and B03 did not bind to U87MG cells (Figure 3A). On U87MG+ cells, we observed strong binding not only of aptamer H02 but also, to a lesser extent, of aptamer G11 (Figure 3B). Because of its binding to U87MG+ cells at 4°C and at 37°C, subsequent analyses were performed using aptamer H02.

Validation of Integrin α5β1 as the Target of Aptamer H02

The SELEX process was performed to guide the selection toward integrin α5β1. However, cell-SELEX-based strategies have already led to the selection of aptamers against “undesirable targets,” meaning other proteins than the expected pre-identified targets.36,37

To validate integrin α5β1 as the target of our SELEX process, surface plasmon resonance (SPR) experiments were performed on a Biacore T200 instrument. Figure 4A shows the experimental scheme. Aptamer H02, with 2’-fluoro pyrimidines to increase its stability in SPR experiments, was captured on a CAP sensor chip via a biotin at its 5’ extremity. The aptamer was captured on a biotin CAPture reagent. Integrins were injected at concentrations ranging from 8–130 nM. We tested human integrins α5β1 and αvβ3 to check the H02 aptamer’s specificity. The surface was then
Figure 2. Aptamers Predicted Secondary Structures and Binding to α5-Expressing Cells

(A) Predicted secondary structure of aptamers H02, G11, B03, G10, and H03. Structures were predicted using the mfold web server.\textsuperscript{67} Nucleotides 1–19 and 50–68 correspond to fixed flanks of the candidate sequences. They are shown in dark red. ΔG values are noted above the structures. (B and C) Monitoring of the binding of five Cy5-labeled aptamers (H02, G11, B03, G10, and H03) at 5 μM on CHO-B2 and U87MG cell lines at 4°C using confocal microscopy. Nuclei, counterstained with Hoechst, are shown in royal blue. Aptamers, coupled to Cy5, are shown in white. (B) Binding on CHO-B2 α5+. α5 integrin is visualized, using the GFP-fused protein, in green. (C) Binding on U87MG α5+. α5 integrin was labeled with the IIA1 antibody and is shown in green. Images were captured at the same setting to allow direct comparison of staining patterns.
CAP chips are designed to capture biotinylated molecules reversibly on the sensor surface, facilitating its regeneration. An SPR cycle thus consisted of injections of biotin CAPture reagent, biotinylated aptamer, integrin, and regeneration solution. Successive cycles were repeated, changing the integrin nature and concentration at each cycle (Figure S3A). Because of 2'-fluoro modifications, aptamer H02 was resistant to degradation over time, and aptamer injections were highly reproducible. The biotinylated aptamer reached the same level at each cycle with as low as 2% variation in responses over 20 cycles (Figure S3A). Therefore, the surface has the same properties during all experiments.

Figures 4B and 4C show the sensorgrams obtained for integrins α5β1 and αvβ3, respectively, after double referencing (subtraction of reference channel and buffer injection). The sensorgrams show that, even when responses are low, aptamer H02 bound specifically to integrin α5β1 in a dose-dependent manner but failed to interact with integrin αvβ3. The equilibrium affinity parameter (K_D) of the interaction between integrin α5β1 and aptamer H02 was 72 ± 11 nM. To ensure that integrin was active, positive controls were used. Figure S3B shows binding by SPR of integrin α5β1 to its natural ligand, fibronectin. We also demonstrated that only aptamer H02 was an α5β1 binder (but not, for example, aptamer B03; Figure S3C).

