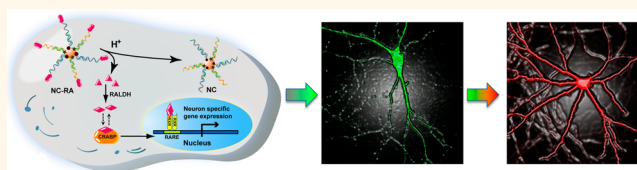


Bioinspired Nanocomplex for Spatiotemporal Imaging of Sequential mRNA Expression in Differentiating Neural Stem Cells

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ABSTRACT Messenger RNA plays a pivotal role in regulating cellular activities. The expression dynamics of specific mRNA contains substantial information on the intracellular milieu. Unlike the imaging of stationary mRNAs, real-time intracellular imaging of the dynamics of mRNA expression is of great value for investigating mRNA biology and exploring specific cellular cascades. In addition to



advanced imaging methods, timely extracellular stimulation is another key factor in regulating the mRNA expression repertoire. The integration of effective stimulation and imaging into a single robust system would significantly improve stimulation efficiency and imaging accuracy, producing fewer unwanted artifacts. In this study, we developed a multifunctional nanocomplex to enable self-activating and spatiotemporal imaging of the dynamics of mRNA sequential expression during the neural stem cell differentiation process. This nanocomplex showed improved enzymatic stability, fast recognition kinetics, and high specificity. With a mechanism regulated by endogenous cell machinery, this nanocomplex realized the successive stimulating motif release and the dynamic imaging of chronological mRNA expression during neural stem cell differentiation without the use of transgenic manipulation. The dynamic imaging montage of mRNA expression ultimately facilitated genetic heterogeneity analysis. *In vivo* lateral ventricle injection of this nanocomplex enabled endogenous neural stem cell activation and labeling at their specific differentiation stages. This nanocomplex is highly amenable as an alternative tool to explore the dynamics of intricate mRNA activities in various physiological and pathological conditions.

KEYWORDS: mRNA · imaging · gold nanoparticle · drug delivery · neural stem cell

Messenger RNA (mRNA) is one of the key factors that regulate cellular functions and is actively involved in numerous activities in the intracellular milieu.¹ The dynamic alteration of specific mRNA at a particular time or location in cells usually implies significant changes of cellular condition.² Aberrant mRNA dynamics is associated with developmental and pathological abnormalities, such as interrupted embryonic development³ and cellular apoptosis.⁴ The dynamics of mRNA expression is a blueprint of cancer progress from benign to malignant stages.⁵ The mRNA expression signature of a given neuron reflects its developmental history, activity, and interaction with other cells and the environment.⁶ Therefore, intracellular imaging of the dynamics

of mRNA expression is of great value for better investigating mRNA biology and exploring specific cellular cascades.

Real-time imaging of mRNA dynamics has recently drawn much attention,^{7,8} as mRNAs have intricate dynamics,⁹ which is evidenced by the delicately timed and spatially controlled processes in cells. The understanding of intracellular mRNA generation, progression, and mutual interaction was mainly dependent either on fluorescence *in situ* hybridization (FISH) of fixed cells or on biochemical separation of subcellular components followed by PCR amplification. Those methods, however, present only the static situation of cells at the terminating time point, when samples were prepared.⁹ In contrast, live cell dynamic imaging provides

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spatiotemporal mRNA profiling, which reveals extensive information on cell progress.

Recently, several methods have been developed for live cell mRNA imaging, such as molecular beacons^{10–15} and nanoflares.^{16,17} Some probes enabled concurrent imaging of multiple mRNAs in living cells to improve detection accuracy.^{16,18,19} Various real-time mRNA imaging probes have been developed,^{19,20} most of which, however, are capable of imaging only the stationary mRNA expression at a single time point in living cells. In addition, those imaging probes were designed to reveal the prevalence of pre-existing mRNAs in cells,²⁰ while the pivotal dynamics of mRNA sequential expression during specific cellular events, such as differentiation or apoptosis, remains largely elusive. Moreover, mRNA dynamic expression is usually initiated by extracellular stimuli.^{21,22} Such environmental cues usually result in rather low stimulation efficiency and a short half-life in media unless used in high concentration, which adversely exerts unexpected cytotoxicity.²³ In addition, the challenge of precise time coordination between stimulating factor function and imaging probe introduction may act as another barrier for real-time mRNA dynamic imaging.

To achieve the goal of dynamic imaging of mRNA sequential expression, the multifunctional probe should be easily delivered into cells, stable upon entry into cells, efficiently and homogeneously distributed in the cytosol, and specific and sensitive to its target. Ideally, it should incorporate a stimulating motif undergoing self-cleavage through endogenous cellular machinery in cells. In this study, we designed a multifunctional nanocomplex (fNC) meeting all the above criteria to enable self-activating and spatiotemporal imaging of the dynamics of mRNA sequential expression in differentiating neuron stem cells.

RESULTS AND DISCUSSION

Design of Multifunctional Nanocomplexes. The nanocomplex consists of a cellular machinery initiated system for successive gene expression stimulation and imaging. To stimulate gene expression, nature has provided us an instrumental guidance: in cells, the metabolism of a visual pigment, in which a small molecule binds to a pocket protein, is reversibly regulated by balanced Schiff base formation and hydrolysis through endogenous subcellular machinery.²⁴ Inspired by this well-established, naturally occurring mechanism, we designed a nanocomplex with the release of a stimulating motif controlled by Schiff base formation and hydrolysis. To construct the mRNA imaging probe, we used gold nanoparticles (AuNPs), a widely used potent delivery vehicle,^{25,26} as the core functionalized with a dense layer of dithiol group terminated recognizing DNA oligonucleotides hybridized with different fluorophore-capped reporter sequences. The fluorophores were initially quenched by a wide-spectrum

AuNP nanoquencher (Figure 1A). Unlike molecular beacons, which require transfection agents for delivery and are susceptible to enzyme degradation, oligo-modified AuNPs enable self-delivery and have high stability to resist enzyme cleavage.²⁷ The termini of recognizing oligonucleotides of AuNP are tethered with stimulating motifs *via* the labile Schiff base to enable self-activation of the fNC inside cells (Figure 1A and B). The fNC is stable at neutral pH, whereas upon cellular uptake, the Schiff base (imine) linkage is hydrolyzed to yield small molecules (retinals) and oligo-modified AuNP imaging probes. In mammalian cells, an enzyme, aldehyde dehydrogenase (RALDH), is responsible for efficient conversion of retinal into retinoic acid (RA), which specifically binds with cellular RA binding protein (CRABP) to activate the downstream gene transcription process for neuron-specific gene expression (Figure 1A).²⁸ Briefly, the low-pH microenvironment triggered the Schiff base conjugated retinal release from gold nanoparticles, and the released retinal was converted into RA by RALDH. The biologically active RA ultimately bound with CRABP to initiate neural stem cell differentiation. During the subsequent cell progression period, the mRNA targets will be chronologically expressed in the cytosol, where they will compete with the fluorophore-tagged reporter sequences and hybridize with recognizing oligonucleotides on AuNP by forming long and stable duplexes. This sequence-specific competitive binding causes the release of a reporter sequence from fNC to generate fluorescent signal proportional to the mRNA target expression level (Figure 1A). The fNC has high colloidal stability in the cytosol,²⁶ endonuclease resistance,²⁷ and stimulating motif self-cleavage capacity. Importantly, this nanocomplex system demonstrates, to the best of our knowledge, the first example of a dynamic imaging study of mRNA sequential expression using robust nanomaterials without transgenic manipulation.

