Structure based innovative approach to analyze aptaprobe–GPC3 complexes in hepatocellular carcinoma

Woo-Ri Shin¹†, Dae-Young Park¹†, Ji Hun Kim²†, Jin-Pyo Lee¹, Nguyen Quang Thai¹, Jin Hwa Ob¹, Simranjeet Singh Sekhon³, Wooil Choi³, Sung Yeon Kim⁴, Byung-Kwan Cho², Sun Chang Kim², Jiho Min³*, Ji-Young Ahn¹*** and Yang-Hoon Kim¹**

Abstract

Background: Glypican-3 (GPC3), a membrane-bound heparan sulfate proteoglycan, is a biomarker of hepatocellular carcinoma (HCC) progression. Aptamers specifically binding to target biomolecules have recently emerged as clinical disease diagnosis targets. Here, we describe 3D structure-based aptaprobe platforms for detecting GPC3, such as aptablotting, aptaprobe-based sandwich assay (ALISA), and aptaprobe-based imaging analysis.

Results: For preparing the aptaprobe–GPC3 platforms, we obtained 12 high affinity aptamer candidates (GPC3_1 to GPC3_12) that specifically bind to target GPC3 molecules. Structure-based molecular interactions identified distinct aptatopic residues responsible for binding to the paratopic nucleotide sequences (nt-paratope) of GPC3 aptaprodies. Sandwichable and overlapped aptaprodies were selected through structural analysis. The aptaprobe specificity for using in HCC diagnostics were verified through aptablotting and ALISA. Moreover, aptaprobe-based imaging showed that the binding property of GPC3_3 and their GPC3 specificity were maintained in HCC xenograft models, which may indicate a new HCC imaging diagnosis.

Conclusion: Aptaprobe has the potential to be used as an affinity reagent to detect the target in vivo and in vitro diagnosing system.

Keywords: Glypican-3, Hepatocellular carcinoma, Cancer diagnosis, Aptaprobe, Structure-based molecular interaction, Aptablotting, ALISA, Aptaprobe-based imaging

*Correspondence: jihomin@jbnu.ac.kr; jyahnt@chungbuk.ac.kr; kyh@chungbuk.ac.kr
†Woo-Ri Shin, Dae-Young Park, and Ji Hun Kim contributed equally
1 School of Biological Sciences, Chungbuk National University, 1 Chungdae-Ro, Seowon-Gu, Cheongju 28644, Republic of Korea
3 Graduate School of Semiconductor and Chemical Engineering, Jeonbuk National University, Jeonju 54896, Republic of Korea

Full list of author information is available at the end of the article
Introduction
Liver cancer is the fifth most common cancer and the second leading cause of cancer-related deaths worldwide [1]. Hepatocellular carcinoma (HCC) is the primary type of liver cancer, accounting for 85–90% of cancers formed by liver cells [2]. Hepatitis B virus, hepatitis C virus, and alcoholic liver disease are responsible for most HCC cases [3, 4]. Although current diagnostic tests and treatments for all cancer types have seen significant developments, the fatality rate for HCC remains very high. The mortality caused by HCC is increasing because early HCC stages are usually asymptomatic, yet when it causes symptoms, it is already at an advanced stage, causing difficulties in treatment [5]. As researchers learn about the mechanisms of HCC, new diagnostic tools are constantly being developed. Biomarkers and imaging studies are commonly used to diagnose HCC. Numerous serum biomarkers are used for diagnosing HCC, including α-fetoprotein (AFP), lipocalin 2 (LCN2), osteopontin (OPN), des-γ-carboxy prothrombin (DCP), and a fucosylated AFP glycoprotein variant (AFP-L3) [6–8]. In particular, AFP is a popular biomarker that has been extensively studied for HCC detection and diagnosis. With a 20 ng/mL cutoff value in serum, the observed serum AFP sensitivity for diagnosing HCC ranges from 60 to 80% [9, 10]. However, it dropped to 40% when it was used for detecting small tumors (< 3 cm). On the contrary, serum AFP level (20–200 ng/mL) also increased in a significant number of patients with chronic liver diseases, including 15–58% patients with chronic hepatitis [11] and 11–47% patients with cirrhosis [12]. In fact, it is not uncommon to find high AFP levels in patients with cirrhosis or HCC and cirrhosis overlap, which causes confusion in assays using AFP biomarkers [7, 13]. Thus, it is essential to identify a biochemical marker with better sensitivity and/or specificity, than AFP for distinguishing HCC from benign liver lesions.

GPC3 is one of six mammalian members of the glypican family (GPC1–GPC6) with heparan sulfate proteoglycans linked to the cell surface through a glycosyl-phosphatidylinositol (GPI) anchor [14, 15]. GPC3 is essential in cell growth, motility, and differentiation, and plays an important role in regulating cancer cell proliferation through interactions with other growth factors [16, 17]. GPC3 protein is highly expressed in HCC, but not in cholangiocellular carcinoma, gallbladder cancer, or benign liver tissues. Thus, it may be a novel serological marker for the early detection of primary liver cancer, particularly HCC [18–20]. Using a human GPC3 protein-specific polyclonal antibody, it has been shown that GPC3 is expressed in up to 70% HCC tumors, but not in normal liver tissues [21]. GPC3 has also been found to be overexpressed in HCC cell lines, including HuH6, HepG2, PLC/PRF/5, Hep3B, HuH7, and HT17 [22]. A recent study showed that high GPC3 expression is not dependent on tumor size. In addition, it has a 56% sensitivity for patients with small early stage tumors [23]. A meta-analysis evaluated GPC3 use in early stage HCC detection (TNM stage I or BCLC 0 and A) and observed the specificity and sensitivity to be 97.0 and 55.1%, respectively [24]. For comparison, the specificity and sensitivity of AFP in similar studies were about 87.6 and 34.7%, respectively [25, 26].

