Preparation of long single-strand DNA concatemers for high-level fluorescence in situ hybridization

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Abstract

Fluorescence in situ hybridization (FISH) is a powerful tool to visualize transcripts in fixed cells and tissues. Despite the recent advances in FISH detection methods, it remains challenging to achieve high-level FISH imaging with a simple workflow. Here, we introduce a method to prepare long single-strand DNA concatemers (IssDNAcs) through a controllable rolling-circle amplification (CRCA). Prepared IssDNAcs were used to develop novel AmpFISH workflows. In addition, we present its applications in different scenarios. AmpFISH shows the following advantages: 1) enhanced FISH signal-to-noise ratio (SNR) up to 160-fold compared with single-molecule FISH; 2) simultaneous detection of FISH signals and fluorescent proteins or immunofluorescence (IF) in tissues; 3) simple workflows; and 4) cost-efficiency. In brief, AmpFISH provides convenient and versatile tools for sensitive RNA/DNA detection and to gain useful information on cellular molecules using simple workflows.

Introduction

Complex biological tissues consist of different types of cells. Characterizing the temporal and spatial heterogeneities of RNA or DNA is critically important to understand cellular functions. Fluorescence in situ hybridization (FISH), a powerful tool, allows the detection of nucleic acid molecules within the intact structure and morphology of cells or tissue sections. FISH has been widely applied in the research1–3 and clinic diagnostic assays4,5. The readout of the FISH signal can be achieved using fluorophore-tagged oligonucleotide probes that bind with specific nucleic acid sequences by complementary pairing. Therefore, the spatial and quantitative information of nucleic acids can be detected in the cells or tissues under a fluorescent microscope using single-molecule RNA fluorescence in situ hybridization (smFISH) imaging, which allows visualizing the spatial distribution of transcripts through hybridization of target genes6. Because it is difficult to detect individual fluorophores by conventional fluorescence microscopy, smFISH usually requires multiple fluorophore-tagged oligonucleotide probes to target different regions of RNA or DNA7 to acquire detectable signals. Despite the accumulation of multiple fluorophore molecules to a certain extent, the accuracy of the smFISH assay is still affected by the low signal intensity with high background signals from the tissue. Moreover, smFISH protocols usually are limited to long RNA molecules (>3 kbp) that can provide more fluorescent-binding sites to achieve enough detectable signals8.

The literature reports several amplification strategies for FISH, such as hybrid chain reaction (HCR)9,10, branch DNA (bDNA)11,12, ClampFISH13, and SABER14. Among these, HCR allows in situ amplification via folding hairpin oligonucleotide pairs that trigger a self-assembly chain. However, HCR may not offer sufficient sensitivity in certain cases15, and designing hairpin structures is a time-consuming process. Moreover, there is a lack of systematic guidelines for hairpin design16. All other amplification strategies for FISH can provide high signal levels; however, they require the synthesis of more expensive modified oligonucleotides, such as bDNA11,12, azide–alkyne13, and inverted bases14. An ideal FISH imaging should
display both cost-efficiency and bright signals that can potentially shorten the exposure time for imaging, thereby achieving high accuracy at low costs.

Here, we established a more cost-efficient strategy for high-level FISH signals. First, a method was prepared to develop a long single-strand DNA concatemer (IssDNAC) through controllable rolling circle amplification (CRCA). Second, based on IssDNAC, several easily implemented workflows of AmpFISH were established. Lastly, we used simple AmpFISH workflows to specifically detect transcripts or chromosomes in the cells and tissues. Our results showed that AmpFISH has several advantages: 1) up to 100-fold signal-to-noise amplification; 2) simultaneous detection of FISH signals with fluorescent proteins or immunofluorescence (IF) in tissues; 3) simple workflow; and 4) cost-efficiency. To sum up, we present a convenient and versatile tool for FISH detection at a high level, providing a promising application for transcript and chromosome assays in the cells or complex and heterogeneous tissues.

Results

Design of amplification method and preparation of hybrid probes for AmpFISH

To establish a novel pipeline for high-level FISH assay, we developed a preparation method for long single-strand DNA concatemers (IssDNAC) using rolling-circle amplification *in vitro*. Here, the concatemer sequence provides multiple repeat sites for binding multiple fluorophore-tagged oligonucleotide probes. The accumulated probes can effectively amplify the FISH signal.

First, we designed two partially complementary DNA oligonucleotides, named padlock and adaptor, as shown in Fig. 1A. At the 5’ terminus of the adaptor, we designed a toehold sequence that was complementary to target molecules. The sequences of 14 to 16 nucleotides at the right arm (B1) and ten nucleotides at the left arm (B2) of the padlock were complementary to the adaptor, respectively. After annealing to the adaptor, the right arm (5’ phosphorylation terminus) and left arm (3’-OH terminus) of the padlock formed adjacent nicking site when they were complementary with the 3’ terminus of the adaptor. Therefore, the right and left arms of the padlock could be ligated by DNA ligase to form a circular ssDNA. When DNA polymerase and dNTP were added to the reaction system, the ssDNA of the adaptor could be extended by displacement amplification, as shown in Fig.1A.