Preparation of Nanocomplexes with Multifunctions. To demonstrate the function and efficacy of this fNC for self-activating and imaging the dynamics of mRNA expression, we applied it, as a proof of concept, to image the dynamics of mRNA sequential expression during neural stem cell (NSC) differentiation (Figure 1B). The differentiation of NSCs plays a central role not only in the developing mammalian nervous system but also in the adult nervous system of all mammalian organisms.²⁹ Transplanted NSCs are capable of ameliorating damage or pathological disorders by their differentiation into neural subtypes in the central nervous system.³⁰ During NSC differentiation stimulated by an external modulator,²¹ the genetic repertoire undergoes remarkable changes, where genes are chronologically regulated at specific developmental stages.³¹ In addition, the proliferation of NSCs undergoing differentiation is in a relatively static condition,

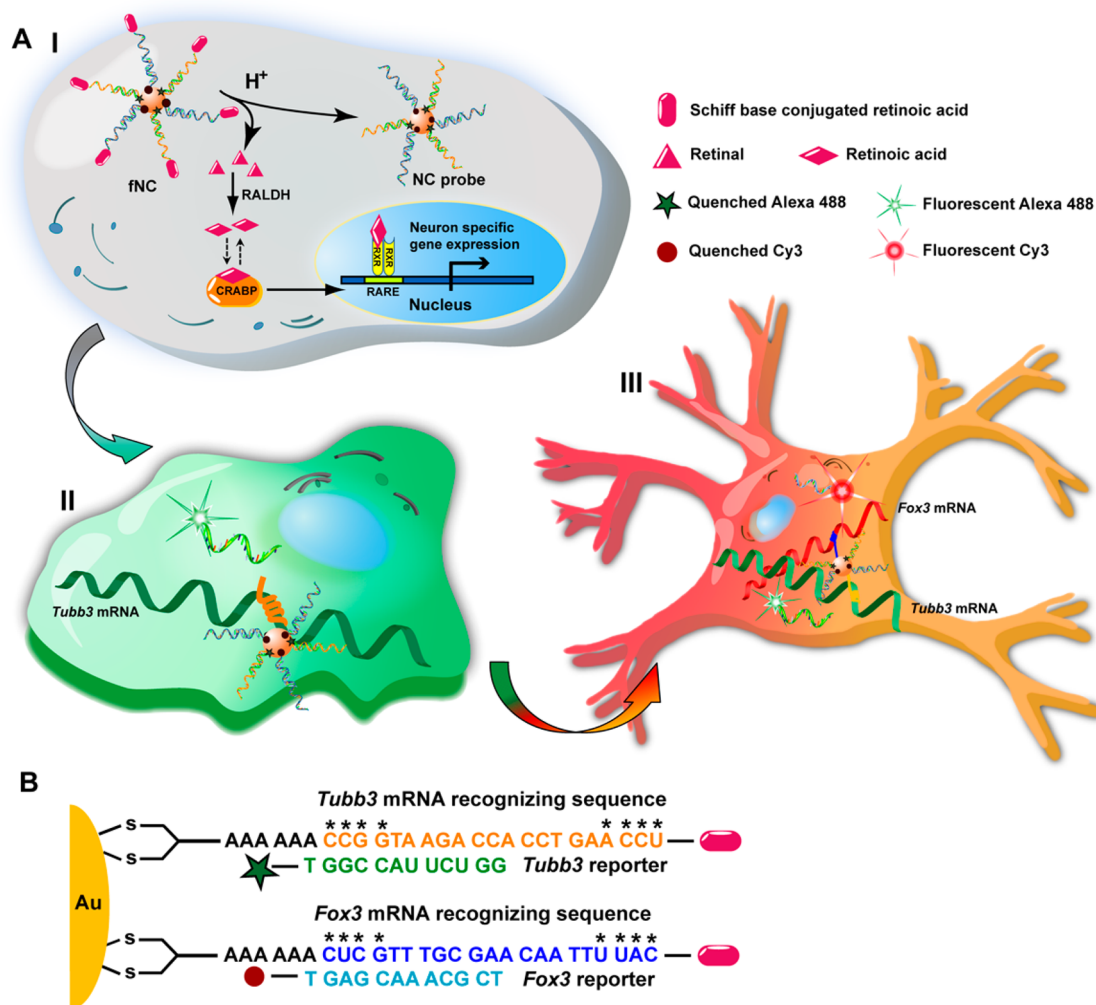


Figure 1. (A) Schematic illustration of a multifunctional nanocomplex (fNC) as an integrated system for tandem activation and imaging of the dynamics of mRNA sequential expression during the process of neural stem cell differentiation. Upon cellular uptake of the nanocomplex, the release of stimulating motif (RA) was controlled by Schiff base formation and hydrolysis through an endogenous mechanism. The released RA efficiently activated the subsequent signaling pathway to stimulate specific gene expression. In the meanwhile, the NC probe underwent sequence-specific recognition and spatiotemporal imaging of sequential mRNA expression during the process of neural stem cell differentiation. RALDH: retinaldehyde dehydrogenase; CRABP: cellular RA binding protein; RARE: RA responsive element; RXR: retinoid X receptor. (B) Design of the recognizing and reporter oligonucleotides for *Tubb3* and *Fox3* mRNA imaging. Oligonucleotides marked with * indicate methylation.

which facilitates imaging of the differentiation process in a fairly restricted environment without interference from the potential cellular division.²¹ Thus, real-time spatiotemporal imaging of the dynamics of mRNA sequential expression can provide insightful information on the biochemical basics of stem cell biology. To visualize the dynamics of mRNA expression by fNCs, we chose two representative neuron-specific mRNA pairs, *Tubb3* and *Fox3*, which are sequentially expressed during NSC differentiation from progenitor cells to neurons. *Tubb3* has been suggested to be one of the earliest markers of signal neuronal commitment,³² while *Fox3* shows dominance in neurons at a later stage.³³ Retinoic acid, commonly used to differentiate stem cell populations into neurons,²² was chosen as the stimulating motif and incorporated in the nanocomplexes. Since mRNA has secondary and tertiary

structures, the designed DNA should be complementary to the outward folding side of an mRNA, which is ready to be approached by other molecules. In addition, the reporter DNA oligos are relatively shorter than the entire complementary sequence of recognizing DNA. This design leads to a lower T_m value and thermodynamic energy, so that the reporter DNA is likely to be replaced by the complementary sequence of the target mRNA, leading to more stable hybridization. The recognizing DNA was terminated at the 3' end by an alkanedithiol group to facilitate highly stable anchoring on the AuNP surface, and the 5' end was flanked with primary amine to react with aldehyde from a retinoic acid analogue, retinal, to form a pH-labile Schiff base.³⁴ The DNA sequence consists of a 6-base-pair (bp) adenine (A_6) spacer and a 21-bp fragment complementary to the corresponding nucleotide