Molecular probes are powerful tools for identifying disease biomarkers and tracking them in their native environments. SELEX technology has been developed as an
in vitro selection method to screen aptamers, which are nucleic acid ligands with prominent functionalities [27]. Aptamers are short single-stranded nucleic acid oligomers (ssDNA, RNA, or amino acids) that can fold into or around a target [28–33]. In addition, aptamers can bind to their targets (nanomolar to picomolar range) with high affinity and specificity and display low to no immunogenicity. They are easier to modify because they are chemically synthesized. Recently, numerous benefits of aptamers as antibody alternatives have been intensively studied for a wide range of applications, such as diagnosis and therapeutic agents like aptaprobe. Aptaprobe is a functionalized-enhanced aptamer for target detection, used as an oligo-recognition molecule in diagnostic platforms for wide array of clinical samples, including blood, urine, and saliva. Reportedly, the aptamer binds to target molecules through its specific shape by a 3-dimensional (3D) structure of the sequence [34–36]. Understanding the interaction between the aptaprobe and its target molecules is a key step in improving aptaprobe-based diagnostic platform technology [29]. In this study, we performed 3D structure-based molecular docking simulations, which were applied to predict the 3D aptaprobe binding structure, followed by validation using in vitro GPC3 detection. Our structure-based simulation study has the following three standpoints: (i) it can provide problem-solving strategies for understanding the 3D structure of aptaprobos, (ii) aptaprobe-binding mechanism on the biological key sites can provide specific local features, including the improvement of aptaprobe-binding strengths and weaknesses without actual tests, and (iii) it provides virtual contact map prediction of intrachain nucleotide-amino acid residues for elucidating the detailed features and roles of the identified contact moieties.

The study includes three major stages: (i) aptamer screening explains the isolation technique of aptamer candidates, (ii) the aptaprobe and aptaprobe–GPC3 complex structural interaction, and (iii) technical platform validations; aptablotting, aptaprobe-based sandwich assay (ALISA), and aptaprobe-based imaging analysis. The first stage has two components: in vitro SELEX and surface plasmon resonance (SPR) affinity analysis. The second stage describes in detail the 3D structural analysis of molecular interactions to provide specific aptaprobos. Next, the platform validation stage describes the experimental results showing the applicability of the selected aptaprobos, and concludes the study by summarizing the results and contributions of HCC-targeted specific diagnosis. The overall flow of this study is shown in Scheme 1.
Materials and methods

Protein and buffer solutions
The aptamers used in all the experiments were synthesized by IDT (Coralville Johnson County, IA, USA). Recombinant human glypicans proteins (GPC3 and GPC5) were purchased from R&D Systems (Minneapolis, MN, USA), and anti-human GPC3 monoclonal human antibody was purchased from G&K Bioscience (Santa Clara, CA, USA). HSA, BSA, human serum, and anti-mouse IgG peroxidase were purchased from Sigma (St. Louis, MO, USA). Horseradish peroxidase (HRP)-conjugated streptavidin was purchased from Thermo Fisher Scientific (Minneapolis, MN, USA). The following buffers were used: TBS-T buffer (0.1% (v/v) Tween 20 in Tris-buffered saline, pH 7.2), blocking buffer (5% non-fat dried milk in TBS-T), and washing buffer (TBS-T buffer).

Molecular docking structural analysis

GPC3 aptamers were identified through a SELEX process; details of the selection can be found in Additional file 1. The secondary structure of GPC3_1–GPC3_12 was obtained using the Mfold web server (http://mfold.alfany.edu/?q=mfold) at 25 °C in a 250 mM [Na+] folding algorithm based on nucleotide sequences. A 3D RNA model was constructed from the secondary structure predicted using the Mfold program [37]. The ssDNA structures of aptamers were built by converting RNA to DNA using Pymol software [38]. The amino acid sequences of GPC3 was obtained from UniProtKB (PS1654.1), which was used to predict GPC3 protein structure using SWISS-MODEL (https://swissmodel.expasy.org/). The docking prediction of the GPC3 protein–aptamer complexes was analyzed using the MOE2016.08 program [39]. All GPC3 protein and aptamer 3D models were minimized using an energy minimization algorithm (gradient; 1.0, Force Field: Amber10: EHT). The Triangle Matcher method was used to analyze the GPC3 protein and aptamer binding region. The results of the interaction distance between the GPC3 protein and aptamers were generated using Pymol, COCOMAPS/CONSRank [40], and LigPlot+ [41]. Finally, the GPC3–aptamer complex (GPC3_2, GPC3_3, and GPC3_9), named “aptaprobe,” was structurally analyzed.

Cell culture
All cell lines, including HepG2 (ATCC HB-8065), Hep3B (ATCC HB-8064), PANC-1 (ATCC CRL-1469), MIA-PaCa-2 (ATCC CRM-CRL-1420), and HeLa (ATCC CCL-2) were purchased from the American Type Culture Collection. All cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Welgene, Gyeongsan, Korea) supplemented with 10% (v/v) fetal bovine serum (FBS; Gibco, Minneapolis, MN, USA) and incubated at 37 °C with 5% CO2.

mRNA expression test

HCC cell lines [HepG2 (ATCC HB-8065), Hep3B (ATCC HB-8064)], pancreatic cancer cell lines [PANC-1 (ATCC CRL-1469), MIA-PaCa-2 (ATCC CRM-CRL-1420)], and HeLa cell line (ATCC CCL-2) were cultured on 100Φ dishes. RNA was extracted using the TRIzol™ Plus RNA Purification Kit (Thermo Fisher Scientific). The extracted RNA was converted to cDNA using a TOP-script™ cDNA Synthesis Kit (Enzymics, Deajeon, Korea). GAPDH and GPC3 genes were PCR-amplified using gene-specific primers [GAPDH (Forward: 5′-GCT TCA GTG CAC CAG GAA-3′, reverse: 5′-TTA GTT CGA CAG TCA GCC-3′), GPC3 (Forward: 5′-TAA TTATGG GCGGACCGTG-3′, reverse: 5′-TCGAGT TCGTG CACCAAG-3′)]. PCR was performed for 25 cycles. The results were confirmed using 2% agarose gel electrophoresis.