H02 Aptamer Stability, Specificity and Affinity for U87MG α5+ Cells
Because aptamer H02 is a non-modified RNA molecule, we tested its stability in the presence of cells at 4°C and 37°C in the buffer used for selection. The results (Figure S4A) confirmed that incubation on cells for 1 h, which corresponds to the maximum contact time of RNAs with cells during different assays, does not induce aptamer H02 degradation. However, this aptamer is extremely rapidly degraded when incubated on cells in a culture medium supplemented with 10% fetal bovine serum (FBS) (Figure S4A). If this aptamer had to be used under more complex conditions than in a simple buffer, then it would have to be modified to increase its nuclease sensitivity. For example the 2'-fluoro-modified H02 aptamer is very stable in contact with cells at 4°C and at 37°C as it is not degraded in the selection buffer and in a complex medium for at least 1 h (Figure S4B).
The specificity of aptamer H02 for α5-overexpressing cells was confirmed by flow cytometry at 4°C by incubation of Cy5-coupled aptamers B03 and H02 at 500 nM for 1 h with detached CHO-B2 cells (Figure 5A) and by incubation of Cy5-coupled aptamers G11 and H02 at concentrations ranging from 0.15–5 μM with detached CHO-B2 α5+ cells (Figure S5). Although no shift in fluorescence was detected for CHO-B2 cells after incubation with the two Cy5-labeled H02 and B03 aptamers, a shift was detected for CHO-B2 α5+ cells with aptamer H02 but not with aptamer B03. Figure S5 also confirmed that aptamer H02 was our best hit from SELEX because it binds much better to α5-expressing cells than aptamer G11.

The equilibrium affinity parameter K_D of the interaction between aptamer H02 and U87MG α5+ cells was determined using flow cytometry (Figure 5B). Binding events associated with the fluorescence signal of different concentrations of aptamer, ranging from 0.15–5 μM, to a constant number of cells were measured. A K_D of 277.8 ± 51.8 nM was determined by plotting the mean fluorescence of U87MG α5+ cells against the concentration of the H02 aptamer.

H02 Aptamer Localization in U87MG α5+ Cells

The localization of aptamer H02 on GBM U87MG α5+ cells was analyzed by confocal microscopy at 4°C and 37°C. Cells were incubated with the Cy5-labeled aptamer H02 and with the anti-α5 IIA1 antibody, followed by incubation of a secondary antibody labeled with Alexa 546. Spots of co-localization were detected between Alexa 546 and Cy5, which reflect spatial proximity between the α5 subunit and the aptamer H02, at 4°C and, to a lesser extent, at 37°C (Figures 6A and 6B). Aptamer H02 detected α5β1 mostly at the plasma membrane and at cell-cell junctions at 4°C, whereas punctuate labeling suggested internalized molecules at 37°C. The weaker co-localization observed with anti-β1 antibody and aptamer H02 (Figure 3B) is explained as the β1 subunit associates with different α subunits to form different integrins, whereas the α5 subunit associates only with the β1 subunit to form the fibronectin receptor.

We next wanted to confirm the internalization of aptamer H02 in α5-expressing cells at 37°C. To this end, adherent U87MG α5+ cells were labeled for 30 min with the Cy5-coupled aptamer H02 at five different concentrations (5, 2.5, 1.25, 0.6, and 0.3 μM). After cell fixation, cells were immunolabeled with the anti-EEA1 antibody to detect early endosomes and then analyzed by confocal microscopy. Figure 6C shows clear co-localization of aptamer H02 at 5, 2.5, and 1.25 μM with the anti-EEA1 antibody in U87MG α5+ cytoplasm, suggesting aptamer H02 endocytosis. A 3D reconstruction of whole z stacks is shown in Video S1.

Fluorescently labeled aptamers were not detected at lower concentrations (0.6 and 0.3 μM). The lower concentration limit of 1.25 μM corresponds to 4.5-fold the K_D of the H02-cell interaction and, theoretically, to 82% receptor occupancy, governed by concentration and affinity. At 4°C, aptamers were not detected at concentrations lower than 5 μM (Figure S6), suggesting a different binding mechanism at 4°C and at 37°C. Figures 5A and S5 show that, in flow cytometry experiments, a difference could be detected at 4°C between H02 and aptamers B03 and G11 at concentrations lower than 1 μM. This difference may be due to the differences inherent to the two different techniques (flow cytometry versus confocal microscopy).
Aptamer H02 is therefore an interesting and promising new tool to differentiate cells according to their α5 subunit expression levels in cytto- and histofluorescence experiments.