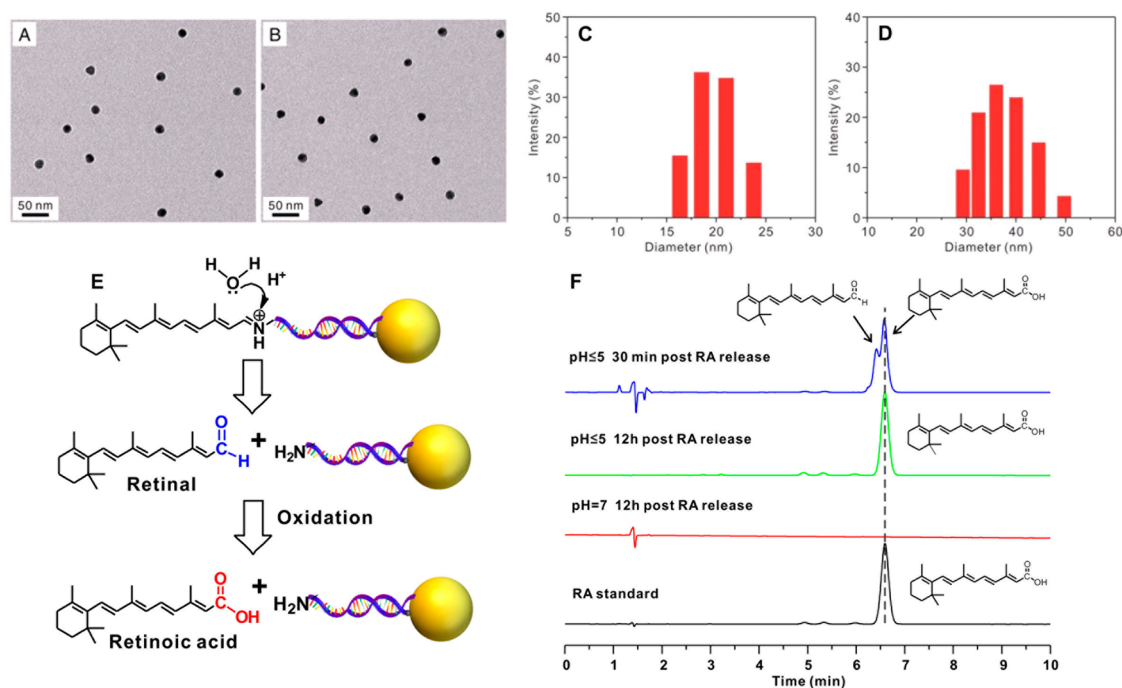


Figure 2. Nanocomplex morphology characterization and retinoic acid release. TEM images of naked gold nanoparticles (A) and multifunctional nanocomplexes (B). Dynamic light scattering (DLS) analysis: (C) naked gold nanoparticles in pure water; (D) multifunctional nanocomplexes in PBS. (E) Principle of Schiff base mediated stimulating motif (retinal) self-cleavage and oxidation under different physiological pH values. (F) HPLC chromatogram of corresponding released retinal or retinoic acid under different pH values and time points. The duplicate peak from the 30 min RA release chromatogram is attributed to the mixture of retinal and retinoic acid in solution.

sequence (*Tubb3*:228–248; *Fox3*:1349–1369) in mRNAs. The recognizing DNA was additionally methylated on selective nucleotides to enhance stability and binding affinity. The reporter sequence with a 5' end fluorophore tag (*Tubb3*: Alexa 488; *Fox3*: Cy3) (Table S1) is complementary to the 12 bp of the corresponding recognizing sequence. The 15 nm AuNP was used as the core. The multifunctional nanocomplex is monodispersed (Figure 2A and B) and highly stable (Figure 2C and D). The UV/vis spectrum of fNC shows a 4 nm red shift of the surface plasmon resonance (SPR) band compared with that of naked AuNPs (Figure S1). This suggests the change of the dielectric constant of the surrounding environment of the AuNP after dense oligonucleotide functionalization. In contrast, the width of the SPR remains unchanged, indicating the monodispersed condition of the fNC. This result is consistent with the TEM imaging (Figure 2A and B). Dynamic light scattering (DLS) measurements (Figure 2C and D) show the colloidal fNC with a diameter of 38 ± 1.6 nm, and the size of the original AuNP was 19 ± 0.6 nm, along with particle surface charge shifting from -15 ± 1.8 mV to -27 ± 2.1 mV, confirming the successful modification with DNA oligonucleotides on the AuNP surface. Retinal conjugation did not dramatically alter the surface charge (-24 ± 2.0 mV). The dense layer of methylated ssDNA oligonucleotides provides additional stability for AuNPs. On average, the fNC was calculated to carry

22 ± 1 *Tubb3*-recognizing DNA oligonucleotides tagged with Alexa 488 and 23 ± 2 *Fox3*-recognizing DNA oligonucleotides tagged with Cy3 (Figure S2A and B). The RA moieties tethered on the termini of oligonucleotides were 35 ± 3 molecules in each particle estimated by the HPLC method (Figure S2C). The calculated quenching efficiency of the gold nanoparticles was 92% for Alexa 488 and 87% for Cy3 (Figure S3).

Low Concentration of Stimulating Motif Released from fNC Activates NSC Differentiation. The fNC was integrated with a stimulating motif function *via* Schiff base. The fNC is stable at neutral pH (pH = 7) such that no free RA is detected, whereas at lower pH (pH < 5) the Schiff base is hydrolyzed into retinal, followed by oxidation into a functional RA molecule (Figure 2E and F).²⁸ In our study, the calculated fNC attaching RA applied for NSC stimulation was about 15 nM, which is much lower than the commonly used $1 \mu\text{M}$ concentration of free RA in media. Immunostaining results confirmed the identical NSC differentiation ability by these two conditions, but 15 nM free RA dissolved in culture medium did not show observable NSC differentiation (Figure S4). Another report showed consistent results that spatially positioned and concentration-controlled RA delivery into NSCs by nanoformulation offered 2 orders of magnitude higher efficiency than free soluble RA to induce an NSC proneurogenic niche in the mouse subventricular zone.²³ This indicates that the

