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Self-assembling nano-probes displaying off/on 19F NMR signal for protein detection and imaging

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ABSTRACT (150 words)
Magnetic resonance imaging (MRI) is one of the most promising techniques for non-invasive visualization of biomarkers and biologically relevant species both in vivo and ex vivo. Although $^1$H MRI with paramagnetic contrast agents, such as Gd$^{3+}$ complexes and iron oxide, is widely used, it often suffers from low contrast due to the large background signals from the abundant distribution of protons in biological samples. Here we report supramolecular organic nanoparticles to detect specific proteins by $^{19}$F-based MRI in a perfect off/on mode. The designed probes are NMR-silent when aggregated but, in the presence of a target protein, are disassembled to produce a sharp signal. This "turn-on" response allowed us to clearly visualize the proteins inside live cells by $^{19}$F MRI and construct an in-cell inhibitor assay. This recognition-driven disassembly of nano-probes for turn-on $^{19}$F-signal is unprecedented and this strategy may extend the utility of $^{19}$F MRI for specific protein imaging. (150 words)

MAIN TEXT (2,000 words)
MRI is superior to optical bio-imaging in living systems for the visualization of deep tissues. Currently, $^1$H MRI is widely used for diagnostic purposes because of its high sensitivity. This high sensitivity is attributed to the abundance of water molecules in the vicinity of contrast agents. However, at the same time, $^1$H MRI often suffers from low contrast-to-noise ratio due to the large background signals from water protons. Therefore, for specific imaging of biomarkers and biologically relevant molecules with higher functional and/or spatial resolution in vivo as well as ex vivo, there is still an obvious need to develop new methodologies. Several approaches are now proposed. In particular, $^{19}$F holds great promise as an alternative nuclide for MRI as it has a high NMR sensitivity next to $^1$H (83% relative to $^1$H) and 100% natural abundance. A more important advantage of $^{19}$F is that essentially no NMR-detectable $^{19}$F is present in animal bodies, eliminating the interference from background signals. As such, when a $^{19}$F-containing probe is applied to a biological sample, only the extrinsic signal from the molecule can be detected. However, $^{19}$F MRI technology is still in its infancy. Despite significant importance for medical diagnosis, strategies for imaging specific proteins/enzymes with high MRI contrast are very limited. To date, targeting or
switching probes for $^{19}$F NMR/MRI detection have been proposed. The former is based on the accumulation of probes in specific regions in tissues through binding to localized components,\textsuperscript{9} while the latter provides signals that can be modulated by enzymatic reaction. Two types of switching $^{19}$F MRI probes reported were enzyme substrates: one displays chemical shift changes\textsuperscript{10} and the other shows a paramagnetic relaxation enhancement (PRE)-dependent signal turn-on, upon enzymatic cleavages.\textsuperscript{11} The switching of these probes relies entirely on the catalytic activity of the enzymes, and thus the probes are spatially diffused away from the target enzyme and also not applicable for non-enzymatic protein targets. Therefore, more universal MRI strategies for protein detection and imaging that do not rely on enzymatic activities are highly desirable.

Here we describe a novel strategy to detect specific proteins with an ‘off/on-type’ $^{19}$F NMR signal using dynamic self-assembled nano-particles. Our idea is based on that the $^{19}$F NMR signal is broadened and attenuated when the molecules assemble into high-molecular weight aggregates, but recovers upon their disassembly.\textsuperscript{2} Because $^{19}$F has a relatively large chemical shift anisotropy (CSA; approximately 39 ppm for $-\text{CF}_3$ group),\textsuperscript{12} the transverse relaxation of $^{19}$F NMR signals is extremely sensitive to the apparent molecular weight, due to the CSA relaxation mechanism. Thus, formation of a large molecular assembly can cause severe broadening of the $^{19}$F NMR signal. Accordingly, small molecule probes were designed which are composed of a $^{19}$F-containing group and a ligand specific to a protein of interest, and concurrently equipped with the ability to form nano-aggregates. The probe alone is NMR-silent due to its self-assembly, but gives a distinct $^{19}$F signal in response to the target protein through the binding-mediated disassembly of the probe (Fig. 1a). Since the signal response is determined by specific protein-ligand interactions, this principle should be applicable to detect the presence of certain proteins including both enzymes and non-enzymatic proteins.