DISCUSSION

Biomarkers are indicators used to establish a diagnosis and prognosis and predict susceptibility to targeted therapies. GBM is the most aggressive form of brain tumors in adults. Despite intensive treatments, the prognosis of GBM patients remains poor. There is an urgent need to incorporate known biomarkers into clinical trials and routine clinical practice, which may assist not only with patient selection but also with adjustment of the treatment schedule based on patient-specific biology. Because differential expression of cell surface proteins often occurs in tumor cells, and considering their accessibility to extracellular ligands, these proteins provide biomarkers of interest in oncology. The identification of molecular probes specific for cell surface protein biomarkers is of great importance. Because of their high affinity and specificity toward their targets, aptamers are attractive and promising tools, alternatives to antibodies for clinical applications. In this study, using a complex and highly stringent SELEX strategy, we showed that it was possible to select an RNA aptamer specific to a pre-identified heterodimeric
Figure 6. Confocal Microscopy Analysis of Aptamer H02 on U87MG α5+

(A) Confocal microscopy analysis with aptamer H02 at 5 μM and the anti-α5 antibody IIA1 on U87MG α5+ cells at 4°C and 37°C. The aptamer is labeled with Cy5 (white). Incubation of antibody IIA1 was followed by incubation with a secondary antibody labeled with Alexa 546 (green). Nuclei are stained with Hoechst (blue). (B) Top: enlargement of the merged image in (A) at 4°C. Bottom: magnified images acquired by zooming in on the indicated areas of the parental image. Scale bars are shown in the lower right corners of the images. (C) Co-localization of aptamer H02 and the endocytosis marker EEA1. Shown are confocal images of U87MG α5+ cells incubated at 37°C with the

(legend continued on next page)
cell surface protein embedded in its natural environment. Integrin α5β1 is a GBM cell surface biomarker.\(^{26}\) Aptamer H02 was able to differentiate between high and low expression of the α5 integrin on cells and tissues. This aptamer is suitable for tumor characterization.

Two main processes have been developed to select aptamers specific for pre-determined cell surface proteins: protein- and cell-based SELEX.\(^{22}\) In protein-based SELEX, a purified protein is used as target, usually full-length or truncated (generally recombinant ectodomains). The major issues with the protein-based SELEX process are that purification of membrane proteins is not easy and that purified proteins may not adopt the same conformational state as in their endogenous cellular environment. Some aptamers identified through protein-based SELEX failed to recognize their target when embedded in whole living cells.\(^{42,43}\) In cell-based SELEX, targets are cell surface proteins based on the same version of the target in its purified form by protein-based SELEX.\(^{50,52}\) A reverse hybrid SELEX combines protein-SELEX followed by cell-SELEX.\(^{51,53,54}\) Consequently, our strategy combines hybrid SELEX and reverse hybrid SELEX (Figure 1). The second originality is that two different cell lines were used compared with one in previous studies. These two cell lines were the human GBM U87MG and the hamster CHO-B2 cell lines. Only three integrins (β2, αvβ3, and α6β4) have so far been used as pre-identified targets for the SELEX processes. Blind et al.\(^{48}\) selected, by protein-SELEX, RNA aptamers targeting a 46-mer peptide corresponding to the complete cytoplasmatic domain of the β2 subunit of integrin αLβ2. Cells infected with vaccinia viruses encoding β2-specific aptamers enabled high-level cytoplasmic expression of RNA aptamers. Intracellular integrin-binding aptamers reduced inducible cell adhesion to the intercellular adhesion molecule 1 (ICAM-1). To target integrin αvβ3, two different protein-based SELEX processes and a cell-based SELEX were used to identify 2′-fluoropyrimidine RNA aptamers. The 2′-fluoropyrimidine modification confers increased nuclease resistance to RNA molecules. Aptamer APT-αvβ3, selected on αvβ3 purified by immunoaffinity chromatography, was able to bind αvβ3 integrin expressed on the surface of live cells and to impair endothelial cell growth and survival.\(^{45,47}\) To select aptamers specific to homodimer αv and β3, Gong et al.\(^{48}\) developed a strategy called multivalent aptamer isolation (MAI)-SELEX. Two distinct selection stages were employed, the first being a classical affinity selection on the purified full-length αvβ3 integrin. The second, for specificity, leads selection to β3 because integrin α1β1β3 served as a protein decoy. Two aptamers specific for αv and β3 were identified with affinities in the low nanomolar range. Takahashi et al.\(^{49}\) applied a process they called isogenic cell (Icell)-SELEX to identify RNA aptamers targeting αv integrins (integrin alpha subunit [ITGA1]), in which isogenic HEK293 cell lines were manipulated for counterselection by microRNA-mediated silencing and for positive selection by overexpression of target proteins. Integrin α6β4 has recently been the target of hybrid SELEX,\(^{50,51}\) a combination of protein- and cell-based SELEX processes, for which five rounds of cell-SELEX on PC-3 cells were followed by 7 rounds of protein-based SELEX on a recombinant α6β4 protein.\(^{52}\) In this last study, despite introduction of counterselection on PC-3 β4 integrin (ITGB4) knockout cells to deplete single-stranded DNA (ssDNA) aptamers specific for cell surface markers other than the β4 subunit, the cell-SELEX process alone was not sufficient to prevent enrichment of non-target-specific aptamers.