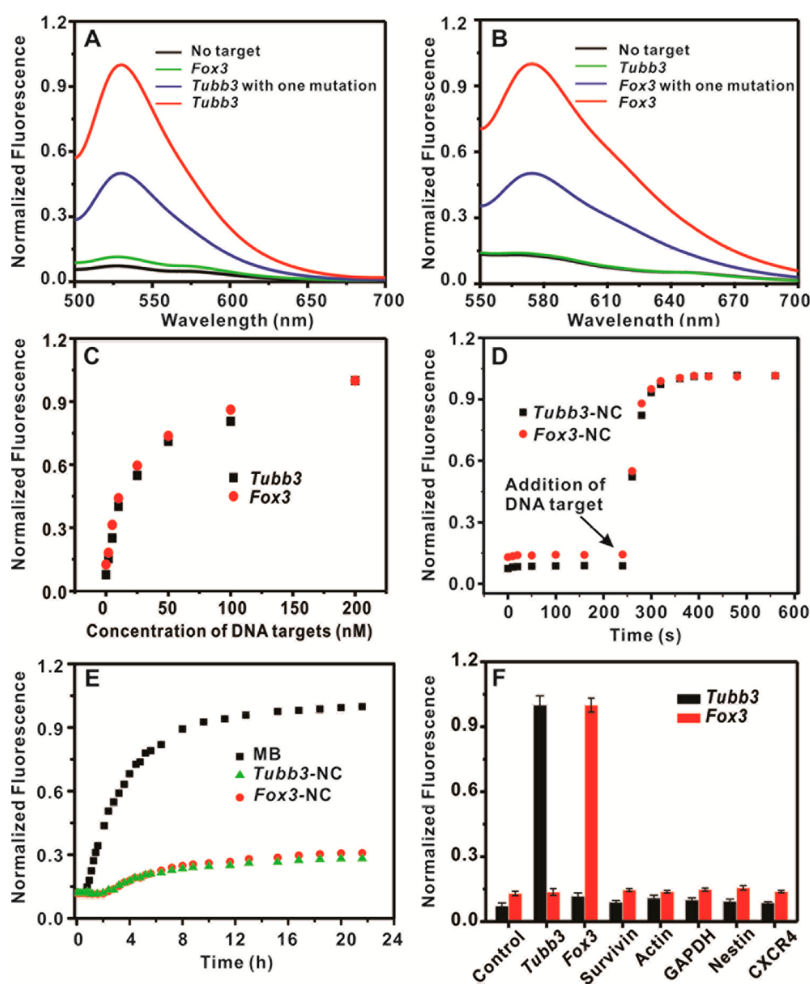


Figure 3. *In vitro* characterization of multifunctional nanocomplexes. (A) Fluorescence spectra of fNCs in the presence of no target (black line), *Tubb3* target (red line), cross-talk *Fox3* target (green line), or *Tubb3* target with one mismatch (blue line). (B) Fluorescence spectra of fNCs in the presence of no target (black line), *Fox3* target (red line), cross-talk *Tubb3* target (green line), or *Fox3* target with one mismatch (blue line). (C) Dynamic response of nanocomplexes to corresponding DNA targets with various concentrations of 0, 2, 5, 10, 25, 50, 100, and 200 nM. (D) Hybridization kinetics of fNCs associated with the complementary targets. The fNCs (1 nM) were mixed with 200 nM of *Tubb3* DNA target (black square) or *Fox3* DNA target (red circle), respectively. (E) Enzymatic stability study of fNCs (red and green) and molecular beacons (black). fNCs (1 nM) and molecular beacons (25 nM) were treated with DNase I (50 U/L). (F) Specificity of the nanocomplex over several DNA targets. The nanocomplex was mixed with *Tubb3* (black) or *Fox3* (red) complementary targets and other noncomplementary targets (targets of surviving, β -actin, GAPDH, Nestin, and CXCR4) measured with a 490 nm excitation wavelength for *Tubb3* and 550 nm for *Fox3* ($n = 3$).

nanoparticle delivery platform^{23,35–37} provides additional benefits to deliver and protect RA in cells to stimulate NSC differentiation. Moreover, this fNC takes advantage of endogenous subcellular machinery to spontaneously trigger RA release for NSC differentiation and simultaneously delivers the imaging probe, which avoids the need for complicated time coordination to image the dynamics of mRNA sequential expression during stem cell differentiation.

Specificity and Stability of fNC as an Imaging Probe. The binding specificity of fNC was first evaluated by adding a completely matched DNA target in fNC solution. To demonstrate the sequence specificity, a single mismatch (mutation) was introduced in the same DNA target. To exclude the possibility of cross-reaction between two targets on a fNC, the cross hybridization

of the fNC with an individual target was also evaluated. The results showed that the fNC responded with a significant signal (over 7-fold change) in the presence of a completely matched DNA target for both *Tubb3* and *Fox3*, while a single mutation in the DNA target resulted in 2.5 times lower signal response in both DNA targets (Figure 3A and B). This indicates that fNC is able to differentiate between the completely matched complementary target and mismatched target. In addition, there is no potential cross-reaction between two DNA targets as evidenced by signal magnitude to the background level after cross hybridization. Figure 3C shows the fluorescence intensity of fNC increases with increasing concentration of DNA target from 0 to 200 nM. This indicates that the fluorescence intensity is highly associated with the concentration of

the DNA target and could represent the target sequence content for sensitive determination of gene expression level. The *in vitro* detection limit of fNC to *Tubb3* and *Fox3* was 1.5 nM and 1.2 nM, respectively. Next, we assessed the hybridization kinetics of fNC to DNA targets and the stability of fNC upon nuclease exposure. Results showed that fNC responded to the target sequence in less than 20 s (Figure 3D). This extremely short time delay should not interfere with real-time imaging data analysis in cells. Moreover, the fNC was resistant to nuclease degradation, whereas the equivalent molecular beacon, widely used for subcellular imaging,¹⁰ underwent degradation over long-term incubation with DNase I (Figure 3E). The off-target gene detection was evaluated by incubating various DNA sequences with fNCs. Figure 3F shows high specificity of fNCs to *Tubb3* and *Fox3* among other off-target genes. As for the stability of the nanocomplex, we introduced this dithiol modification to increase the stability. For the DNA duplex dissociation, the data shown in Figure 3C demonstrated that the nanocomplex is sensitive to spiked complementary DNA oligos, and the hybridization rate is very fast (Figure 3D). However, if mismatched DNA oligo was mixed with the nanocomplex, the nanocomplex remained stable at spiked DNA oligo concentrations as high as 200 nM (Figure 3F). Hence, the nanocomplex presented high stability in various conditions. Collectively, our results show that fNC is a robust probe for *Tubb3* and *Fox3* mRNA pair detection.

Cytotoxicity of fNCs to Neural Stem Cells. Next, we evaluated the possible cytotoxicity of fNCs to neural stem cells by conducting an MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay to evaluate cell proliferation. No detectable cytotoxicity to NSCs at concentrations of 0.1–1 nM over 48 h incubation was found (Figure S5), suggesting that fNCs are biocompatible for imaging of intracellular NSC gene expression.