To establish a user-friendly reaction, we used an all-in-one tube strategy to amply adaptor ssDNA (see Materials and Methods). After padlock and adaptor were annealed, T4 DNA ligase was used to ligate the nicking site of the padlock. To extend the adaptor ssDNA, we utilized DNA polymerases with displacement ability using the rolling circle. At first, two kinds of DNA displacement polymerases, phi29 and Bst 2.0, were tested. After extended by DNA polymerases, the products were run through agarose gel electrophoresis, as shown in Fig 1B. Both DNA polymerases could extend to form clear amplification bands in the gel. Phi29 DNA polymerase provided rapid extension, whose amplification length reached up to 10 kb (amount to dsDNA marker) in only 10 min, and its extension speed was consistent with that reported in the previous literature\textsuperscript{22}. However, overlong DNA molecules extended by phi29 DNA
polymerase were not suitable for FISH assay, because a molecule with large molecular weight results in slow diffusion and poor penetration ability and could affect the efficiency of FISH\textsuperscript{23}. Therefore, we excluded phi29 DNA polymerase in subsequent experiments. In contrast, Bst 2.0 DNA polymerase showed relatively slower amplification at 50 °C, which extended approximately 0.5 kb in length after 1 h (Fig. 1B). To investigate its extension ability under different conditions, we amplified the ssDNA adaptor under three reaction temperatures (45 °C, 50 °C, and 55 °C) with three reaction times (0.5 h, 1 h, and 2 h). The results of the electrophoresis assay showed that their extension length increased slightly as the time or temperatures increased, indicating that Bst 2.0 DNA polymerase provided a relatively controllable extension (Fig. 1C). Therefore, the IssDNAc lengths can be more easily modulated by changing the reaction time only. Similar to Bst 2.0 DNA polymerases, Klenow (exo\textsuperscript{–}) exhibited the ability of controlled extension at 37 °C (Supplementary Fig. 1). After gel electrophoresis, we recycled the extended IssDNAcs from agarose gel using a gel extraction kit to obtain purified extended ssDNAc for the FISH assay. The schematic representation of IssDNAc preparation workflows is depicted in Fig. 1D.

**AmpFISH effectively amplifies fluorescent signals in cultured cells**

To evaluate the feasibility of ssDNAc as a FISH probe, we first used AmpFISH in HeLa cells that were transiently transfected with the pCAG-EYFP plasmid. A set of probes containing four IssDNAcs with ~750 bp in length could target EYFP mRNA. They were mixed and added to the cells transiently transfected with the pCAG-EYFP plasmid. A workflow of AmpFISH that could retain fluorescent protein for cultural cells was developed, allowing simultaneous detection of EYFP and AmpFISH (see Materials and Methods). The imaging results showed that the AmpFISH signal perfectly localized to the cells positive for EYFP fluorescence, whereas cells with negative EYFP fluorescence did not exhibit FISH signal (Fig. 2A). These results indicate that IssDNAc can be effectively used to assess EYFP mRNA expression with high specificity in cultured cells. As shown in Fig. 2A, intensities of the AmpFISH signal were not completely consistent with those of EYFP. We presume that the turnover pattern of rapid mRNA\textsuperscript{24} was different from that of EYFP protein.

To quantitatively compare the intensities or signal-to-noise ratio (SNR) of AmpFISH, we performed a series of FISH experiments for exogenous Cas9 mRNA assay in stable cell lines transfected with the pX330-Cas9–2A-Cerulean-2A-PuroR plasmid. Conventional smFISH (without extension), primary, secondary, and tertiary amplifications were performed (Fig. 2B). Among these, the secondary amplification included branched structures that were formed by secondary concatamers by binding to primary concatamers. Tertiary amplification formed high-level branching based on secondary amplification, as shown in the upper panel of Fig 2B. We quantified and compared the signal of different amplifications. The results showed that smFISH presented an extremely faint signal (Fig. 2B, the most right panel). Compared with the SNR signal of smFISH, the primary, secondary, and tertiary amplification AmpFISH were enhanced by 18.9-fold, 33.2-fold, and 168.3-fold improvement, respectively (Fig. 2B and 2C).

**AmpFISH enables robust chromosome assay**
To check the feasibility of AmpFISH in chromosome assays, we detected the chromosome ploidy of cultural cells. Aneuploidy alterations are hallmarks of tumor aggressiveness and increase with malignant progression\textsuperscript{25}. The chromosome ploidy can be used for clinic cancer assay. Because centromere satellite repeats provide multiple binding sites, we only used one IssDNAc to target the centromere of chromosome 8. After hybridization, tomography of the cell nucleus was performed by confocal microscopy (Supplementary Movie), and subsequently, three-dimensional (3D) images were merged, as shown in Fig. 3. The results showed that HeLa cells have a triploid karyotype of chromosome 8, which is consistent with that reported in a previous study\textsuperscript{26}. The results revealed that AmpFISH efficiently labeled interphase chromosomes of HeLa cells and is a promising assay for assessing chromosome ploidy.