To allow discovery of highly selective but also conformation-dependent aptamers and to guide the selection toward α5β1 integrin, the complex SELEX strategy we adopted presents two originalities compared with other SELEX strategies toward cell surface biomarkers. In our study, a hybrid SELEX combining successive rounds of cell-SELEX, protein-SELEX, and then again cell-SELEX was performed. Usually, in hybrid SELEX, the first rounds of selection are performed by cell-SELEX, and then rounds of selection are performed on the same version of the target in its purified form by protein-based SELEX.\(^{50,52}\) A reverse hybrid SELEX combines protein-SELEX followed by cell-SELEX.\(^{51,53,54}\)

Cy5-labeled aptamer H02 at five different concentrations (5, 2.5, 1.25, 0.6, and 0.3 μM). After aptamer labeling (shown in white), cells were fixed, permeabilized, and then labeled to detect nuclei (DAPI, blue), actin (Phalloidin-Atto 488, green), and early endosome EEA1 (EEA1 immunolabeling, red). Shown in the first row are merged images. Shown in the second row are magnified images of selected areas (white squares) of the parental images. These images show co-labeling of EEA1 and aptamerH02. Shown in the third row are separate EEA1 and aptamer labeling. Images were captured at the same setting to allow direct comparison of staining patterns.
Figure 7. Aptafluorescence on GBM Cells and Tissues

(A–D) Aptafluorescence on GBM cells (A and B) and tissues (C and D). (A) Western blot analysis. One representative western blot of the GBM cell lines (U87MG α5+, U87MG α5−, LN319, LN229, SF763, LN18, LN229, LNZ308, LN18, LNZ308, U373, LN443, and T98G) used in this study is shown at the top. Histograms, at the bottom, show the quantification of α5 integrin expression normalized to GAPDH levels (mean ± SEM of 3 independent experiments). GAPDH was used as a loading control. (B) Immuno-quantification versus apta-quantification of confocal fluorescence experiments on ten GBM cell lines. Immuno-quantification was performed with an anti-α5 antibody, followed by a secondary antibody coupled to Alexa 546. Aptafluorescence was performed with the Cy5-labeled aptamer H02. Quantification of mean fluorescence intensity (a.u.) was performed on at least 5 randomly selected images per cell line. The correlation coefficient is 0.78, with p < 0.0001. (C) Immuno-fluorescence (top panel) and apta-fluorescence (bottom panel) of patient-derived tumor xenografts TC7 and TC22, showing high and low levels of α5 integrins, respectively. For immunofluorescence, detection of α5 (in white) was performed with an antibody, followed by a secondary antibody coupled to Alexa 647. For aptafluorescence, detection of α5 (in white) was performed with the Cy5-labeled aptamers H02 and G11. DAPI staining is shown in blue. Images were captured at the same setting to allow direct comparison of staining patterns. One representative image per condition is shown. (D) Quantification of immuno-fluorescence (top panel) and apta-fluorescence (bottom panel) by confocal microscopy. Histograms show quantification of 10 to 26 different images per condition. Statistical analyses were done with Student’s t test (***p < 0.0001 and **p < 0.003).
Aptamer H02 was selected after 18 rounds of a stringent SELEX process. This aptamer was the most represented sequence. It is not degraded in contact with cells under the conditions used for the experiments. As for aptamers G11 and B03, the predicted secondary structure of aptamer H02 is highly stable in imperfect hairpins. Using label-free SPR experiments, integrin α5β1 was validated as the target of aptamer H02 (Figure 4B). Aptamer H02 was also identified as a binder of α5-expressing cells on cells used for positive selection or on other GBM cells in aptacytochemistry assays (Figures 6, 7A, and 7B). U87MG α5+ cells incubated with aptamer H02 allowed internalization of aptamer H02 at 37°C by endocytosis of the α5β1 cell surface receptor. A Kd value of 277.8 ± 51.8 nM was determined for the interaction between aptamer H02 and U87MG α5+ cells. This affinity value is of the same order of magnitude than the 100–400 nM that have been determined for aptamers characterized toward other integrins by cell- or hybrid SELEX. By SPR, a Kd of 72 nM was determined for the interaction between aptamer H02-2F and integrin α5β1. The 3.8-fold difference in binding affinity between the aptamer-recombinant protein and aptamer-cell interaction can certainly be explained by the different techniques used (flow cytometry versus SPR) and by conformational differences between cell surface proteins and soluble recombinant proteins. However, it is not due to the use of the H02-2F aptamer in SPR versus the non-modified aptamer in flow cytometry because the H02-2F and non-modified H02 aptamers have the same affinity for U87MG α5+ cells (data not shown). Only a few studies compared aptamer binding with isolated proteins and with tumor cell surface protein biomarkers.22 The 5-fold difference, which is of the same order of magnitude than the difference observed in our study, has been described for the OX40-aptamer interaction and can be explained by conformational differences.23