Evaluation of fNCs as mRNA Imaging Probes in Highly Differentiated NSCs. We then investigated the fNC's ability to detect intracellular mRNA in differentiated NSCs. NSCs were first stimulated by free RA (1 μ M) to differentiate into neural subtype cells over 8 days before incubation with fNCs. Figure 4A shows the intracellular *Tubb3* mRNA (green) and *Fox3* mRNA (red) distribution in single cells detected by fNCs after 4 h incubation. At day 8 after RA stimulation, NSCs are partially differentiated into neurons, in which β III tubulin (encoded by *Tubb3*) and NeuN (encoded by *Fox3*) proteins are coexpressed in cells.³⁸ This is consistent with our results in Figure 4A, showing the co-localization of *Tubb3* and *Fox3* mRNAs in the cytoplasm. In addition to the static imaging of mRNA expression, we then studied the kinetics of fNCs on detecting *Tubb3* and *Fox3* mRNAs in a well-differentiated NSC colony under identical conditions. Results (Figures 4B and S6)

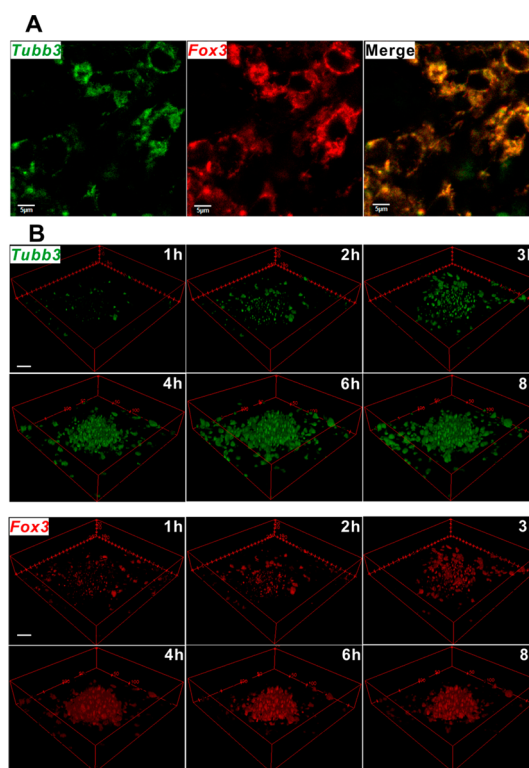


Figure 4. (A) Fluorescence imaging of *Tubb3* (green color) and *Fox3* (red color) mRNAs by fNCs at day 8 in differentiated NSCs stimulated by free RA in advance. Scale = 5 μ m. (B) Fluorescence recovery kinetics of fNCs in detecting *Tubb3* (green color) and *Fox3* (red color) mRNAs at day 8 in differentiated NSCs stimulated by free RA in advance. The images were reconstructed into 3D format without compromising the originality for better illustration. Scale = 20 μ m.

showed dominant *Tubb3* and *Fox3* mRNA expression levels at 4 h incubation with fNCs, and fluorescence intensity became almost saturated after 6 h.³⁹ In contrast, in the initial 3 h, minimal fluorescent signal could be detected, probably due to the relatively slow kinetics of fNC endocytosis and the endo/lysosome escape progress. The dark field imaging results (Figure S7) also confirmed this endocytosis progress. In addition, it was reported that the spherical DNA oligos on the gold nanoparticle surface undergo 3D structural rearrangement during cellular uptake. This spatial structure change supports the significant endocytosis and efficient intracellular release through a lipid-raft-dependent, caveolae-mediated pathway.⁴⁰ The mRNA imaging observed with fNCs in differentiated NSCs validated the time-dependent trend of endocytosis of engineered gold nanoparticles in cells²⁶ and confirmed the efficient mRNA detection by fNCs in NSCs.

Self-Activating and Spatiotemporal Imaging Dynamics of mRNA Sequential Expression during NSC Differentiation. We applied fNCs to spatiotemporally image the dynamics of mRNA sequential expression during NSC differentiation from neural stem cells to neurons in the self-activating condition. Results in Figure 5A and B show that fNCs could efficiently stimulate NSC differentiation