**AmpFISH enables robust transcript assay in tissues**

Next, we assessed whether AmpFISH could robustly assay transcripts in tissues, apart from in cultured cells. We used mouse brain tissue to check the feasibility of AmpFISH for tissue assay. We first tested the primary amplification probes for *Gad1* mRNA via methanol penetration. However, the primary amplification probes provided an extremely faint signal with a high tissue background (Supplementary Fig. 3A). Next, further branching amplification via secondary amplification probes was performed. The secondary amplification enhanced the FISH signal compared with primary amplification (Supplementary Fig. 3B).

Given that brain tissue is rich in lipid components that could contribute to a high-signal background\textsuperscript{27}, we optimized AmpFISH workflows to clear the background. We evaluated varying concentrations of sodium dodecyl sulfate (SDS) and methanol permeabilization (with/without) in four experimental workflows, as shown in Fig. 4A. The signal intensity and the SNR from four workflows were compared (Fig. 4B and 4C). Treatment with 0.2% or 1% SDS and methanol permeabilization improved SNR by three to four folds when compared with no SDS and methanol treatment and by ~two folds when compared with methanol treatment only.

To further verify the accuracy of our experimental workflow, we detected EGFP mRNA using methanol and 1% SDS treatment described above in brain tissue slices of *Thy1*-EGFP transgenic mouse. Fig. 4D (left panel) shows that the fluorescence signal of EGFP was retained through treatment with 1% SDS and 100% methanol, allowing to assess the accuracy of AmpFISH by analyzing the distribution of AmpFISH signal and fluorescence signal from proteins in the same slice. Fig. 4D (middle and right panels) show that the AmpFISH signal (red) perfectly matched with the EGFP fluorescence (green) in the same cells, demonstrating that the AmpFISH workflow that we developed exhibited specificity in the brain slices.

**AmpFISH in combination with IF in brain slices**

Immunofluorescence (IF) can visualize cellular antigens such as proteins based on the antigen-antibody reaction. IF is extensively used in the field of clinical diagnostics, research, and drug development. To study multiple applications for AmpFISH, we optimized the AmpFISH workflow to make it compatible with IF. Briefly, SDS was replaced by HCl, proteinase K, and H\textsubscript{2}O\textsubscript{2} in the optimized workflow to treat the
tissues (see Materials and Methods). To assess the sensitivity of AmpFISH and IF under the optimized workflow, we simultaneously detected protein and mRNA of calcium/calmodulin-dependent protein kinase II a (CaMKIIa) by a combination of IF and AmpFISH in the same slices from the caudate putamen (CPU) area of the brain. As shown in the upper panels of Fig. 5A, up to 90% of AmpFISH (red) and IF signals (green) overlapped. Moreover, the proportion of overlap accounting for positive AmpFISH cells (91.68% ±2.07%) was slightly lower than that for positive IF signal (98.50% ±1.14%) (Fig. 5B), suggesting that the sensitivity of AmpFISH should be better than that of IF in our workflow.

To demonstrate the ability of AmpFISH to identify the neuronal cell population, we used the established AmpFISH/IF workflow to detect the proportion of subpopulation in the two midbrain regions, the ventral tegmental area (VTA) and substantia nigra pars compacta (SNC), respectively. Tyrosine hydroxylase (Th) protein and CaMKIIa mRNA were detected simultaneously in the same slice from the two midbrain regions, as shown in the below panels of Fig. 5A. We compared the proportion of Th+/CaMKIIa+ subpopulation to Th+ population in the two brain regions. The results showed that this proportion was significantly higher in the VTA area (25.27% ±0.46%) than in the SNC area (13.97% ±1.04%) (Fig. 5C). The projections and functions of neurons correlate with their expressed molecules in certain cell types. Whether different proportions of Th+/CaMKIIa+ cells in VTA and SNC correlate with their functions in different brain regions needs to be further studied.

Discussion

In this study, we provided a novel methodology to detect transcripts and chromosomes with a high imaging efficiency and simple workflows. We first developed a controllable rolling-circle amplification (CRCA) to prepare IssDNAc that could bind to a high number of fluorescent oligo-probes. Based on IssDNAc, we established simple workflows for AmpFISH that are easily implemented.

Previously published studies reported that the padlock DNA probe could be efficiently extended in situ by phi29 DNA polymerase via RCA, in which the enzyme was added to tissues. However, the in situ signal amplification of RCA may be limited. To improve the SNR following in situ amplification, the brain slices usually require further tissue clearing and hydrogel protocols to improve imaging resolution. However, this process could perturb tissue components. AmpFISH overcame the problem of in situ amplification and efficiently enhanced the SNR through branching amplification, with no requirement for further tissue clearing and hydrogel protocols.