In the field of precision oncology, histological detection of specific biomarkers is a crucial diagnostic tool. Immunochemistry is a cheap, easy method for detection of tumor biomarkers. Aptahistochemistry is a new option, still rarely described, for which aptamers, as a new class of probes, are used instead of antibodies. In our study, we used aptamer H02 directly end-labeled with a single cyanine 5 fluorescent dye. Aptamer H02 was able to specifically interact with α5-overexpressing tumor tissues from patient-derived xenografts (Figures 7C and 7D) because it efficiently differentiates TC7 (tissue with high α5 expression) from TC22 (tissue with low α5 expression). Aptamer H02 is an effective molecular probe for labeling histological tissue sections and detection of the α5β1 biomarker on tumor cells. Aptamer probes may become powerful tools for pathologists to characterize tumor tissues because the protocols are simple to implement, straightforward to automate, and can be applied to paraffin-embedded cancer tissue as well as to frozen tumor tissues.26-28 Aptahistochemistry could certainly be easily extrapolated to biomarker multiplexing detection28 to assess co-localization of different markers on the same tumor section. Aptamer H02 targeting integrin α5β1 as well as other aptamers targeting other GBM biomarkers might therefore help to better characterize GBM inter-tumoral as well as intra-tumoral heterogeneity, which would have implications for personalized targeted therapies. A recent review describes the technicalities of the current applications of aptahistochemistry. Aptaffluorescence will probably reduce the cost, time, and cross-reactivity concerns compared with indirect immunofluorescence approaches generally based on primary and then secondary antibodies. Conjugation of dye on aptamers is easy and reduces the risk of disrupting the aptamer structure compared with antibodies generally labeled with multiple tags. Compared with an antibody, the aptamer’s smaller size (10-fold reduction in size) allows better penetration in tissues, particularly in applications for which epitope accessibility is reduced, such as in fixed tissues. A further advantage of aptamers is that the target of interest is not limited to molecules that produce an immune response in the host animal, as for antibodies.29 Chemical synthesis of aptamers virtually eliminates the issue of batch-to-batch variation.