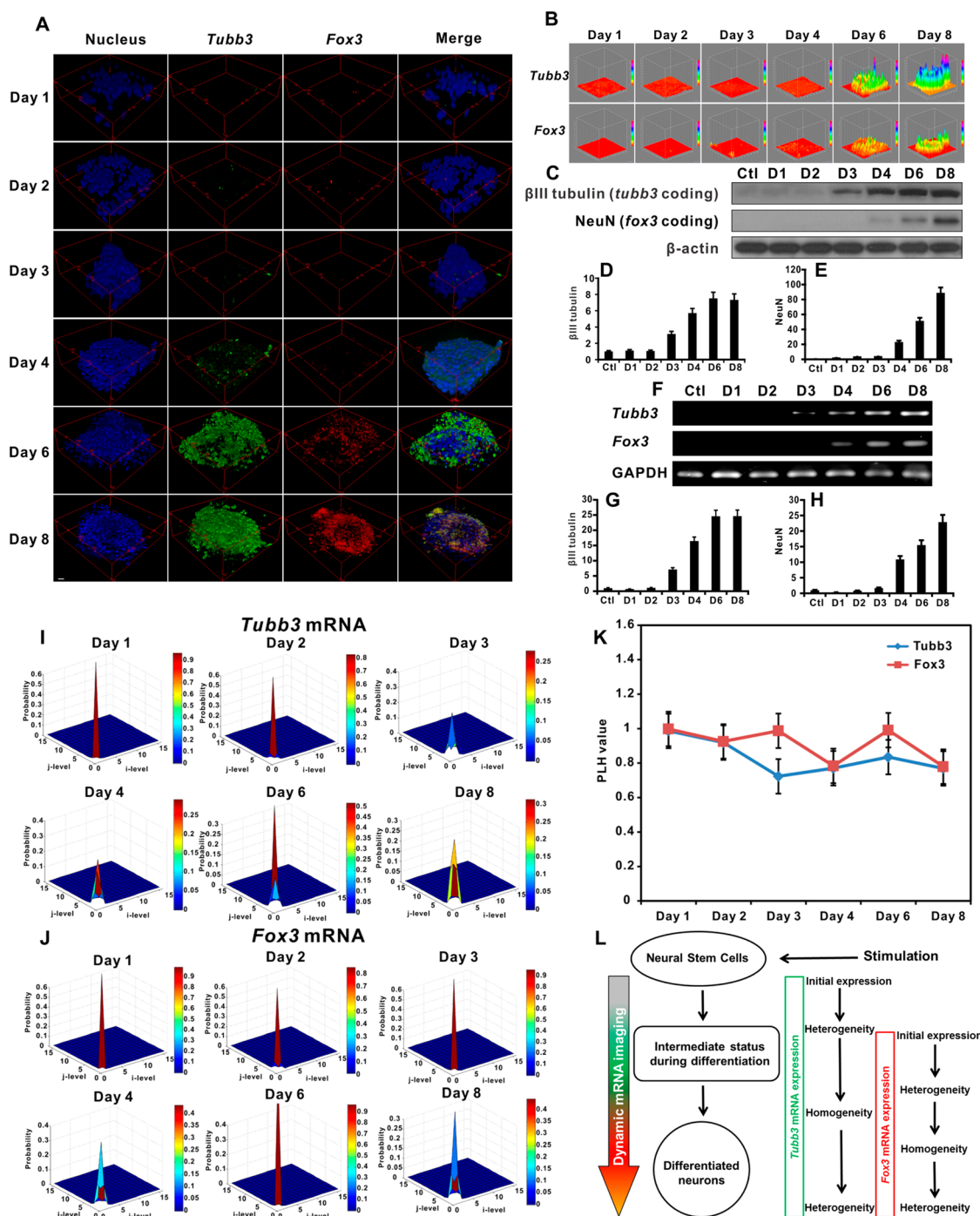


Figure 5. (A) Self-activating and spatiotemporal imaging of *Tubb3* and *Fox3* mRNA sequential expressions during NSC differentiation in living cells. Green: *Tubb3* mRNA; red: *Fox3* mRNA; blue: nuclei stained with Hoechst 33342. Scale = 20 μ m. (B) Hot maps of *Tubb3* and *Fox3* mRNA sequential expression. (C–E) Western blot analysis of *Tubb3* coding β III tubulin and *Fox3* coding NeuN protein expression levels at different NSC differentiation stages. (F–H) PCR analysis of *Tubb3* and *Fox3* mRNA expression levels at different NSC differentiation stages. (I) Texture feature heterogeneity analysis of *Tubb3* mRNA expression imaging. (J) Texture feature heterogeneity analysis of *Fox3* mRNA expression imaging. (K) Photolithographic hologram (PLH) value of mRNA heterogeneity analysis of *Tubb3* and *Fox3* expression ($n = 5$). (L) Mechanistic insights of successive mRNA expression initiation and dynamic imaging during neural stem cell differentiation and associated texture feature pattern progress of the heterogeneity of mRNA expression.

followed by sensitive and specific imaging of the dynamics of *Tubb3* and *Fox3* mRNA sequential expressions, respectively. From the imaging result, we observed that *Tubb3* mRNA expression is about 2 days earlier than *Fox3* mRNA expression, and both peak at

day 8. This imaging result is in line with the inherent functions of *Tubb3*, which encodes β III tubulin protein, an early marker of signal neuronal commitment,³² and *Fox3*, encoding NeuN, which is expressed in neurons.³³ The imaging results were further validated by Western

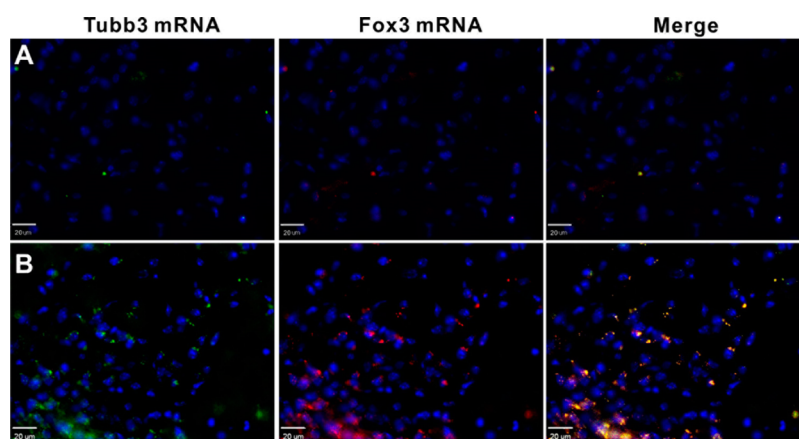


Figure 6. *In vivo* self-activation and imaging of endogenous neural stem cells by fNCs. (A) Images of nanocomplexes without RA-treated mouse brain. (B) Images of nanocomplexes with treated mouse brain. Blue: nucleus; green: *Tubb3* coding mRNA expression; red: *Fox3* coding mRNA expression. Scale bar = 20 μm .

blots for corresponding coding protein expressions (Figure 5C–E) and PCR analysis for mRNA levels (Figure 5F–H). Our data also demonstrated the enhanced differentiation stimulation efficiency of RA delivered by nanoscale vector.²³ In contrast, fNCs without the RA motif can hardly stimulate NSC differentiation, nor can they image any mRNA expression upon incubation over 8 days (Figure S8).

Texture Feature Analysis of the Dynamics of mRNA Expression Network. Dynamic gene expression imaging provides advantages over conventional static single-point imaging to provide a continuous overview of gene expression dynamics and facilitates analysis of gene expression progress in real time. To correlate the mRNA expression pattern network, we employed a texture feature analysis method⁴¹ to investigate the gene expression pattern obtained from fNC imaging during NSC differentiation (Figure 5I,J). In the pictorial data, the texture feature represents important information on the statistic distribution of gray-scale variation in a group of defined pixels.⁴¹ In our study, the texture feature analysis is able to specifically identify the spatial relationship of a single pixel with the adjacent pixels in the two-dimensional direction and generally describes the “roughness” and distribution pattern of the gene expression condition in a single colony. For both *Tubb3* and *Fox3* mRNAs, the homogeneous pattern (the overall uniform gene expression level of a cell colony or cluster derived from the almost identical gene expression level of individual cells in a specific time point) on the first day is attributed to the stem-like condition of NSCs, indicating neither gene was expressed at that stage. However, *Tubb3* expression became less homogeneous at day 2 due to the early onset expression of mRNA. The homogeneity increased gradually in the following 2 days, as more nascent NSCs initiated the onset of *Tubb3* expression. Heterogeneous (the overall nonuniform gene expression level of a cell colony or cluster derived from the

distinctive gene expression level of individual cells at a specific time point) colonial cell differentiation at day 4 and after led to the decline of *Tubb3* heterogeneity (Figure 5K and L). *Fox3* mRNA showed a similar dynamic expression texture pattern but delayed for 2–3 days (Figure 5K and L). These imaging and texture pattern analysis results are in high agreement with observed growth patterns during stem cell differentiation.⁴²

***In Vivo* Self-Activation and Imaging of Endogenous Neural Stem Cells by Nanocomplex.** We stereotactically injected the fNC directly into the lateral ventricle of the mouse brain to investigate the *in vivo* behavior of the nanocomplex. Figure 6 shows the parallel comparison of endogenous NSC activation and labeling effects post fNC injection. fNC without RA could not extensively activate NSC differentiation, resulting in a sporadic imaging signal distribution (Figure 6A). In contrast, fNC led to strong fluorescent signals for both *Tubb3* and *Fox3* mRNAs (Figure 6B). The area adjacent to the lateral ventricle showed little, if any, detectable fluorescent signal (Figure S9). This negligible fluorescent signal may come from the tissue background noise or the diffusion of fNCs taken up by mature neurons. Overall, our results clearly demonstrate that (1) the synthesized nanocomplex is capable of activating endogenous NSC by the cellular machinery triggered RA release, followed by efficient mRNA imaging; (2) the attachment of RA on the distal end of nanoparticles is critical to regulate NSC differentiation ability (Figure 6A). Importantly, the required RA concentration is much lower (15 nM) than the commonly used 1 μM to regulate NSC differentiation in animals. The *in vivo* experiment confirmed the high efficiency of nanocomplexes for simultaneous gene regulation and imaging. Unfortunately, we could not perform noninvasive real-time imaging *in vivo* due to the limitation of microscopic light penetration depth to the lateral ventricle in the brain. New techniques implanting microfibers into the deep compartment of the mouse brain would help

the spatiotemporal imaging of the mRNA expression dynamics of endogenous differentiating neural stem cells in the presence of the fNCs.