Cell viability and mitochondrial respiratory competency assay

Cell viability and mitochondrial respiratory competency were determined by MTT assay and Seahorse XF analyzer mito stress test assay (Agilent Technologies, Santa Clara, CA, USA). For MTT assays, HepG2 cells were seeded in 48-well plates and incubated overnight for cell adhesion. They were then incubated for 24 h with 10–0.125 μM GPC3_3 aptaprobe (two-fold dilutions) and wells containing untreated cells were used as controls. At the end of incubation, the media was replaced with fresh media containing MTT and further incubated at 37 °C for 4 h. Finally, the formazan crystals were dissolved in dimethyl sulfoxide (DMSO) and the absorbance at 570 nm was recorded using a SPECTRAMAX M2e ELISA reader (Molecular Devices, San Jose, CA, USA). The Seahorse XF analyzer mito stress test assay was performed on HepG2 cells (1.8 × 10⁴ cells/well) treated for 1 h with various GPC3_3 aptaprobe concentrations. The OCR and ECAR were measured before and after treating the cells with mitochondrial toxins oligomycin (ATP synthase inhibitor, 1.5 μM) and carbonyl cyanide-4-phenyl-hydrazone (FCCP, uncoupling agent, 0.5 μM), and rotenone/antimycin A (Rot/AA, uncoupling agent, 0.5 μM) to determine the baseline and stress measures of oxidative phosphorylation and glycolysis, respectively. The OCR and ECAR values were normalized to the total protein concentration. All experiments relative to the control were expressed as the mean±standard error of mean (SEM) from at least triplicate studies.
Confocal microscopy image and flow cytometry

The binding of aptamers to HeLa, HepG2, Hep3B, PANC-1, and MIA-PaCa cell lines (1 x 10^6 cells/mL) was monitored using confocal microscopy. After washing twice with cold washing buffer, the cells were incubated with 200 pM of FAM-labeled GPC3 aptapropbes (GPC3_3, GPC3_2, and GPC3_9) at 37 °C for 1 h. Subsequently, the cells were washed twice, imaged using a Zeiss LSM 710 upright confocal microscope (Oberkochen, Germany), and analyzed using a BD FACS Calibur™ Flow Cytometer (Franklin Lakes, NJ, USA).

HCC xenograft model generation

HCC xenograft models were created using HepG2 cells. Briefly, 1 x 10^7 cells were injected subcutaneously into the right flank of 6-week-old BALB/c nude mice (Orienvbio, Seongnam, Korea). The tumor was allowed to grow till 100-200 mm^3 (measured with a Vernier caliper). The tumor volume was calculated using the formula: V(volume) = L(length) x W(width) x H(height)/2. The mice were monitored weekly for tumor growth and GPC3_3 aptaprobe in vivo imaging system (IVIS) imaging as described below.

Aptaprobe-based blotting assay: aptablotting

The analysis was performed using a previously described protocol with some modifications (Fig. 2A) [29]. GPC3 aptapropbes, GPC3_2, GPC3_3, and GPC3_9 were synthesized with 5' biotinylation by IDT. GPC3, HSA, and BSA (5 μg) were loaded into each well and 5 μg GPC3 was spiked into human serum. All proteins were separated using 10% SDS-PAGE and transferred to a nitrocellulose (NC) membrane (GE Healthcare). The transferred proteins were stained with Ponceau S for 1 min. The stained membrane was destained with water, then air-dried. The membrane was blocked for 1 h and washed thrice with TBS-T. GPC3 aptamers were used to detect only the GPC3 protein. The blocked membranes were incubated with 1 μg/mL GPC3 aptamer in blocking buffer (5% milk, 0.1% BSA in TBS-T) for 3 h at 4 °C and washed thrice with TBS-T. The GPC3 aptamers were detected using 1:2000 diluted HRP-conjugated streptavidin (Thermo Fisher Scientific) in blocking buffer for 1 h at 4 °C. The HRP signal was developed using an enhanced chemiluminescence (ECL) system. In aptaprobe-dot blotting, the serially diluted proteins (100, 50, 10, 1, and 0 pmol) were immobilized on NC membranes using the method mentioned above.

Aptaprobe-based quantitative detection of GPC3: aptaprobe-linked immobilized sorbent assay (ALISA)

ALISA was performed in streptavidin-coated 96-well plates for quantitative GPC3 protein detection using GPC3 aptamers. Biotinylated GPC3_2 and GPC3_9 aptapropbes (1 μM/well) were used to immobilize a 96-well plate for 1 h at 25 °C. The plate was washed thrice and non-specific binding was blocked with TBS-T containing 2% BSA. The same concentration (500 nM/100 μL) of GPC3, GPC5, BSA, and HSA was then added to each well and incubated for 1 h at 25 °C with gentle shaking. Then, 5, 10, 25, 50, 100, 250, and 500 ng/100 μL GPC3 protein were added to each well. The unbound proteins were discarded and washed thrice with a washing buffer. To quantify the signaling target protein, the FAM-modified GPC3_3 (1 μM/well) was added to individual wells for 30 min at RT. Following the reaction, the unbound and non-specific bound FAM-modified GPC3_3 was removed and the plate was washed with washing buffer. The fluorescence intensity of each well was then measured at 495 nm excitation wavelength and 520 nm emission wavelength using a SPECTRAMAX M2e ELISA reader.

Aptaprobe-based fluorescence imaging in vitro targeting

HepG2 cells (1.5 x 10^5 cells/well) were seeded in 6-well plates onto coverslips (Marienfeld Superior). After incubating for 24 h, the cells were incubated with fresh media containing 200 pM FAM-modified GPC3_2, GPC3_3, and GPC3_9 separately for 4 h at 37 °C in a 5% CO2 incubator. The cells were washed three times with DPBS to remove the unbound aptamers, fixed with 4% paraformaldehyde, and stained with 4',6-diamidino-2-phenylindole (DAPI). Coverslips with fixed cells were mounted on glass slides with a mounting solution. Fluorescence images were obtained using a confocal laser scanning microscope (Zeiss, Oberkochen, Germany).

HCC-positive and HCC-negative cell imaging with GPC3_3

HCC (HepG2 and Hep3B), pancreatic cancer (PANC-1 and MIA-PaCa-2), and HeLa cell lines (3.0 x 10^4 cells/well) were cultured in 24 wells onto coverslips. After incubation for 24 h, the cells were incubated with fresh media containing 0–200 pM FAM-modified GPC3_3 for 4 h at 37 °C in a 5% CO2 incubator. Confocal imaging was performed as described above.

In vivo fluorescence imaging of the xenograft tumor

HCC xenograft model mice were imaged in vivo to minimize the expression of fluorescence in the body by fasting for 12 h. The mice were randomly assigned to two groups (n = 3 per group): Group I, intravenously injected with 2 mg/kg Control (Cy5.5-labeled random sequence); Group II, intravenously injected with 2 mg/kg GPC3_3Cy5.5 (Cy5.5-labeled GPC3_3). The Cy5.5-labeled
samples were detected 30 min after injection by placing the mice in the supine position using the IVIS Lumina system III (Perkin Elmer, Waltham, MA, USA). The mice were euthanized 1 h after injection, laparotomy was performed, and in situ imaged using Cy5.5 filters, which were normalized to the background signal per manufacturer’s protocol. The liver, lung, spleen, kidney, heart, tumor, small intestine, and large intestine tissue of mice were harvested.