Conclusion
In conclusion, we characterized a new, original, and powerful aptamer tool to detect GBM tumoral cells and tissues expressing integrin α5β1. These detections might be extended for use in other cancers where α5β1 has proven to be a therapeutic target, such as colon cancer, ovarian cancer, breast cancer, lung tumors, and melanoma.30 For clinical translation, the structure of aptamer H02 will have to be confirmed, and aptamer H02 will be improved further by truncations to obtain the minimal active fraction and by increasing its resistance toward nucleases via modifications of its nucleic acid backbone and extremities. Internalized, an aptamer targeting integrin α5β1 might open roads for α5β1-specific therapeutic payload delivery. Endocytosis may be crucial for targeting and increasing the therapeutic efficacy of GBM drugs. Linked to a cytotoxic agent, an aptamer to integrin α5β1 could serve as a carrier for targeted therapeutic delivery. Such aptamers were very recently called “charromers.” Charromers internalized with integrin α5β1 would be very powerful carriers to deliver therapeutic agents into targeted cells.

MATERIALS AND METHODS
Materials
The ssDNA library was synthetized and purified by Eurogentec (Seraing, Belgium). All RNA aptamers and chemicals were purchased from IBA and Sigma-Aldrich (Hamburg, Germany), respectively, unless otherwise stated. The sequences of all primers, the library, and aptamers from this study are described in Table S2.

Cell Culture and Transfection
Cell culture medium and reagents were from Lonza (Basel, Switzerland) or Gibco (Thermo Fisher Scientific, Waltham, MA, USA). U87MG cells were from the ATCC. U373MG and T98G cells were from ECACC (European Collection of Authenticated Cell Cultures, Sigma). LN319, LN229, LN443, LN18, and LNZ308 cells were kindly provided by Prof. Monika Hegi (Lausanne, Switzerland), SF763 by Frédéric André (Marseille, France), and CHO-B2 by
Wolfram Ruf (La Jolla, CA, USA). All GBM cells were routinely cultured in Eagle’s minimum essential medium (EMEM), 10% heat-inactivated FBS, and 2 mM glutamine. For U373MG and T98G, 1% non-essential amino acids and 1 mM sodium pyruvate were added to the medium. For CHO-B2 cells, EMEM was substituted for DMEM (high glucose). U87MG cells were stably transected to overexpress (U87MG z5+) and repress (U87MG z5−) the human z5 gene, as described previously. CHO-B2 cells lacked the z5 subunit. They were stably transfected by pcDNA3.1 plasmid provided by Dr. Ruoshlati (La Jolla, CA, USA) to overexpress the human z5 integrin gene in fusion with the gene for GFP by using jetPRIME (Polyplus transfection) according to the manufacturer’s instructions and named CHO-B2 z5+ cells.

Expression and Purification of z5β1-Fc
The recombinant soluble human z5β1-Fc integrin (a gift from Martin Humphries, Manchester, UK) was produced from NSO culture supernatant and purified via the Fc domain on protein A-Sepharose as described previously. The purity of the protein was verified by Coomassie staining of SDS-polyacrylamide gels.

SELEX Strategy
The RNA library was obtained by transcription from a starting ssDNA library (Eurogentec) containing 30 random nucleotides (N30) and flanked by primer annealing sites: 5’-GTGTGAC CGACCGTGGTGC-N30-GCAGTGAAGGCTGGTAACC-3’. Two primers, P3’ (5’-GTGTGACCGACCGTGGTGC-3’) and P5’ (5’-TA ATACGACTCAGATAGTTACCGCCCTGACTGC-3’), containing the T7 transcription promoter (underlined) were used for PCR amplification as described previously. Synthesis of the RNA library and transcription followed by DNase I (Roche) treatment have been described previously. The RNA pool was gel purified by denaturing (7 M urea) gel electrophoresis on an 8% polyacrylamide gel. Unbound sequences were then incubated with the starting RNA library (1 nmol) at 37°C for 2 min. Eluted RNA was recovered, reverse transcribed, PCR amplified, and transcribed back into RNA with the T7 transcription primer (Cy5-labeled aptamer), denatured at 95°C, and incubated with protein A-Sepharose beads alone prior to positive selection. Unbound sequences were incubated under agitation on z5β1-Fc-loaded beads for 20 mins. Eluted RNA was recovered, reverse transcribed, PCR amplified, and transcribed back into RNA for the subsequent round as described above. For rounds 9 and 10, counterselection was also performed on negative control immunoglobulin G (IgG; cetuximab, Merck Serono). Beginning with round 11, another cell-SELEX process was performed as described above. Cells for counterselection and positive selection and SELEX conditions are described in Figure 1 and Table S1. After the 14th round of selection, competitor yeast tRNA was added (Table S1). At the end of SELEX, the sequences of the 18th pools were cloned with the pGEX-4T vector and sequenced. The sequences were compared by MultAlin. Prediction of secondary structure was obtained using the mfold software.