Each 200 mL of the amplification system was used to extend the IssDNAc, and 1.2 mL (100 mM) of the adaptor and 1 µL (100 mM) of padlock oligonucleotide were added to the system, respectively. The average amount of recycling IssDNAc was 3 mg to 5 mg. A sample of the FISH experiment required 0.5 mg for each IssDNAc, meaning that one tube of ssDNAc probes can be used for 6 to 10 samples. The cost of ssDNAc preparation ranged from $0.28 to $2.1 per assay (Supplementary Text), which is dramatically lower than the cost of commercial probes. Moreover, we used 1 mL of the hybridization...
buffer to hybrid the samples. We will further develop a hybridization workflow with a smaller volume buffer for hybridization to increase its cost-effectiveness.

Penetrants, such as SDS, methanol, H$_2$O$_2$, HCl, and proteinase K, improve the permeability of the tissue. However, we found that an excellent effect cannot be achieved if all reagents were added. The excessive penetration may also lead to the loss of RNA. Another important factor for AmpFISH is PFA fixation. Both over fixation and no fixation should be avoided. Pre-fixation is required in our workflows during transcardial perfusion, and no to 1 h post-fixation is acceptable. In addition, the sample should always be attached to the glass slide; otherwise, it cannot be used. For instance, in the previously described IF/FISH protocol\textsuperscript{14}, in which proteinase K was directly used to digest the tissue slices after sectioning, we found it was difficult to attach the tissue slices to the glass slide after digestion by proteinase K. In our developed workflow, one more 10-min PFA fixation was performed after slice sectioning, easily allowing the binding of the tissue slice to the glass slide.

The AmpFISH workflows are compatible with fluorescent proteins, which improve its application in certain scenarios. For example, in combination with transgenic mice expressing GFP, AmpFISH can determine the gene expression by overlapping the signal. In addition, the spatial distribution of molecules can be obtained by studying the GFP contour. Although the signal intensity of GFP is partly reduced after penetration, most signals of fluorescent protein can still be detected. Proteinase K should be avoided in the protocol because the GFP signal will be lost completely. We penetrated the tissue by adding TritonX-100 and SDS instead of proteinase K. In contrast, moderate proteinase K was used to repair the antigen but not SDS in AmpFISH/IF experiment.

We noted that the sensitivity of IF greatly depends on its corresponding antibody, which may vary in titers when they are from different manufacturers. For example, we used two kinds of anti-Th antibodies from different manufacturers, which exhibited a significant difference in the IF signal for Th (Supplementary Fig. 4). In contrast, AmpFISH provided a more stable assay than IF and it does not depend on antibodies that are not always commercially available, making it a promising alternative under certain situations. Although some signal-enhanced methods for IF, such as Immuno-SABER\textsuperscript{31}, Immuno-RCA\textsuperscript{32}, etc. were developed, all of them require antibodies for specific proteins, which are not always commercially available or work stably. In contrast, AmpFISH probes can much easier and cheaper to be synthesized. Moreover, AmpFISH exhibited sensitivity and accuracy in both cells and tissues. And it is potentially suitable for imaging of multiplex transcripts. Further work is required to explore these possibilities. All together, AmpFISH is a convenient and versatile tool for FISH detection at a high level, offering a unique practical tool for developmental biology research, disease prognosis, and novel therapeutic developments.

**Materials And Methods**

**Mice**
All animal procedures were conducted according to the Guide for the Care and Use of Laboratory Animals of Huazhong University of Science and Technology and approved by the Animal Care Committee of Hubei Province, China. Animals were individually housed in a 12 h (7 am–7 pm) light/dark cycle, with food and water ad libitum. To avoid the circadian expression of gene expression, we sacrificed mice at approximately 10 am each time. The mice used for Fig. 4D were transgenic mice of Tg(Thy1-EGFP)MJrs (Jackson Labs stock #007788) (8–9 weeks). In other experiments, mice used were male C57/BL6 (8–9 weeks).

**NIH3T3 cell and HeLa cell culture**

NIH3T3 and HeLa cells were purchased from Wuhan University Preservation Center (Wuhan, China). All cells were grown in Dulbecco's modified Eagle medium (DMEM, Gibco) supplemented with 10% fetal bovine serum (FBS, Sigma), 50 U/mL penicillin, and 50 mg/mL streptomycin (Gibco, cat. no.15070) at 37 °C under 5% CO₂. To transiently transfect EYFP in HeLa cells, we used polyethylenimine (25 Kd, Polysciences, Inc., cat. no. 23966) as the delivery reagent to transfec the pCAG-EYFP plasmid. After three days of transfection, the transfected HeLa cells were used for conducting AmpFISH experiments.

To compare the signal magnification of AmpFISH (Fig. 2B), we established a stable NIH3T3/Cas9 cell line. The stable NIH3T3/Cas9 cell line was transfected with pX330-Cas9-2A-mCerulean-2A-puroR plasmid (Note: mCerulean is the variant of cyan fluorescent protein and puroR is puromycin resistance gene; the map and sequence are shown in Supplementary Fig. 2). After one month of puromycin screening, the stable NIH3T3/Cas9 cells with cyan fluorescent signal were established. We seeded these cells on round glass slips with 18 mm diameter that can be placed inside 12-well plates. When the cells density reached 60 to 80% confluency, the cells were used for FISH.