For the proof-of-principle experiment, we initially chose human carbonic anhydrase I (hCAI) as a target protein. Probe 1 was synthesized which contains a 3,5-bis(trifluoromethyl)benzene (FB) derivative carrying six magnetically equivalent $^{19}$F nuclei connected to a benzenesulphonamide moiety, a ligand specific for hCAI (dissociation constant $K_d = \text{ca. } 3 \, \mu\text{M}$),\textsuperscript{13} via a relatively hydrophobic linker (Fig. 1b and Supplementary Methods). When 1 was dissolved in a buffer solution containing
trifluoroacetic acid (TFA; internal standard at -75.6 ppm), no $^{19}$F signal was observed. However, a sharp signal appeared at -62.6 ppm upon addition of hCAI (Fig. 1d). The signal intensity increased linearly in proportion to the concentration of hCAI, and was saturated at a 1:1 molar ratio of probe 1 and hCAI (Fig. 1e). By the subsequent addition of a strong inhibitor, EZA (ethoxzolamide, 2; Fig. 1c), into the above solution, the signal disappeared again (Supplementary Fig. 1a). These data indicate that 1) probe 1 alone is NMR silent, 2) the appeared $^{19}$F signal can be assigned to the probe 1 bound to the ligand-binding pocket of hCAI, and 3) the signal can be turned off in a reversible manner when the probe is expelled from the protein (that is, when the target protein is incapable of binding the probe). We next examined the target-specificity of probe 1 under miscellaneous conditions. Addition of 1 to a mixture of four proteins different from hCAI (hemoglobin, bovine serum albumin, concanavaline A, and chymotrypsin) did not give any sharp signal. On the other hand, the $^{19}$F signal was clearly observed in a mixture containing the four proteins and hCAI (Supplementary Fig. 1c). We also found that the hCAI-induced appearance of the $^{19}$F signal took place in 70% fetal bovine serum solution which contains many biological substances including proteins, lipids, and small molecules (Supplementary Fig. 1d). Overall, these results clearly demonstrate that probe 1 can selectively detect hCAI by turned-on $^{19}$F signal even in crude samples.

The self-assembly/disassembly properties of probe 1 were investigated by various measurements. Spherical or oval aggregates of 1 with a size ranging from 200 to 500 nm in diameter were observed by atomic force microscopy (AFM) (Fig. 2a). Transmission electron microscopy (TEM) and scanning electron microscopy (SEM) data supported the formation of aggregates of approximately 200 nm in size (Fig. 2b and Supplementary Fig. 2a). The UV-visible absorption spectrum of 1 in buffer solution showed a broad visible light scattering around 500–700 nm due to the aggregates of 1 (Fig. 2c and Supplementary Fig. 2b). The scattering was decreased 10-fold by the addition of hCAI, indicating that the aggregates were efficiently collapsed by hCAI. The aggregates with a mean size of 250 nm in diameter were consistently shown by dynamic light scattering (DLS) measurements in buffer solution containing 1 alone (Fig. 2d), whereas negligible DLS intensity was obtained after adding hCAI to the above solution. The molecular weight (Mw) of the nano-aggregate of 1 roughly estimated by the observed diameter is $10^7$ Da, while the Mw of the
complex between hCAI and probe 1 is $3 \times 10^4$ Da. We thus conclude that the binding of
probe 1 to hCAI induces the disassembly of the nano-aggregates, which dramatically
decreases the apparent Mw. This decrease effectively reduces the $^{19}$F relaxation rate, so
that a sharp $^{19}$F signal can be observed. By the concentration dependence of DLS
measurement, the critical aggregation concentration (CAC) of the self-assembly of 1
was revealed to be $< 5 \mu$M (Supplementary Fig.2c).

On the basis of this principle, we next produced a turn-on $^{19}$F probe 3 for trypsin
(TPS). Instead of benzenesulfonamide, benzamidin, a typical inhibitor for TPS ($K_d = \text{ca.}$
20 $\mu$M)$^{15}$ was linked to the FB moiety as a suitable ligand. Almost no NMR signal was
detected in buffer solution containing probe 3 alone, whereas a new signal was
intensified upon addition of TPS (Fig. 3a). In the presence of benzamidin (4, Fig. 1c),
such signal intensification did not occur. Similarly, biotin-tethered probe 5 showed a
clear off/on $^{19}$F NMR response to a non-enzymatic protein, avidin ($K_d$ of biotin, ca. $10^{15}$
M)$^{16}$ (Fig. 3b). Furthermore, the orthogonality of probe 1 and 5 was investigated. It was
shown that 1 responded to hCAI but not to avidin, and vice versa for 5 (Supplementary
Fig.3). These data demonstrate the general applicability of the present strategy for
designing turn-on supramolecular nano-probes to detect target proteins by $^{19}$F NMR
spectroscopy.