Flow Cytometry
Flow cytometry was performed with individual aptamers directly coupled to Cy5 at their 3’ end. For comparison of the binding profile of different aptamers to cells, aptamers were used at a final concentration of 500 nM. For determination of equilibrium binding affinities, aptamer H02 was used at concentrations of 0.15, 0.3, 0.6, 1.25, 2.5, and 5 μM. After washing with EDTA (0.2 M), 300,000 cells were incubated for 30 min at 4°C with Cy5-labeled aptamers. As a control, cells were incubated with a 1:100 dilution of an anti-z5 antibody (mouse anti-human CD49e, IIA1 antibody, BD Chemigen) for 30 min, followed by 30-min incubation with Alexa 647-conjugated affine pure goat anti-mouse IgG (Jackson ImmunoResearch) at 10 μg/mL. After washing, cells were analyzed using a FACScalibur flow cytometer (Becton Dickinson), and the mean fluorescence intensity (counting 10 000 events) was measured using Flowing software 2.5.1. For KD determination, experiments were repeated three times, and data were evaluated using GraphPad Prism (version 5.04).

SPR Analyses of Aptamer H02-Integrin Interactions on a CAP Sensor Chip
All experiments were performed on a Biacore T200 instrument (GE Healthcare) at 25°C. The sensor surface and other Biacore consumables were purchased from GE Healthcare. Integrins z5β1 and z9β3 were from R&D Systems. Running buffer was composed of PBS (10 mM), NaCl (150 mM), and MgCl2 (1 mM), filtered through a 0.22-μm membrane, and supplemented with surfactant P20 (0.005% w/v). The biotin CAPture reagent, composed of streptavidin conjugated with an oligonucleotide, was stably hybridized to the complementary sequence of a CAP sensor chip following the biotin CAPture kit instructions (GE Healthcare). The biotinylated aptamer was denatured at 95°C for 3 min, incubated on ice, and then injected onto the biotin CAP reagent at 100 nM for 5 min at a flow rate of...
10 μL/min. Five different concentrations of integrin (ranging from 8–130 nM) were injected into the flow cells at 10 or 30 μL/min for 300 s. Dissociation followed for 300 s. After each measurement, the sensor chip was washed with an injection of 6 M guanidine hydrochloride in 0.25 M NaOH as recommended by the manufacturer. The reference surface was treated similarly except that aptamer injection was omitted. Binding curves were double-reference-subtracted from the buffer blank and reference flow cell (without the aptamer). The equilibrium response was recorded 5 s before the end of integrin injection. The K_D was determined by fitting the equilibrium response versus the [integrin] curve to a simple 1:1 interaction model with the Biacore T200 evaluation software (GE Healthcare).

**Fluorescence-Based Assays on Cell Lines**
The adherent CHO-B2 and GBM cell lines (U87MG α5+, U87MG α5−, LN319, LN229, LN443, SF763, LN18, LNZ308, U373, and T98G) were plated on sterile glass slides for one night at 37°C in culture medium, washed three times, and then saturated for 1 h at RT in selection buffer containing 2% BSA. Cy5-labeled aptamers were denatured at 95°C for 3 min and incubated on ice for 5 min and then on cells in selection buffer for 30 min on ice or at 37°C at concentrations dependent on the assay (5, 2.5, 1.25, 0.6, or 0.3 μM). Cells were then washed in selection buffer, fixed for 8 min in 4% paraformaldehyde (PFA), permeabilized for 2 min with 0.2% Triton, and washed again. Sequentially, when immunocytochemistry was performed, the primary antibodies used were the anti-α5 antibody (mouse anti-human CD49e, IIA1 antibody, BD Chemigen, 1/200), the anti-β1 antibody (mouse anti-human CD29 antibody, clone TS2/16, BioLegend, 1/500), or the anti-EEA1 (early endosome antigen 1) antibody (anti-mouse clone 14/EEA1, BD Transduction Laboratories, 1/1,000). Primary antibodies were added for 1 h at RT or overnight (O/N) at 4°C, followed by a secondary antibody coupled to Alexa Fluor 647 (A21245, Life Technologies). Horseradish peroxidase (HRP)-coupled secondary antibodies were used as housekeeping protein to serve as the loading control for cell lysate samples. Analyses were performed on three independent experiments.