**FISH probe design**

In order to design amplification probes, the toehold sequences in the adaptor were designed using the Picky 32-bit versions software. The sequences of mouse transcripts (Mus_musculus.GRCm38.cdna.all.fa.gz) were downloaded from ftp://ftp.ensembl.org/pub/release-102/fasta/mus_musculus/cdna/. The parameters of Picky were set as GC content 30–70%, oligo size 30–40 for toehold part, and Tₘ value > 75 °C. The complementary sequence to the padlock circle was added to the 3' end of the toehold sequence. Finally, the whole adaptor sequences were further assessed by the UNPACK online software (http://www.nupack.org/partition/new). To minimize the probability of secondary structures, we tried to avoid using guanine in the padlock. The Tₘ values were calculated using the following formula:

\[ T_m = 81.5 + 16.6 \log M + 0.41 \%G+C - 0.72 \% \text{formamide} \]
where $T_m$ is the melting temperature in degrees celsius, M is the monovalent salt molarity, (% G+C) is the percentage of guanine and cytosine in the DNA strand of interest, and (% formamide) is the percentage of formamide added.

**Preparation of FISH probes**

The oligonucleotide and fluorophore-tagged oligonucleotides (see Supplementary table) were synthesized by Tianyi Huiyuan Biotech. Ltd. Wuhan, China. The synthesized ssDNA of adaptors were purified by desalting, the synthesized ssDNA of padlock with 5' phosphorylation were purified by high-performance liquid chromatography (HLPC). The adaptor (0.6 µL, 100 µM) and padlock ssDNA (0.5 µL, 100 µM) were mixed and added to the EP tube with 42.9 µL of 1× annealing buffer (10 mM Tris, 50 mM NaCl, pH 8.0). The EP tubes with mixture were put in a dry bath incubator (MK200–2, Allsheng Instruments Co., Ltd., Hangzhou, China) at 95 °C for 3 to 5 min. Next, the power to the incubator was turned off to spontaneously cool down the mixture until the temperature reached room temperature (RT). Then 5 µL of 10×T4 ligase buffer and 2 µL of T4 DNA ligase (Takara Corp., Dalian, China) were added to the mixture. The total volume was made to 50 µL by adding ddH$_2$O. After overnight ligation reaction, 5 µL of 10×Bst 2.0 polymerase buffer, 3 µL of Mg$_2$SO$_4$ (8 mM), 3 µL of dNTP (10 mM/each), 1 µL of Bst 2.0 DNA polymerase (New England Biolabs, Ipswich, MA, USA), and 38 µL of ddH$_2$O were added to form the extension reaction system with 100 µL reaction volume. The extension reaction was set for 2 h at 50 °C. After the extension, LssDNAs were separated by electrophoresis in 1.0% agarose gel using the GelRed dye. Next, DNA gel images were acquired using the Bio-Rad Universal Hood II imaging system. The DNA bands were excised from the agarose gel with a clean, sharp scalpel. DNA was extracted using the QIAquick Gel Extraction Kit following the manufacturer's protocol. The purified ssDNAs were analyzed and quantified by NanoDrop™ 2000/2000c spectrophotometer.

**Coverslip functionalization**

Glass coverslips were placed in 12-well plates and soaked in 0.1% diethylpyrocarbonate (DEPC) water overnight. Next, cells were washed with autoclaved DEPC water. Next, the coverslips were treated with methacryloxypropyltrimethoxysilane (Bind-silane) for 1 h and further treated with the poly-L-lysine solution for 5 min. The poly-L-lysine solution was removed and coverslips were dried at 60 °C for 1 to 2 h.

**FISH in fixed cell chambers**

To fix the cultured cells, cells were rinsed in 1× phosphate-buffered saline (PBS) at room temperature, and then immediately fixed in 4% paraformaldehyde (PFA) for 10 min. Finally, the cells were rinsed in 1×PBS. For permeabilization, the cells were washed with the PBST buffer (1×PBS with 0.5% TritonX-100) for 10 min, then washed with 1×PBSTw (1×PBS with 0.1% Tween-20) for 1 min, and subsequently washed with 2×SSC/TritonX-100 (2×SSC with 0.1% Triton X-100) for 2 min. The primary amplification probe sets were denatured at 95 °C for 3 to 5 min. Each primary amplification probe at 0.5 µg/mL final concentration was added to the hybridization solution (2×SSC, 1% Triton X-100, 40% formamide, 10% dextran sulfate, 20 mM RVC). The cells were gently shaken overnight at 42 °C and then washed with 1×PBS containing 40%
formamide (3 × 10 min) and then with 2×SSCT buffer (3 × 10 min) at 42 °C. For secondary or ternary amplification hybridization, the probes were added to the hybridization solution (2×SSC, 1% Triton X-100, 20% formamide, 10% dextran sulfate), 20 mM RVC, 0.1 mg/mL salmon sperm DNA, and 0.5 µg/mL of each amplification probe with gentle shaking at 37 °C for 4 to 5 h. Next, the cells were washed with 1×PBS containing 20% formamide (3 × 10 min) and then with 2×SSCT (3 × 5 min) at 37 °C.