**Statistical analysis**
All statistical analyses were performed using GraphPad Prism or R statistical software [42]. Two-tailed unpaired Student’s t-test with Welch correction or one-way analysis of variance were used for multiple comparisons. Statistical significance was defined as $P < 0.05$ in all comparisons and calculated as described in the manuscript or figure legends.

**Results and discussion**

**Highly specific and efficient GPC3-binding DNA aptamers**
Based on our aptamer screening procedure, specific GPC3-binding aptamers were carefully isolated in this study. Single-strand DNA aptamer candidates against the recombinant GPC3 protein were generated using the modified SELEX protocol [43]. The details of this protocol are available in Additional file 1. The eluted ssDNA concentration in each selection round was monitored using a NanoDrop spectrophotometer, which reflected the binding capacity between the selected ssDNA pool and the GPC3 target. ssDNA concentrations in each round increased until the sixth round of selection (Additional file 1: Fig. S1A). Changes in ssDNA concentration indicate that the ssDNA pool was enriched after each selection round. After a negative round, ssDNA aptamer concentrations of the eighth, ninth, and tenth selection rounds were 488.4, 491.9, and 405.9 ng/μL, respectively. At the tenth round, the ssDNA aptamer concentration decreased, it means the binding efficiency of ssDNA for the GPC3 was decreased. To provide the binding efficiency of ssDNA for the GPC3, we performed the Post-SELEX and Real-time PCR (Additional file 1: Fig. S1B). The samples from ninth round SELEX showed a lower $C(t)$, $C(t)$ = 58.83, as compared to other samples, $C(t)$ = 88.35 for eighth round, 19.51 for tenth round and 19.75 for seventh round, indicating the presence of higher amounts of ssDNA aptamer for GPC3. It is suggesting that the ninth round was the optimal state. The GPC3 bound ssDNA pool was amplified, cloned, and sequenced. A total of 76 clones were selected for DNA sequencing and 12 different DNA sequences were identified (Table 1).

To determine the characteristics of GPC3-binding ssDNA aptamer candidates, an SPR assay was used to analyze the specific binding of 12 aptamers with different sequences to GPC3 and determine the binding affinity ($K_D$ values) of the ssDNA aptamer (Additional file 1). $K_D$ values of ssDNA aptamer candidates were measured using BIA evaluation software (GE Healthcare). The results indicated that the $K_D$ values of these ssDNA aptamer candidates could be classified into three distinct groups. Of these ssDNA aptamer candidates, GPC3_1–GPC3_12 had low $K_D$ values (Table 1 and Additional file 1: Fig. S2). The $K_D$ values of ssDNA aptamers ranged from $1.2 \times 10^{-10}$ to $1.06 \times 10^{-8}$ M. In addition, the results obtained from SPR confirmed that GPC3_3 had the highest affinity for GPC3 based on $K_D$ values ($1.2 \times 10^{-10}$ M), which would aid the detection of GPC3 targets.

**Table 1** Sequences and affinity value of GPC3 aptamers

| Clone  | Selected sequence | $ka$ (1/Ms) | $kd$ (1/s) | $K_D$ (M) |
|--------|-------------------|-------------|------------|-----------|
| GPC3_1 | CATGATCAATCCGGTACATATTTATCCTCAATTCACA | $2.09 \times 10^4$ | $6.45 \times 10^{-5}$ | $3.09 \times 10^{-9}$ |
| GPC3_2 | AATCCATAGTTCTGTGTTTCCAGTTTCTTTATGTCG | $2.28 \times 10^5$ | $9.84 \times 10^{-4}$ | $4.31 \times 10^{-9}$ |
| GPC3_3 | CCTATCCCTATTATTTTCTTTTGGATTTGTCG | $3.67 \times 10^5$ | $4.40 \times 10^{-5}$ | $1.2 \times 10^{-10}$ |
| GPC3_4 | CCATAAATCTTTTTCTCGTCATTTAGTTATCCCTACG | $1.11 \times 10^6$ | $7.14 \times 10^{-5}$ | $6.46 \times 10^{-9}$ |
| GPC3_5 | GCCTAATATATGTATTTCCATTACTCGTTTTTTACATT | $2.20 \times 10^5$ | $8.53 \times 10^{-4}$ | $3.88 \times 10^{-9}$ |
| GPC3_6 | TAGTTTTTTAGGGTTTTTGTGGTTGCTATGTGGTGG | $2.00 \times 10^5$ | $1.35 \times 10^{-5}$ | $6.71 \times 10^{-10}$ |
| GPC3_7 | ACTTTGTAGTGTGTATTGATTGTCTTTTCTTCCTCAG | $5.94 \times 10^4$ | $6.30 \times 10^{-5}$ | $1.06 \times 10^{-9}$ |
| GPC3_8 | ATACACATTATCTGTCATTTTTATATCCCTACG | $2.75 \times 10^5$ | $5.00 \times 10^{-5}$ | $1.82 \times 10^{-9}$ |
| GPC3_9 | ACCTCTTTTTTTCCTCGTATGTTTTTCTTCTCG | $2.5 \times 10^5$ | $5.17 \times 10^{-5}$ | $2.07 \times 10^{-10}$ |
| GPC3_10 | TTTCTGTGATAATTCTTCTATTTCATTCTACG | $1.72 \times 10^5$ | $6.12 \times 10^{-5}$ | $3.57 \times 10^{-9}$ |
| GPC3_11 | TTTTCTTCGTTATATCTTTCTATTCTAGTGG | $1.80 \times 10^6$ | $9.80 \times 10^{-4}$ | $5.43 \times 10^{-10}$ |
| GPC3_12 | CTTTTCTGTGTTTTTATAGCTTCTTCCAGTTTTCTGGAAC | $1.09 \times 10^6$ | $8.91 \times 10^{-4}$ | $8.16 \times 10^{-10}$ |

Affinity value derived $ka$, $kd$, and $K_D$ values of 12 selected GPC3 aptamers by measuring the surface plasmon resonance (SPR; Biacore ×100)
3D structure analysis of the molecular interaction between DNA aptamer and GPC3 protein

The GPC3 protein is a member of the glypican family, which is attached to the cell surface and has a glycosylphosphatidylinositol (GPI) anchor region [44]. The protein comprises two subunits, the ~40 kDa α-subunit (light blue) and the ~30 kDa β-subunit (deep blue), which is cleaved between Arg358 and Ser359 [45] (Additional file 1: Fig. S3A and S3B). The GPC3 protein structure was predicted using the Glycan-1 structure model, 4YWT, in SWISS-MODEL by piecewise homology (Additional file 1: Fig. S3C). The GPC3 protein structure was used to determine the structural interactions with the DNA aptamer. The important GPC3 protein residues are shown in Additional file 1: Fig. S3B, C. The predicted GPC3 model consists of 16 α-helices and three major lobes [46].