Notably, probe 1 was also capable of detecting hCA within live cells. hCA is a
cytosolic protein and naturally expressed inside human red blood cells (RBCs) (at a
concentration of approximately 170 $\mu$M)$^{17}$ A suspension of RBCs was incubated with 1
for a few minutes, and after collecting cells, in cell $^{19}$F NMR analysis was conducted.
Note that no hemolysis occurred during the experiments. A signal was clearly observed
at -62.6 ppm (Fig. 4a, top), with a chemical shift of which is identical to that obtained
using purified hCAI and 1 (Fig. 1d), although there was a slight peak broadening. In
contrast, no signal appeared when 1 was incubated with RBCs in the presence of EZA
(Fig. 4a, bottom). Clearly, probe 1 is cell-permeable and can specifically detect
endogenous hCA with turn-on signal response even inside cells.

The perfect turn-on probe 1 allowed us to visualize the target protein using a $^{19}$F
MRI phantom. At first, $^1$H and $^{19}$F MR images were acquired after mixing hCAI with
probe 1 under in test tube conditions in the absence (sample 1) and presence of the
inhibitor, EZA (sample 2) or SBA (4-sulfamoylbenzoic acid, 7, Fig. 1c)$^{13}$ (sample 3). In
$^1$H MRI, all of the three samples gave indistinguishable images (Fig. 4b, top). On the
other hand, a distinct $^{19}$F MR image was obtained from sample 1, whereas no MRI signal was detected in sample 2 and 3 (Fig. 4b, bottom). These results are consistent with the $^{19}$F NMR data described above (Fig. 1d and Supplementary Fig. 1a). More significantly, a clear $^{19}$F MR image was observed in sample 4 containing RBCs and probe 1 (Fig. 4c). The MR image was completely diminished by co-incubation of EZA (sample 5), confirming that the positive MR image obtained above is indeed due to probe 1 bound to hCAI inside cells. Interestingly, sample 6 containing RBCs, probe 1 and SBA still gave a MR image similar to that of sample 4 despite the presence of the inhibitor. Given that the SBA efficiently blocks the binding of probe 1 to hCAI under in test tube conditions (Fig. 4b, bottom), this result is reasonably ascribed to the less cell-membrane permeability of anionic SBA compared to neutral EZA. Therefore, the present system serves as a cell-based inhibitor screening platform which can visualize the potency of inhibitors in cellular contexts.

In conclusion, we have developed the supramolecular $^{19}$F-containing nano-probes which can detect specific proteins spatially by the $^{19}$F MRI technique with sharp turn-on-type switching. The simple principle for the off/on response, i.e. self-assembly and recognition-driven disassembly of the nano-probe, should be applicable to the design of other turn-on probes for various target proteins by appropriately replacing the ligand module, as shown in the case of TPS probe 3 and avidin probe 5. It should also be noted that, in contrast to existing MRI contrast agents, this method does not require any metals that have potential toxicity. In the practical viewpoint, it may be fair to consider that the sensitivity of $^{19}$F is considerably low compared to $^1$H MRI and thus the present system is not yet sufficient for in vivo or clinical application using conventional bench-top spectrometers (which generally have strengths of ~1 tesla). Functional $^{19}$F probes are also very limited in number. However, we believe that with further advances in both instrumentation and chemistry, $^{19}$F-based NMR/MRI technique may become a more powerful modality for practical diagnosis as well as basic research in the future. It is essential for chemists to establish general and useful design concepts for new $^{19}$F probes that facilitate functional and molecular MRI. (1854 words)

METHODS (800 words)

$^{19}$F NMR analysis of probe 1 with various concentrations of hCAI
Human carbonic anhydrase I (hCAI, 5.0 mg) was dissolved in 50 mM HEPES buffer (1.0 mL, pH 7.2, 10% D$_2$O (v/v), 0.2 mM TFA). The concentration of hCAI was determined by absorbance at 280 nm using the molar extinction coefficient (49 000 M$^{-1}$cm$^{-1}$)$^{19}$ and 100 µM stock solution was prepared. Probe 1 (3.0 mg, 3.8 µmol) was dissolved in 75 µL of DMSO as the stock solution, and slowly added to the hCAI solution (0.6% DMSO (v/v)). These samples were analyzed by $^{19}$F NMR with TFA as an internal standard (-75.6 ppm).