For fluorescence detection, tissue slices were rinsed once in 2×SSCT at RT and then stained with DAPI for 30 min. Next, the samples were washed with 1×PBS (3 × 10 min) containing 0.1% TritonX-100 at room temperature. Fluor hybridization solution (2×SSC, 20% formamide, 10% dextran sulfate, 150 nM fluorescent probes) was added and held at 37 °C for 30 min. Then, samples were washed with 2×SSC (3 × 10 min) containing 20% formamide at 37 °C. The round glass slides with samples were placed into a magnetic chamber (L-shape tubing type Chamlide CM-B18-1, Live Cell Instrument, Seoul, South Korea) for imaging.

**FISH for centromere detection in HeLa cells**

HeLa cells were seeded in 12-well chambers. After growing to 50 to 70% confluence, HeLa cells were rinsed with 1×PBS and then fixed by 4% PFA solution for 10 min. Next, the cells were rinsed with 1×PBS.

The cells were treated with 2×SSCT containing 70% formamide at 70 °C (3 min) and then transferred to ice-cold 70% ethanol (5 min), 90% ethanol (5 min), and 100% ethanol (5 min) successively.

After fixation, the cells were rinsed with 1×PBS (1 min), next with 1×PBST (1×PBS/0.5% Triton X-100) for 10 min, then with 1×PBSTw (1×PBS buffer/0.1% Tween-20) for 2 min. The cells were incubated in 0.1 N HCl (5 min) and washed with 2×SSC/0.1% TritonX-100 (2 × 5 min). Next, the cells were incubated in 2×SSCT with 50% formamide (10 min) and transferred to fresh 2×SSCT with 50% formamide at 60 °C (at least 1 h). The wells were loaded with 500 µL of a solution containing 2×SSCT, 50% formamide, 10% (wt./vol.) dextran sulfate, and 400 ng/µL RNase A, primary probe (final concentration 500 ng/mL, denaturation at 80 °C for 3 min). The samples were incubated overnight at 44 °C. After hybridization, samples were washed with 2×SSCT at 50 °C (3 × 10 min). The samples were stained with DAPI and washed with 1×PBS (3 × 10 min).

The samples were incubated with fluorescent probes (150 nM final concentration) in the fluorescent hybridization solution (1×SSC, 30% formamide, 10% dextran sulfate at 37 °C (30 min). The cells were then washed (3 × 10 min) at 37 °C with wash buffer (1×SSC, 30% formamide).

**FISH in fixed tissue chambers**

The round glass slips with 18 mm diameter were autoclaved at 210 °C for 8 h. The glass slips were treated with methacryloxypropyltrimethoxysilane (Bind-Silane) and were further treated with poly-L-lysine solution. Adult mice were anesthetized with a mixture of 2% chloral hydrate (160–200 mg/kg) and xylazine (15–20 mg/kg) and transcardially perfused with 1× DEPC-PBS for 10 min, followed by treatment with 4% PFA in PBS. The brains were removed and immersed in 30% sucrose overnight at 4 °C until the
brains sunk to the bottom. Then brains were placed onto OCT and frozen at −80 °C with isopentane. Next, 16 to 20 mm sections were prepared using a cryostat (Leica CM1700). The brain slices were attached to the pretreated glass slips, then fixed with 4% PFA in PBS at RT for 10 min, dehydrated through gradient methanol/PBS (50%–75%–95%) 3 min each, treated with methanol at −20 °C overnight, and then placed at −80 °C for 15 min before hybridization.

For hybridization, slices were taken from −80 °C and equilibrated to room temperature for 5 min. These were washed with 1xPBS/0.1% TritonX-100 (3 × 5 min) containing 0.1 U/µL RNase inhibitor. Primary amplification probes were denatured at 95 °C for 3 to 5 min and then directly added to the hybridization buffer (2×SSC, 1% Triton X-100, 40% formamide, 10% dextran sulfate, 20 mM RVC, 0.1 mg/mL salmon sperm DNA, 1% SDS). The final concentration of each primary amplification probe was 0.5 µg/mL. The slices were incubated at 42 °C with gentle shaking overnight, and then washed with 1xPBS/40% formamide (3 × 10 min) at 42 °C and then with 2×SSC/0.1% Triton X-100 at room temperature. For secondary amplification hybridization, secondary amplification probes were denatured at 90 °C for 3 to 5 min and were added to the hybridization solution (2×SSC, 1% Triton X-100, 25% formamide, 10% dextran sulfate, 20 mM RVC, 0.1 mg/mL salmon sperm DNA, 1% SDS) to form 0.5 µg/mL for each probes. Samples were incubated for 4 to 5 h with gentle shaking at 37 °C. Samples were washed 3 × 10 min with 1xPBS containing 25% formamide at 37 °C. The samples were then washed with 2×SSCT (3 × 5 min) at 37 °C and 2×SSC/0.1% TritonX-100 (3 × 5 min) at RT.