These aptamers have complex secondary structures, including protruding loops and stems. The interaction conformation between the GPC3 protein and each aptamer candidate is shown in Additional file 1: Fig. S4A. The GPC3 protein was fixed in the same position for analysis, and each aptamer binding site of the GPC3 protein is shown in a different color. We also analyzed the detailed GPC3 protein–aptamer binding interaction using COCOMAPS/CONSRank and LigPlot, and the amino acid sequence of the GPC3 protein is shown in Additional file 1: Table S1. We further analyzed the distance of contact sequence between all GPC3 and 12 aptamer candidates, and indicated them using the heatmap; dark purple color indicates >4 Å distance, while the bright color indicates <4 Å distance or closer [47] (Additional file 1: Fig. S4B). All aptamer candidates showed a strong bright color in distinct areas of the GPC3 protein sequences, showing that the interaction between the GPC3 amino acid residues and DNA aptamers is in the proximation position. We analyzed the heatmap to measure the binding distance between the GPC3 amino acid residues. Heatmaps representing intermolecular contacts for individual residues indicated that frequent contact of the GPC3/GPC3_3 aptamer (green) interaction was comparable to that of the GPC3/GPC3_9 aptamer (magenta) interaction. However, the interaction mode of the GPC3/GPC3_2 aptamer (light purple) was completely different from those of the GPC3/GPC3_3 and GPC3/GPC3_9 aptamers (Fig. 1A). To characterize the GPC3–DNA aptamer interaction comprehensively, visual contact performances using the COCOMAPS web server were obtained (Fig. 1B). Despite the nucleotide sequence differences, two DNA aptamers (GPC3_3 and GPC3_9) bind to GPC3 with similarly. In addition, we studied the relative contribution of hydrogen bonds and hydrophobic interactions of GPC3 at the DNA aptamer binding sites. All 12 DNA aptamer candidates were analyzed for hydrogen bonding at the GPC3 binding interface using LigPlot+ software (Additional file 1: Table S1). The average distance of the hydrogen bonds is shown in Fig. 1C. This result demonstrates that the GPC3_3 aptamer had the shortest distance, which may help stabilize the GPC3 protein–aptamer complex by mediating strong hydrophobic interactions. Hydrophobic interactions can influence binding affinity and molecular functional activity [48, 49]. These results correspond well with those described in the SPR affinity analysis (Table 1 and Additional file 1: Fig. S2). Based on the GPC3_3 structural interactions, which the binding region is comparable to GPC3_3, and GPC3_2, which is totally different, referred to “Aptaprobese.” We then attempted to determine the functional binding of GPC3 aptaprobese (GPC3_3, GPC3_2, and GPC3_9) and confirmed their structure-based functional differences. As shown in Additional file 1: Table S1, the results of the structural interaction of the aptaprobese binding with GPC3 protein involves 3D interaction between the binding site on GPC3 and the nucleotide paratopic sequences (nt-paratope), the binding site on the aptaprobese. The binding geometry of GPC3_3 aptaprobese indicated that five nt-paratopes (T3, C13, A25, A97, and C99) were responsible for binding to the GPC3 aptaprobese: the specific amino acid residues, Lys453, Lys467, and Val532, have a significant interaction with the 5′-end nt-paratope and Asp193 and Cys197 bind with the 3′-end nt-paratope (Fig. 1D). GPC3 consists of an N-terminal secretory signal peptide and a C-terminal core domain. The GPC3 core forms a complex with Wnts, activating downstream signal transduction and stimulating HCC proliferation [50]. GPC3_3 aptaprobese bind directly to the GPC3 core region, suggesting that the aptaprobe

Fig. 1 Interaction of the GPC3_3, GPC3_2, and GPC3_9 aptamers with GPC3 protein. A Heatmap and graph of the binding distance between GPC3 amino acids and GPC3_3, GPC3_2, and GPC3_9 aptamers. Dark purple color indicates >4 Å distance, while the bright color indicates <4 Å distance or closer. B GPC3 protein–aptamer contact map obtained by COCOMAPS. A significant contact region of GPC3 protein–DNA aptamers (GPC3_3, GPC3_2, and GPC3_9) are shown in light purple, green, and magenta, respectively. Six regions in overlapped area of aptamer (GPC3_3 and GPC3_9) have been highlighted through blue rectangles. C Average distance of hydrogen bond (O–H) length (Å) between GPC3 protein and each DNA aptamer candidate (GPC3_1–GPC3_12). Each value of 12 DNA aptamer candidates was calculated by COCOMAPS/CONSRank and LigPlot+ (see *Materials and methods*). The GPC3 protein is depicted as a ribbon structure with stick structure of D GPC3_3 (green), E GPC3_2 (light purple), and F GPC3_9 (magenta). Each color represents areas in GPC3 protein contacted by nt-paratopes.
Fig. 1 (See legend on previous page.)
may alter the GPC3–Wnt signaling pathway. Since GPC3 expression increases in HCC, aptamers can be suggested as potential therapeutics for HCC.

Figure 1E illustrates that the GPC3_2 aptaprobe binds to GPC3 at two binding sites different from GPC3_3 binding sites (Fig. 1D). The GPC3_2 aptaprobe comprises two physically connected paratopic moieties (six nt-paratopes of GPC3_3 aptaprobes: T72, T78, T80, A4, A87, and G88) that can simultaneously interact with two different aptatopes, His412, Asn127, Asn124, His121, and Gln106 (Additional file 1: Fig. S5A, C; Met1, Ala2, Leu192, Asp193, Ile194, Glu196, Cys197, Phe445, Pro455, Ser460, Glu490, Leu523, Tyr528, Leu530, Asp531, Asp533, and Asp534). Taken together, GPC3_2 and GPC3_3 were finally chosen for ALISA as capture and reporter, respectively. The results of this study highlight that the molecular docking can successfully predict the interaction between nt-paratopes and aptaprobes, which is promising for high-throughput aptamer screening. We present three GPC3 detection methods using: GPC3_3, GPC3_2, and GPC3_9 aptaprobes.