**AFM, TEM and SEM observations**

AFM imaging; a solution of probe 1 was spin coated onto a freshly cleaved mica surface and dried in vacuo. The sample was imaged by a tapping-mode AFM on a SEIKO SPA-400. TEM imaging; a solution of probe 1 was deposited on a thin carbon support film and dried in vacuo. The sample was imaged by a JEOL JEM-1025, operating at 100 kV, without any contrast agent. SEM imaging; a solution of probe 1 was deposited on a silicon wafer and dried in vacuo. The sample was imaged by a JEOL JFC-1600, operating at 15 kV, with the addition of platinum spray as a conductive material.

**Measurements of optical density and dynamic light scattering**

The optical density was measured at 25°C in 50 mM HEPES buffer (pH 7.2, 0.2 mM TFA) using a quartz cell (1 cm). The DMSO stock solution of probe 1 was slowly added to the buffer solution (0.6% DMSO (v/v)) to be 25 µM. The measurements of dynamic light scattering were performed in the same conditions using a tubular-type cell. All measurements were performed in triplicate.

**$^{19}$F NMR analyses of probe 3 and probe 5**

Trypsin (TPS, 5.0 mg) was dissolved in 50 mM Tris-HCl buffer (1.0 mL, pH 8.5, 300 mM NaCl, 0.2 mM TFA, 10% D$_2$O (v/v)). The concentration of TPS was determined by absorbance at 280 nm using the molar extinction coefficient (36 700 M$^{-1}$cm$^{-1}$)$^{20}$ and 100 µM stock solution was prepared. The probe 3 (1.0 mg, 1.3 µmol) was dissolved in 26 µL of DMSO and 26 µL of 50 mM Tris-HCl buffer (pH 8.5) as the stock solution (25 mM), and slowly added to the TPS solution (0.2% DMSO (v/v)). Avidin (5.0 mg) was dissolved in 50 mM HEPES buffer (1.0 mL, pH 7.2, 500 mM NaCl,
0.2 mM TFA, 10% D₂O (v/v)). The concentration of avidin was determined by absorbance at 280 nm using the molar extinction coefficient (35 700 M⁻¹cm⁻¹)²¹ and 100 μM stock solution was prepared. The probe 5 (3.0 mg, 3.8 μmol) was dissolved in 72 μL of DMSO as the stock solution, and slowly added to the avidin solution (0.6% DMSO (v/v)). Conditions for ¹⁹F NMR measurements were same as probe 1.

**HCA Inhibitor assays in RBCs**

A 2 mL solution of probe 1 (200 μM) and inhibitor (EZA or SBA, 0 or 1 mM) in HBS buffer (20 mM HEPES, 107 mM NaCl, 6 mM KCl, 1.2 mM MgSO₄, 2 mM CaCl₂, 11.5 mM glucose, pH 7.4) was added to 2 mL of sedimented red blood cells (RBCs), and the suspension was incubated at room temperature for a few minutes. After centrifugation (1,500 rpm × 5 min), the supernatant was removed and re-suspended in buffer for ¹⁹F NMR measurement (50 mM HEPES buffer (pH 7.2), 100 mM NaCl, 0.17 mM TFA, 20% D₂O (v/v)). 0.7 mL and 2.5 mL of the suspension were used for ¹⁹F NMR and MRI measurement at 25°C, respectively.