For fluorescent hybridization, samples were rinsed in 2×SSCT and then stained with DAPI (Invitrogen, no. D1306) for 1 h at RT. The samples were washed with 1xPBS (3 × 10 min) with 0.1% TritonX-100 at RT. Fluor hybridization solution (2×SSC, 20% formamide, 10% dextran sulfate) and 150 nM fluorescent probes were incubated at 37 °C for 30 min. The samples were washed 3 × 10 min with wash buffer (2×SSC/20% formamide) at RT. Round glass slides with samples were mounted onto a magnetic chamber (L-shape tubing type Chamlide CM-B18-1, Live Cell Instrument, Seoul, South Korea) for imaging.

**IF and FISH**

The animal brain slices were obtained and dehydrated as described above. The sections were taken from −80 °C, equilibrated to RT for 15 to 20 min, and washed with PBSTR (1xPBS/0.1% TritonX-100, 0.1U/µL RNase inhibitor) for 3 × 5 min. The sections were then treated with protein K (1.5 ng/mL) at RT for 15 min, HCl (0.1 N) at RT for 20 min, and 2% H₂O₂ at RT for 20 min, washed with PBSTR for 5 min, followed by incubation in 1× first hybridization buffer (40% formamide, 2×SSC, 1% TritonX-100, 20 mM RVC, 0.1 mg/mL salmon sperm DNA, 10% dextran sulfate, and probes at 500 ng/mL per oligo) at 42 °C with gentle shaking overnight. Then, the sections were washed with 40% formamide buffer (40% formamide/2×SSC) at 42 °C for 20 min thrice and PBSTR for 5 min, followed by incubation in 1× second hybridization buffer (25% formamide/2×SSC, 1% TritonX-100, 20 mM RVC, 0.1 mg/mL salmon sperm DNA, 10% dextran sulfate, and probes at 0.5 mg/mL per oligo) at 37 °C with gentle shaking overnight. The next day, the sections were washed with 25% formamide buffer (2×SSC, 25% formamide) at 37 °C for 3 × 20 min and PBSTR for 5 min, and then incubated with DAPI at RT for 30 min. Next, the sections were washed with
PBS at RT (3 × 10 min). The sections were incubated in the fluor hybridization solution (2×SSC, 20% formamide, 10% dextran sulfate, 150 nM fluorescent imager strands) at 37 °C for 30 min, and washed by 20% formamide buffer (20% formamide, 2×SSC) at 37 °C for 3 × 10 min.

For IF labeling, the sections were incubated with blocking buffer (1×PBS with 0.3% TritonX-100 and 10% goat serum) at RT for 30 min. Primary antibody for CaMKIIα was purchased from GeneTex Inc. (No. GTX127939), and anti-Th antibody was purchased from Sigma-Aldrich (No. AB152) and Abcam (No. ab6211). Primary antibodies were diluted at a ratio of 1:500 with blocking buffer, and samples were incubated overnight at 4 °C. Next, samples were washed with 1×PBS (3 × 10 min) and then labeled with secondary antibody (goat anti-rabbit Alexa 488, 1:500) for 2 h at RT. Finally, the samples were washed with 1×PBS (3 × 10 min).

**Imaging set-up**

For FISH imaging in HeLa and stable NIH3T3/Cas9 cells, the data were acquired by Nikon Ni-E microscopy with a 25× water-immersed objective lens (numerical aperture: 1.1); 405 nm and 561 nm lasers were used. A set of filters UV-2E/C (EX: 325–375, DM: 400, EM: 425–475) was used to visualize DAPI staining. G-2E/C (EX: 528–553, DM: 565, EM: 570–620) was used to visualize the TAMRA dye signal.

For GFP FISH imaging in HeLa cells and FISH/IF imaging in brain slices, data were acquired by Zeiss LSM 710 microscope; 405 nm, 488 nm, and 561 nm lasers were used. The following set of filters were used: filter set 49 (EX: G365, DM: 395, EM: 422–498) for DAPI dye, filter set 38 (EX: 470/40, DM: 495, EM: 495–553) for GFP fluorescent protein and Alexa 488, and filter set 43 (EX: 545/25, DM: 570, EM: 563–660) for TAMRA dye signal. Color images were captured with a Zeiss AxioCamICc5 camera.

**Quantification and statistical analysis**

The signal puncta were quantified using the Fiji software (NIH). The signal intensity of puncta was quantified in a 16-bit figure using a previously described method. Labeled neurons from each sample were randomly selected. Using the selection tool of ImageJ software, an area covering the signal spot was selected within the defined region, and the integrated density (IntDen) of the signal was measured using an ImageJ plug-in. To measure the background signal, a straight line was brought across the background area close to the signal spot, and the mean gray value was measured. The total mean value of the background of the rectangular selection was subtracted from the integrated density of the signal to obtain the total signal density in the rectangular selection. The average intensity of the signal without background was calculated by dividing the total signal density by the pixel number of the rectangular selection.

$$\text{Signal intensity} = \frac{\text{IntDen} - (\text{area} \times \text{mean of background})}{\text{Area}}$$
For statistical analysis, one-way analysis of variance (ANOVA) followed by Bonferroni correction was performed using GraphPad Prism 5 (Fig. 4C) and an unpaired t-test with Welch's correction (Fig. 5C). Differences between the datasets were considered significant at $p < 0.05$.