CONCLUSION

In summary, we have developed a multifunctional nanocomplex integrating stimulating motif self-releasing capacity and dynamic mRNA expression imaging function. Our results demonstrated the high stability of fNCs to enzymatic degradation, the high specificity to target genes, and potent self-cleavage RA delivery to stimulate NSC differentiation. Moreover, fNCs enabled the spatiotemporal imaging of the dynamics of mRNA sequential expression in the complicated cascades of differentiating NSCs in a self-activating manner. In comparison to static mRNA imaging techniques at a

single time point, fNCs provide a temporal tool for evaluating mRNA expression that can enhance our understanding of genetic regulation processes. Importantly, this conceptual fNC platform can be easily reprogrammed by amending corresponding oligonucleotide sequences and stimulating motifs to detect other sophisticated biological events, such as embryo development and cancer progression. We envision that this robust multifunctional nanocomplex would provide an alternative opportunity to explore dynamics of complicated biological events in living cells. An inherent limitation of this nanocomplex design is that it cannot image the downregulation of interested genes during neural stem cell differentiation, even though it demonstrated superior imaging capacity of gene upregulation in this report.

MATERIALS AND METHODS

1. Synthesis of Nanocomplexes. Citrate-modified gold colloidal nanoparticles (15 ± 0.7 nm) were synthesized using published procedures.⁴³ Two dithiolated DNA oligonucleotides were mixed separately with corresponding reporter strands at a molar ratio of 1:1.2 in PBS buffer, heated to 75 °C, and maintained for 30 min, then slowly cooled to room temperature and stored in the dark to allow hybridization. These preformed DNA duplexes were mixed together and added to a gold colloid solution (3 nM) at final concentrations of 540 nM of Alexa 488–*Tubb3* DNA duplex and 560 nM of Cy3–*Fox3* DNA duplex, then shaken overnight. After 12 h, phosphate buffer saline (10×; pH = 7.4) was added to the mixture to bring the NaCl concentration gradually to 0.3 M during a period of 8 h. Then the resulting nanocomplexes were centrifuged at 12 000 rpm for 15 min three times and were redispersed in PBS buffer. Particles were sterilized using a 0.2 μm acetate syringe filter (GE Healthcare). (PBS buffer components used in this paper: 10 mM phosphate, pH 7.4, 138 mM NaCl, 2.7 mM KCl, and 2 mM MgCl₂.)

2. Synthesis of Multifunctional Nanocomplexes with Retinal. The resulting nanocomplexes of 0.05 nmol in 1× PBS (pH 7.4) were mixed with 10 nmol of retinal in DMSO and shaken overnight in the dark. Then the resulting fNCs were purified using centrifugation (12 000 rpm, 15 min, three times) and were redispersed in PBS buffer.

3. DNA Loading Determination. The concentration of fNCs was determined by using UV–vis absorption spectroscopy ($\lambda = 524$ nm, $\epsilon = 3.64 \times 10^8$ M⁻¹ cm⁻¹).⁴⁴ DNA oligonucleotides were set free from fNCs by dissolving their gold backbones in a 0.1 M KCN solution. The concentration of DNA duplexes was determined by fluorescence measurements and comparing to a standard curve (Figure S2). The loading number of DNA oligonucleotides was calculated by dividing the concentration of dye-containing oligonucleotides by the concentration of gold colloidal nanoparticles.

4. Hybridization Experiment. For multiplex analysis detection, the nanocomplex of 1 nM in PBS buffer was treated with the *Tubb3* targets (single-base mismatched or complementary DNA targets) or *Fox3* targets (single-base mismatched or complementary DNA targets) at various concentrations of 0, 2, 5, 10, 25, 50, 100, and 200 nM. After incubation for 1 h at 37 °C, the fluorescence was observed at appropriate excitation wavelengths. The fluorescence of Alexa 488 was excited at 490 nm and measured at 530 nm (range from 500 to 700 nm), and the fluorescence of Cy3 was excited at 550 nm and measured at 570 nm (range from 550 to 700 nm). The nanocomplex had a linear range from 0 to 25 nM, and the limit of detection was calculated by the derived calibration curve, as defined by IUPAC. All experiments were repeated at least three times.

5. Specificity Experiments. The completely complementary DNA targets for designed nanocomplexes and other cancer marker targets as controls were spiked in 1 mL of buffer containing 1 nM of nanocomplex, while the DNA targets were 200 nM. All experiments were repeated at least three times.

6. Nuclease Assay. Nanocomplexes were diluted to a concentration of 1 nM in buffer (10 mM PBS, pH 7.4, 2 mM MgCl₂, 1 mM CaCl₂, and 50 mg/L bovine serum albumin). After allowing the samples to equilibrate for 10 min, DNase I (Amplified grade, Invitrogen) with a final concentration of 50 U/L was added immediately before reading. Molecular beacons were tested in an analogous manner at a concentration of 25 nM. The fluorescence of these samples was monitored for 24 h and was collected at predetermined intervals during this time period.

7. Cytotoxicity of fNCs. The cytotoxicity of fNCs to neural stem cells was evaluated by the MTT assay and cell proliferation curve. In the MTT assay, 100 μL of complete culture medium containing 5000 NSCs was cultured in a 96-well plate for 24 h before experiments. Various concentrations of fNCs (with or without RA) were incubated with cells for 48 h. The medium was replaced with 100 μL of fresh media containing 10 μL of MTT solution (5 mg/mL), followed by incubation for another 4 h. The media was then removed, and 100 μL of dimethyl sulfoxide (DMSO) was added into each well to dissolve the formed formazan crystals. The absorbance at 570 nm was recorded using a microplate reader. The absorbance from the control cells was set as 100% cell viability ($n = 6$). For the cell proliferation curve, 0.25 million cells were initially cultured in a 24-well plate in complete medium with various concentrations of fNCs (without RA). The medium was changed every other day until day 8. The cell number in each well was counted by cytometer every day ($n = 4$).

8. Characterization of Nanocomplexes Using TEM, UV Spectra, and DLS. Transmission electron microscopy (TEM) measurements were carried out on a JEOL 200X operated at 200 kV. UV–vis spectra were measured using a Shimadzu UV1701. Dynamic light scattering analysis was performed using a Zetasizer Nano ZS (Malvern Instruments Ltd., UK).