Aptablotting assay for the GPC3–aptaprobe interaction

The GPC3 protein is gradually expressed in the serum and tissues of HCC patients [51]. Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) is the most popular biomarker detection method, which identifies a specific biomarker protein in a complex matrix, such as tissue lysate, blood, or other clinical samples. This motivated us to use aptamers for GPC3 detection. Aptablotting has previously shown that the aptamer can help overcome the obstacles of current immunoblot assays [29]. In this study, we designed an aptablot assay for GPC3 detection. GPC3_3, GPC3_2, and GPC3_9 aptaprobes were chosen for establishing an aptablotting assay and chemically synthesized as biotin-labeled affinity aptamers (Fig. 1A). Subsequently, we tested whether the aptamer non-specifically react with various proteins, such as human serum proteins, human serum albumin (HSA), and bovine serum albumin (BSA). The purified recombinant protein GPC3 was used as a control (Fig. 1A). The GPC3 proteins and test samples were separated using 10% SDS-PAGE and then transferred onto a nitrocellulose (NC) membrane. We stained the membrane with Ponceau S that indicates 5 ug of each protein, GPC3, HSA, and BSA in approximately 50–70 kDa. Following a simple blocking step, the aptablotting assay required only a single aptaprobe affinity detection using each biotinylated aptaprobe. It is important to note that three GPC3 aptaprobes could recognize their target in both SDS-denatured environments and complex biological samples. In contrast, GPC3 aptaprobes did not react with the serum abundant proteins HSA or BSA.

The aptaprobes could recognize GPC3 in its native state. Figure 2C and Additional file 1: Fig. S6 show the specificity and sensitivity of the aptamer probe to GPC3 concentration in the dot blot assay. The GPC3 protein, a member of the glypican family protein, is not specific to antibodies and aptamers with specificity [52]. When GPC3 was included to represent the complex protein mixture (human serum: H. serum; bovine serum: B. serum), it was clearly demonstrated that the aptamer-bound GPC3 proteins selectively and the observed binding signals were not due to non-specific adsorption to either NC membrane or abundant serum proteins. However, anti-GPC3 antibody showed a strong signal in human serum without GPC3, which was generated by the reaction of human serum with second antibody for detection because the first GPC3 antibody was sourced from humans. Finally, all aptaprobes, GPC3_3, GPC3_2, and GPC3_9, exhibited similar binding behaviors toward native GPC3 proteins. In particular, GPC3_3 reacted with 1 pmol GPC3 in both H. serum and B. serum.
Realizing that the isolated aptaprobes tended to recognize their target structure well in SELEX conditions where they were originally isolated or not, the aptablotting performances in SDS-denatured environment were far from their appropriate condition. Some proteins are known to be resistant to SDS-induced denaturation [53, 54]. It has been shown that aptamer binding was observed in the SDS-PAGE and blotting assays, suggesting that sufficient SDS amount and intrinsic structural rigidity contribute to SDS resistance [55, 56]. We concluded that the high level of GPC3 structural conservation from SDS denaturation may be used as the basis for the aptablotting assay to identify both SDS-denatured and native GPC3 proteins.

The aptablotting assay can directly detect GPC3, whereas western blotting detection by antibody-based experiments is generally stepwise with incubation and labeling, such as primary and secondary antibody use. In addition, negatively charged nucleic acid aptamers bind poorly to the PVDF/NC membrane, indicating that the aptaprobe-involved aptablotting assay can be improved and contribute to their ability to detect and image targets with various signal read-out fluorescent or colorimetric tags. It is expected that the immunoblotting with primary and secondary antibodies might lead to several non-specific signals in addition to the GPC3 signals (Additional file 1: Fig. S6C, see red arrow). This means that the GPC3 aptaprobe should not cross-react with non-target molecules in the sample mixture, and our aptablotting protocol simplifies the immunoblotting workflow. To be an effective cellular target binding reagent, an aptaprobe should ideally be achieved following binding to its aptatopes expressed on the cell surface of various cell lines.

Quantitative-ALISA platform for detecting GPC3

Interest in the use of aptamers in ALISA is on the rise [8, 31, 38]. ALISA relies on the formation of a sandwich configuration between two layers of the aptaprobe pair (capture and reporter probes), which binds to the target to create a sandwich complex. This indicates that aptaprobos should bind to distinct aptatopes, and their
target recognition should not be mutually exclusive. To identify the aptaprobe pair, individual aptatopes should be tested pairwise in a sandwich matrix so that each aptatope is occupied with other candidates, both as capture and reporter aptatopes. However, this conventional method is extremely time-consuming and labor-intensive to complete, and less time-consuming approaches are required. It should be noted here that molecular docking can be considered as a valuable strategy to provide high-quality and functional aptaprobes and contribute to understanding the binding configuration of aptaprobe-target ligands.

ALISA-quantitative detection was performed as shown in the sandwich assay platform, which is easy to use and does not require any coupling of stepwise secondary antibodies for signal readouts (Fig. 3). To confirm the use of aptaprobe pairs for detecting GPC3 in ALISA, three blanks (C1: microwell plate, C2: non-specific reporter aptaprobe binding, and C3: non-specific target binding without capture aptaprobe), and two negative controls (C4: biotinylated GPC3_9 and fluorescein amidite (FAM)-labeled GPC3_3, and C5: biotinylated GPC3_2 and FAM-labeled GPC3_3) were used. The experimental test groups were set as [Test1] to [Test4] (Fig. 3A).

The normal sandwich assay platform was subjected to quantitative analysis after treatment with 5, 10, 25, 50, 100, 250, and 500 ng in 100 μL GPC3 protein. Therefore, the fluorescence signal combined with the reporter aptaprobe was confirmed only in the GPC3_2–GPC3–GPC3_3 configuration condition [Test4]. This is because GPC3_9 and GPC3_3 have the same binding sites in GPC3, which interferes with GPC3 binding when they were used for the ALISA [Test3]. The blanks, control (C1–C5) and Test1–Test3 groups had no effect on signal response (Fig. 3A). Increasing GPC3 concentration increased the signal response, and ALISA demonstrated
identified < 1 ng/mL GPC3 in clinically relevant serum levels. The results of the ALISA followed the linear equation $Y = 616.92 \log_{10}(X) - 342.75$ ($R^2 = 0.9958$) (Fig. 3B) and corresponded significantly to the binding interaction results (Fig. 3C).