**Confocal Imaging**
Images were acquired using a confocal microscope (Leica TCS SPE II, 63× magnification, oil immersion).

Mean fluorescence intensity on cells and tissues was measured using ImageJ software. Statistical analysis of data was performed with Student’s t test. Data were analyzed with GraphPad Prism version 5.04 and are represented as mean ± SEM.

**SUPPLEMENTAL INFORMATION**
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**AUTHOR CONTRIBUTIONS**

**Fluorescence-Based Histochemical Assays of Patient-Derived Xenografts**
Two patient-derived heterotopic xenografts (PDXs) were selected for *in vivo* analysis.68 TC7 and TC22 GBM-PDXs presented high and low levels of α5 integrins, respectively. PDX mouse models were established using tissues surgically removed from patients as described previously.69 Integrin α5 was apta- and immunostained using formalin-fixed paraffin-embedded xenografts mounted on glass slides. Sections were deparaffinized, rehydrated, and subjected to an antigen unmasking protocol. Briefly, sections were boiled at 100°C for 10 min in target retrieval solution (pH 9) (S2367, Dako), cooled down to RT for 20–40 min, and rinsed in H2O. For aptafluorescence, slides were rinsed for 5 min in selection buffer, dried, incubated in blocking buffer (2% BSA in selection buffer) for 1 h at RT, rinsed in H2O and then in selection buffer, and dried. RNA molecules were denatured at 95°C for 3 min and incubated on ice for 5 min before dilution in selection buffer to a 1 μM final concentration. Aptamers were incubated on tumor sections for 1 h at RT in a humid chamber, washed in selection buffer, dried, fixed in 4% PFA, and then washed three times in PBS. For immunofluorescence, slides were rinsed for 5 min in PBS-T (0.1% Tween 20 in PBS), dried, and then incubated in blocking buffer BB-1 (5% goat serum in PBS-T) for 1 h at RT in a humid chamber. Overnight incubation with anti-integrin α5 mAb 1928 (6B8516, Millipore, 1/200) in BB-1 was followed by a 5-min wash in PBS-T and by an incubation step with a 1/100 dilution of the goat anti-rabbit secondary antibody coupled to Alexa Fluor 647 (A21245, Life Technologies). Immun- and apta staining was followed by DAPI (10 μg/mL) staining for 30 min at RT to visualize cell nuclei. The stained samples were then washed in PBS, and coverslips were mounted onto tissue sections using fluorescent mounting medium (S3023, Dako).

**Western Blot**
Cells were lysed (1% Triton X-100, NaF [100 mmol/L], NaPPI [10 mmol/L], and Na3VO4 [1 mmol/L] in PBS, supplemented with complete anti-protease cocktail; Roche), and 10 μg of protein was separated by SDS-PAGE (Bio-Rad) and transferred to polyvinylidene fluoride (PVDF) membranes (Amersham). Blots were probed with antibodies to α5 integrin (H104, Santa Cruz Biotechnology) and to glyceraldehyde 3-phosphate dehydrogenase (GAPDH; Millipore). Horseradish peroxidase (HRP)-coupled secondary antibodies were from Promega. Proteins were visualized with enhanced chemiluminescence using the LAS4000 imager, and densitometry analysis was performed using the ImageJ software (GE Healthcare). GAPDH was used as housekeeping protein to serve as the loading control for cell lysate samples. Analyses were performed on three independent experiments.

**AUTHOR CONTRIBUTIONS**
CONFLICTS OF INTEREST
The authors declare no competing interests.

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