9. In Vitro RA Release of fNCs. The prepared fNC (5 nM final concentration, diluted from a 40 nM stock solution in corresponding buffer into a final volume of 200 μL) was incubated in either PBS buffer (pH = 7) or sodium acetate buffer (pH = 5) for a predetermined time (30 min or 12 h), followed by ultracentrifugation at 14 000 rpm for 30 min to precipitate DNA oligo attached gold nanoparticles. The supernatants containing released RA or derivatives were harvested for HPLC analysis. For the HPLC analysis, we used a gradient mobile phase with water/acetonitrile (20%/80%–10%/90%) over 12 min at 360 nm UV detection wavelength.

10. Quenching Effect of AuNPs of Fluorophores Conjugated on Oligos. In each experiment, different recognizing DNA-modified gold nanoparticles were added gradually to 50 nM of corresponding reporter in PBS buffer. The fluorescence signal was scanned by a fluorescence spectrometer. The quenching effect was calculated as the following equation: QE = quenched fluorescence/initial fluorescence \times 100%. The quenched fluorescence represents the remaining fluorescence from the initial recorded fluorescence signal minus the fluorescence after addition of DNA-Au NPs. When 1 nM of DNA-Au NPs was used, the calculated quenching efficiency of gold nanoparticles was 92% for Alexa 488 and 87% for Cy3. This quenching effect difference of AuNPs is attributed to the various absorption spectral overlap of gold nanoparticle with distinct emission spectra of different fluorophores.

11. Real-Time Imaging and Kinetics of fNCs without RA on *Tubb3* and *Fox3* mRNAs in Differentiated NSCs at Day 8. Mouse neuroectodermal neural stem cells (NE-4C, ATCC CRL-2925) were cultured in eight-well Lab-Tech Chamber plates (NUNC, Thermo Scientific) for 24 h prior to experiment. The next day, free retinoic acid dissolved in DMSO was diluted into 1 μ M working solution with complete medium. The cell culture medium was changed to the freshly prepared free-RA-containing medium for 48 h incubation with NSCs. Then, fresh medium without RA was changed every other day until day 8. On day 8, an fNC stock solution was diluted with complete medium to a 0.5 nM working solution before incubation with differentiated NSCs. The kinetics of fNC to image mRNA targets in differentiated NSCs was obtained by continuously imaging a single cell colony over 8 at 1 h intervals by laser scanning confocal microscopy (LSFM) (Olympus, FV10). Single living cell imaging results were obtained under similar conditions, while the imaging focus was set to the periphery of the cell colony with high magnification to ensure single-cell resolution. The 3D imaging results were achieved by postprocessing with Matlab software (Cambridge, MA, USA) and reconstructed with ImageJ software (NIH).

12. Self-Activating and Spatiotemporal Imaging of mRNA Sequential Expression Dynamics during NSC Differentiation. Mouse neuroectodermal neural stem cells were cultured in eight-well Lab-Tech Chamber plates (NUNC, Thermo Scientific) for 24 h prior to experiments. A fNC of 0.5 nM was used in this experiment. The stock solution of the fNC was diluted with complete medium and incubated with NSCs. The first day after incubation, 0.5 μ L of Hoechst 33342 stock solution was added into each well for living cell nucleus staining 30 min prior to imaging. A marker was made on the chamber to ensure the consistently same cell colony imaging over 8 days. On the second day prior to imaging, cells were washed with HBSS solution (pH = 7.4) twice, followed by changing the fresh medium. The culture medium was changed every other day until day 8, and imaging was performed every day. Hoechst 33342 was added whenever necessary to illustrate the cell nucleus and cell colony location. For the negative control experiment, 0.5 nM of fNC without the RA motif was incubated with NSCs and imaged over 8 days using the same above protocol. The 3D imaging results were achieved by postprocessing with Matlab software and reconstructed with ImageJ software.

13. Western Blot Study of β III Tubulin and NeuN for Imaging Result Validation. NSCs were incubated with 0.5 nM fNCs at predetermined time points. Then, cells were washed with ice cold PBS (calcium and magnesium free) three times before being treated with RIPA buffer (Pierce) with protease inhibitor cocktail (Roche) for 10 min on ice. Cell lysates were harvested and subjected to centrifugation at 13 000 rpm at 4 $^{\circ}$ C for 10 min. The resultant supernatants containing protein were harvested, followed by 4–12% gradient SDS-PAGE gel separation. Nitrocellulose membranes were used for transferring separated protein bands. The membranes were blocked by 5% skim milk for 1 h at room temperature and then immunoblotted with anti β III tubulin (1:1000) and anti NeuN (1:1000) primary monoclonal antibodies (Cell Signaling Technology, Inc.). HRP conjugated goat anti-rabbit secondary antibody (1:3000) was used to visualize immunoreactive bands. Anti β -actin rabbit mAb (Cell Signaling Technology, Inc.) (1:2000) was used as the housekeeping control.

14. qPCR Analysis of *Tubb3* and *Fox3* mRNA Expression Levels during NSC Differentiation. NSCs were incubated with 0.5 nM fNCs at predetermined time points. Total RNAs were extracted from treated NSCs using Tripure isolation reagent (Roche). RNA (1 μ g) was reversely transcribed into cDNAs using SuperScript First Strand Synthesis kit (Invitrogen), followed by quantitative real-time PCR analysis using a SYBR Green Master Mix kit (Applied Biosciences) on a StepOne Plus real-time PCR system (Applied Biosystems, Foster City, CA, USA). The primers used in this study are *Tubb3*: 5'-CTTTCTGCTCTAGCCGCGT-3' (forward); 5'-CTCATCGCTGATGACCTCC-3' (reverse). *Fox3*: 5'-GGCAATGGTGGGACTCAAAA-3' (forward); 5'-GGGACCCGCTCCTTCAAC-3' (reverse). *GAPDH* was used as the housekeeping control: 5'-GGTCTCCTCTGACTTCAACA-3' (forward); 5'-AGCCAAATTCGTTGCATAC-3' (reverse).

15. In Vivo Animal Model. All experiments were performed in accordance with NIH guidelines for the care and use of laboratory animals. Adult Balb/c mice (7–8 weeks) were anesthetized under 2% isoflurane, and the brain was placed on a stereotaxic apparatus. The following stereotaxic coordination parameters were used for the direct injection of nanocomplex (5 μ L, 10 nM) into the lateral ventricle: rostral: 0.5 mm, lateral: 0.75 mm, ventral: 2.5 mm. After completing injection, the micropipette was kept for 5 min before being withdrawn. Mice were placed on a warm pad until awakened. At a predetermined time, mice were perfused using 3.7% polyformaldehyde (20 mL) before being sacrificed. Brains were dissected and sliced into 15 μ m tissue slides for imaging using fluorescence microscopy.

Conflict of Interest: The authors declare no competing financial interest.

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Supporting Information Available: This material is available free of charge via the Internet at <http://pubs.acs.org>.

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