**Aptaprobe-based fluorescence imaging**

GPC3 has three parts: signal peptide for membrane translocation (SP), cysteine-rich domain (CRD), and glycosylphosphatidylinositol (GPI). The Notum cleavage site breaks the bond between amino acids 562 and 563 [50], causing proteins other than GPI to be secreted into the serum (Additional file 1: Fig. S3). FAM-labeled GPC3_3 and GPC3_2 aptaprobies successfully achieved targeted imaging in HepG2 cells, which are known to overexpress GPC3 protein on their cytoplasmic membrane (Fig. 4). The green fluorescence intensity was shown in Additional file 1: Fig. S7B to analysis aptaprobe binding. However, HepG2 cells were incubated with the FAM-labeled GPC3_9 and confocal imaging results demonstrated that the fluorescence signal was reduced (Additional file 1: Fig. S7). This may be due to the non-binding of the GPC3_9 aptaprobies that did not bind to their potential binding site (GPI site, Additional file 1: Table S1) by the cell membrane-bound region. However, FAM-labeled GPC3_3 and GPC3_2 successfully achieved targeted imaging in HepG2. FACS experiments supporting the results of confocal imaging, blocked GPC3_3 binding. This is consistent with the results of previous confocal imaging experiments (Fig. 4). As mentioned earlier, GPC3 is a heparan sulfate proteoglycan, and the relationship between protein glycosylation alteration and liver diseases has been reported [57]. Our GPC3_3 aptaprobies may be used to monitor glycosylation alterations during HCC progression.

The aptaprobe-binding capacity can be maintained within the actual cellular environment. We used two pancreatic cancer cells, PANC-1 and MIA-PaCa, as negative controls for aptamer binding, as these cells do not express GPC3. To confirm GPC3 expression, the mRNA expression levels were determined preferentially. GPC3 gene expression was observed in only two HCC cell lines, HepG2 and Hep3B, but not in the pancreatic cancer cells (Additional file 1: Fig. S8). To verify extracellular GPC3 targeting, GPC3-positive and -negative cell lines were incubated at 37 °C for 30 min with FAM-labeled GPC3_3

![Confocal imaging and flow cytometry with GPC3 aptaprobies.](image)

**Fig. 4** Confocal imaging and flow cytometry with GPC3 aptaprobies. Fluorescence image of HepG2 cells using 200 pM of fluorescein amidite (FAM)-labeled GPC3 aptaprobe (GPC3_3, GPC3_2, and GPC3_9). Signals were observed on the cytoplasmic membrane on HepG2 cells. Flow cytometry assays of the binding of individual FAM-labeled aptaprobies with HepG2 cells (target cells). Only HepG2 cell is shown in red and each aptaprobe-bound HepG2 is shown with colored green curves.
An intense fluorescent signal was observed in GPC3-positive cells, indicating that GPC3_3 specifically targeted the surface of the liver cancer cell membrane. FACS experiment using FAM-modified aptamers confirmed the result of confocal imaging. A shift ratio of <1% was observed in the GPC3-negative cell lines (HeLa, PANC-1, and MIA-PaCa), while a shift ratio of approximately 99% was observed in the GPC3 positive cell lines (HepG2 and Hep3B) (Fig. 5 and Additional file 1: Fig. S7C). These results indicate that the aptaprobe can distinguish the GPC-3 positive cancer cell lines from the GPC-3 negative-cell lines. Chemical modifications (Cy3 and FAM) are beneficial for applications in HCC imaging. In the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide salt (MTT) assay, HepG2 cells incubated with GPC3_3 retained their high survival properties, indicating the low cytotoxicity of GPC3_3 (Additional file 1: Fig. S9A). Corresponding to confocal imaging, the HepG2 cells did not show any nuclear or cytoplasmic morphological changes. Additionally, the intracellular metabolic potential and mitochondrial respiration of the aptaprobe were confirmed. Since at least 90% cellular energy is produced by mitochondria, we investigated whether the aptaprobe affects mitochondrial structure, which reflects a disruption in cellular bioenergetics. We utilized the Seahorse XF analyzer mito stress test, which allows the real-time measurement of both extracellular acidification rate (ECAR), a measure of glycolysis, and oxygen consumption rate (OCR), a measure of oxidative phosphorylation [58, 59]. A 1 h treatment with 0.15 to 10 μM GPC3_3 aptaprobe slightly increased baseline OCR than that in the control, and no difference in baseline ECAR (Additional file 1: Fig. S9B, C). The 10 μM aptaprobe concentration did not affect toxicity, cytoplasmic morphology, or mitochondrial respiration. Taken together, the aptaprobe has low (or no) cytotoxicity and
can be used in various in vitro and in vivo bio-imaging experiments.

Finally, we tested the ability of the GPC3_3 aptaprobe to bind GPC3-expressing tumor in HCC xenograft models in vivo. We established HCC xenograft tumors grown subcutaneously in BALB/c nude mice (Additional file 1: Fig. S10). To avoid the overlay between the tumor and other tissues in fluorescence imaging, an approximately 169.76 mm³ tumor was successfully induced between the right flank and thigh (Additional file 1: Fig. S11A). To confirm that the GPC3_3 aptaprobe targets the target GPC3 protein in vivo, the GPC3_3 aptaprobe and the same size of the random sequence was labeled at the 5′ end with Cy5.5 fluorescence and intravenously injected to measure the fluorescence distribution. The retention of the Cy5.5 signal was assessed using the IVIS system. We examined GPC3_3 aptaprobe binding to induce tumor intravenous injection. Figure 6A shows the fluorescence image of Balb/c nude mice bearing subcutaneous HCC tumors after intravenous injection of ControlCys and GPC3_3Cys. All injected formulations were imaged immediately after injection, indicating the whole body through blood circulation after intravenous injection. The GPC3_3Cys signal shifted to the right flank and thigh, where the induced tumor was localized 30 min after injection. The fluorescence signal disappeared 2 h after injection, indicating that GPC3_3Cys can be excreted through kidney, liver, and other metabolic organs. While after 2 h the injection of GPC3_3Cys, no obvious fluorescent signal was observed at the tumor sites, indicating that GPC3_3Cys does not accumulate in tumors in vivo. The fluorescence intensity of GPC3_3Cys also exhibited a plateau at the beginning and then gradually decreased after 2 h. The fluorescence signal intensity of normalized tumor area with GPC3_3Cys was significantly higher than that of ControlCys signal (1.4 × 10⁹ vs 2.45 × 10⁹ counts; P = 0.02; Fig. 6B). After 60 min injection, it reacts the maximum of 2.60 × 10⁹ counts, then gradually decreases to 1.78 × 10⁹ counts after 2 h. Then, the biological distribution of ControlCys and GPC3_3Cys in mice were photographed after injection 30 min and 60 min. The GPC3_3 aptaprobe (as indicated by fluorescence) was mainly distributed in the liver and kidney (Additional file 1: Fig. S11B), consistent with the biodistribution data of ControlCys [60–62]. The accumulation of GPC3_3Cys and ControlCys in liver and kidney tissue was similar at both time points, while the GPC3_3Cys accumulated in tumor tissue in the first 30 min after injection and it seems started being cleared from the tissue after 1 h. The tumor tissue of 60 min were measured by increasing the fluorescence intensity, we still found that the fluorescence signal