**Declarations**

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**Author Contributions**

DC, CX carried out mRNA hybridization in brain slices and cells and FISH imaging; SW prepared the FISH probes, DL conducted the DNA hybridization, KZ screened the stable cell line, JY carried out data analysis. All authors participated in planning of experiments, discussion and contributed to the preparation of the manuscript. QL and JY supervised the project.

**Conflict of interest**

The authors declare that they have no conflict of interest.

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**Figures**
Preparation of lssDNAc (A) Schematic representation of the extension reaction for lssDNAc. Both nucleotides of 5’ and 3’ termini of the padlock can complementarily bind to the 3’ termini of the adaptor.

Next, the padlock is ligated at 5’ and 3’ termini by DNA ligase to form a closed circle. The adaptor ssDNA can be extended using the closed padlock as a template using RCA. (B) The amplification products were assayed via agarose gel electrophoresis. The extended products of adaptor ssDNA were amplified by

**Figure 1**
phi29 DNA polymerase (lane 1 for 10 min, lane 2 for 0.5 h) and Bst 2.0 DNA polymerase (lane 3 for 0.5 h, lane 4 for 1 h). (C) The ssDNA adaptors were amplified to form products of different lengths by Bst 2.0 DNA polymerase (2–14 lane) under three reaction temperatures (45 °C for 2–5 lanes, 50 °C for 6–9 lanes, and 55 °C for 10–13 lanes) with different amplification times (0.5 h, 1 h, 1.5 h, 2 h), and in the absence of DNA polymerase (lane 1). (D) The schematic diagram depicting the preparation for lssDNAc and hybridization workflow.

Figure 2
AmpFISH enables to precisely label (A) and effectively amplify fluorescent signals (B-C) in cultured cells. (A) A set of primary amplification probes for targeting the EYFP mRNA are used in HeLa cells transiently expressing EYFP. FISH signal (red, signal labeled by Cy5, in the third panel from left in A) and fluorescent protein signal (green signal in EYFP panel, the second panel in A) were imaged in the same view field (scale bar: 20 μm). (B) Fluorescent signal labeled by TAMRA dye was amplified by AmpFISH in stable NIH3T3/cas9 cells. The confocal imaging for smFISH (unextended probe), primary, secondary, and tertiary amplifications are arranged from left to right, respectively (scale bar: 20 μm). (C) Comparative analysis of signal signal-to-noise ratio (SNR) of different hierarchies amplification.

Figure 3

AmpFISH efficiently labeled interphase chromosomes of HeLa cells. Most centromeres of HeLa chromosome 8 show three FISH signal puncta.
Figure 4

AmpFISH is adopted to robustly assay transcripts in mouse brain slices. (A) Schematic diagram depicting four different experimental workflows. (B) FISH imaging of Gad1 gene in mouse brain slices using different pipelines depicted in (A) (scale bar: 10 μm). (C) Statistical analysis of signal-to-noise ratio (SNR) of AmpFISH from four experimental pipelines in (B). One-way analysis of variance (ANOVA), F(3, 72) = 10.83, P < 0.0001; Bonferroni’s multiple comparison test, no treatment vs. 0.2% SDS, t = 4.206, P < 0.001; no treatment vs. 1% SDS, t = 4.810, P < 0.001; methanol vs. 0.2% SDS, t = 3.030, P < 0.05; methanol vs. 1% SDS, t = 3.633, P < 0.01. (n = 19, from three repeat experiments). (D) The specificity of AmpFISH via secondary amplification hybridization was verified in brain slices by expressing EYFP protein in Thy-EGFP transgenic mouse (scale bar: 25 μm).
Figure 5

AmpFISH assay in combination with IF. (A) IF and AmpFISH assays were verified mutually. IF for CaMKIIα protein and AmpFISH for CaMKIIα mRNA were double-labeled in the same slice from the CPu brain region (scale bar: 10 μm). (B) The TH+CaMKIIα+ cell subtypes were detected in the slice from the VTA brain region through a combination of IF with AmpFISH (scale bar: 10 μm). (C) The proportion of positive double-labeled cells was analyzed. “Overlap” represents cells with both IF-and AmpFISH-positive signals. Overlap/FISH represents the proportion of overlapping cells with a positive AmpFISH signal (91.68% ±2.07%, n = 5). In the same way, Overlap/IF represents the proportion of overlapping cells with a positive IF signal (98.50 ±1.14%, n = 5). The total number of “overlap” cells, cells with AmpFISH signal, and cells with IF signal were 170, 185, 172, respectively, from five different brain slices. (D) Statistical analysis and comparison of cell populations in the SNC and VTA brain slices. The proportion of TH+CaMKIIα+ cells (“overlap”) accounting for TH+ cells in the VTA area (25.27% ±0.46%) is significantly higher than that in the SNC area (13.97% ±1.04%) (n = 4) (unpaired t-test with Welch’s correction, two-tailed, P = 0.0006).
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