![Fig. 6](image-url)
was also observed in the tumor tissue of the GPC3_3 aptaprobe injected mice group, but no ControlCy5.5 signal was not observed in the control group (Fig. 6C). This indicates that the GPC3_3 aptaprobe, which has high affinity and specificity to the GPC3 protein, is quickly distributed and targeted to HCC tumor tissue. This verifies that the aptaprobe can have high specificity and discriminate the target protein in complex biological samples. It should be emphasized that the relatively weak fluorescence of GPC3_3 aptaprobe in the tumor fluorescence image might be due to fluorescence quenching by in vivo biomolecules because the tumors were isolated > 1 h after the injection of GPC3_3 aptaprobe. After intravenous injection of GPC3_3 aptaprobe, the signal in the tumor region dramatically increased, leading to the visualization of the tumor at 1 h post injection (Fig. 6A). The fluorescence signal of the aptaprobe in the tumor region appears faster than that in the in vivo imaging experiment using other materials; such nanoparticles appeared within 72 h, and biological molecules (peptides and aptamers) were observed within 1 to 2 h (Additional file 1: Table S2). This should be mainly attributed to the fact that fluorescence signal imaging usually relies on in vivo experimental conditions. These data highlight that aptaprobe-based fluorescence imaging allows fast detection of tumors with high specificity and targeting over other materials.

A recent study developing an in vivo tumor-targeting system using a tumor-specific aptamer showed a significant accumulation of aptamer in the tumor target site, followed by varying observation times based on their aptamer [63–66]. In comparison to existing aptamer-based in vivo imaging systems, additional research examining the nonspecific uptake of the proposed imaging probe into the liver and kidney is required to improve aptaprobe-based imaging’s optimal biodistribution. The aptaprobe injection method, concentration, injection time, tumor types, and tumor stage can all influence aptaprobe accumulation and biodistribution. The next phase in the development of this multimodal diagnostic aptaprobe for in vivo imaging will be to find a potential solution to improve biodistribution, such as a study to avoid natural buildup as nonspecific uptake into the liver and kidney by changing experimental settings.

**Conclusion**

GPC3 is a promising biomarker for early HCC diagnosis. Aptameres were successfully isolated using the SELEX methodology, and their binding moieties were predicted through MOE docking analysis. To explore the potential use of GPC3 aptamers in HCC diagnosis, Aptablotting, ALISA, and intra-/extracellular GPC3 imaging were performed. Aptameres do not cross-react with non-target molecules in a complex biological mixture, and aptablotting undoubtedly simplifies the immunoblotting workflow. Here, we showed through structure-based molecular interaction that aptaprobe pairs are significant for designing quantitative ALISA platforms. FAM-labeled GPC3 aptameres were successfully used to fluorescent stain Hela, HepG2, Hep3B, PANC-1, and MIA-PaCa cells, which will provide insight into the development of aptaprobe-based atomic imaging diagnostic technology. Our GPC3 aptaprobe can potentially be used as an affinity reagent to detect in vivo and ex vivo HCC diagnosing systems such as Aptablotting, ALISA, and aptaprobe-based imaging. Therefore, it is reasonable to speculate that our aptaprobe strategy might be useful in HCC diagnosis.

**Supplementary Information**

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**Additional file 1:** Figure S1. SELEX round eluted aptamer concentration. Figure S2. Surface plasmon resonance (SPR) analysis of the interaction of GPC3 and GPC3 aptamer candidates (GPC3_1 to GPC3_12) using a Biacore X100. Figure S3. Schematic of the GPC3 protein. Figure S4. Comparing the GPC3 protein and aptamer complex conformation by docking simulation. Figures S5–S6. Ligplot molecular interaction analysis of aptameres with the GPC3. Figures S6. Aptablotting assay and immunodot blot assay. Figures S7. Confocal imaging and fluorescence intensity quantification of FAM-labeled GPC3 aptameres in cell line. Figures S8. mRNA expression test to confirm GPC3 expression of in Hela cells, liver cancer cells (HepG2 and Hep3B), and pancreatic cancer cells (PANC-1 and MIA-PaCa). Figures S9. Cell viability using 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide salt (MTT) assay and Seahorse analysis for HepG2 cells treated with different aptaprobe concentrations. Figures S10. Schematic demonstrating the experimental workflow of in vivo GPC3_3 labeling in an intravenously hepatocellular carcinoma (HCC) xenograft model. Figures S11. Representative images of HCC xenograft model. Table S1. Amino acid residue involved in hydrophobic contact of the between GPC3 protein and aptamer candidate. Table S2. Features of biomolecules for interventional fluorescence imaging of tumors in preclinical studies.

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**Author contributions**

YHK, JYA, and JM conceived the project. WRS and DYP performed the experiments with help from JHK, JPL, NQ, IH, and WC. All authors wrote the manuscript with help from SSS, SYK, BKC, and SCK. All authors read and approved the final manuscript.

**Availability of data and materials**

Research data refer to the results of observations or experimentation that validate research findings. Data includes “raw” data and processed data.
Declarations

Competing interests

The authors declare no conflict of interest.

Author details

1 School of Biological Sciences, Chungbuk National University, 1 Chungdae-Ro, Seowon-Gu, Cheongju 28644, Republic of Korea. 2 Department of Biological Sciences, Korea Advanced Institute of Science and Technology, 291 Daehak-ro, Yuseong-gu, Daejeon 34141, Republic of Korea. 3 Graduate School of Semiconductor and Chemical Engineering, Jeonbuk National University, Iksan 570-970, Republic of Korea.

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