**¹H and ¹⁹F MRI in test tube or in RBCs**

¹H images of sample 1-3 (in test tube) were obtained by gradient echo with repetition time (TR)/echo time (TE) = 100/6 ms, flip angle = 30, field of view (FOV) = 16 x 4 cm, slice thickness = 5 mm, matrix size = 256 x 256 and 1 average. ¹⁹F images of sample 1-3 (in test tube) were obtained by fast spin echo with TR/TE = 1500/5.5 ms, echo train length = 32, FOV = 16 x 4 cm without slice selection, matrix size = 128 x 32, the depth of sample tube was 20 mm, voxel size was approximately 31 mm³, and 1200 averages. ¹⁹F images of sample 4-6 (in RBCs) were obtained by gradient spin echo with TR/TE = 1000/2.4 ms, flip angle = 90, FOV = 32 x 8 cm without slice selection, matrix size = 128 x 32, the depth of sample tube was 30 mm, voxel size was approximately 188 mm³, and 400 averages. The sine window function was applied to the ¹⁹F images. All the images were acquired at 25°C. (787 words)
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AUTHOR CONTRIBUTIONS

I.H. conceived of the project. Y.T., T.S., S.T. and I.H. designed the experiments. Y.T. performed all the experiments, with help from H.T. and M.S. on $^{19}$F NMR measurements. M.N. and T.M. performed the MRI experiments. The manuscript was written by Y.T., S.T. and I.H., with editing by all the co-authors.
REFERENCES

1 Figures:

2 Figure 1. The turn-on-type $^{19}$F NMR probes for protein imaging. (a) Schematic illustration of the present strategy for an off/on switching mode of $^{19}$F NMR. (b) Chemical structures of probe 1, 3 and 5 for hCAI, TPS and avidin, respectively. (c) Chemical structures of inhibitors. For hCAI, EZA (ethoxzolamide, 2) and SBA (4-sulfamoylbenzoic acid, 7); for TPS, Benzamidin (4); for avidin, Biotin(+) (6). (d) Turn-on $^{19}$F NMR signal of probe 1 (50 μM) in the presence (top) or absence (bottom) of hCAI (50 μM). The signal to noise ratio (SNR) was 60.1. (e) Dependence of the $^{19}$F-signal intensity (-62.6 ppm) on the hCAI concentration: probe 1 (50 μM) in 50 mM HEPES buffer (pH 7.2, 0.2 mM TFA as an internal standard for peak intensity and chemical shift, 10% D$_2$O (v/v)) at 25°C.
Figure 2. Microscopic and spectroscopic characterization of the self-assembled nano-particles of probe 1. (a) AFM image of the self-assembled probe 1 (25 μM). The scale bar is 500 nm. (b) TEM image of the self-assembled probe 1 (25 μM). The scale bar is 500 nm. (c) Optical density at 600 nm of aqueous solution containing probe 1 (25 μM) in the absence (left) or presence (right) of hCAI (25 μM). Experiments were performed in triplicate to obtain mean and standard deviation values (shown as error bars). (d) DLS analysis of particle size distribution of the self-assembled probe 1 (25 μM). All experiments were performed in 50 mM HEPES buffer (pH 7.2, 0.2 mM TFA).
Figure 3. Turn-on $^{19}$F NMR detection of trypsin and avidin by probe 3 and 5. (a) $^{19}$F NMR spectra of 3 (100 µM) in the absence or presence of TPS (50 µM) and benzamidin (500 µM) in 50 mM Tris-HCl buffer (pH 8.5, 0.2 mM TFA, 300 mM NaCl, 10% D$_2$O (v/v)). The SNR was 12.2. (b) $^{19}$F NMR spectra of 5 (100 µM) in the absence or presence of avidin (50 µM) and biotin (500 µM) in 50 mM HEPES buffer (pH 7.2, 0.2 mM TFA, 500 mM NaCl, 10% D$_2$O (v/v)). The SNR was 57.3.
Figure 4. $^{19}$F NMR spectra and MR images in red blood cells (RBCs). (a) $^{19}$F NMR spectra of probe 1 in the absence (top) or presence (bottom) of EZA in RBCs. The SNR was 14.2. (b) MR images of probe 1 in test tube (top; $^1$H, bottom; $^{19}$F). (c) MR images of probe 1 in RBCs. Conditions: For in test tube (sample 1-3); 1 (100 μM), hCAI (50 μM), inhibitors (EZA or SBA) (0 or 500 μM) in 50 mM HEPES buffer (pH 7.2, 0.2 mM TFA). For in RBCs (sample 4-6); A solution of probe 1 (200 μM) and inhibitors (0 or 1 mM) in 2 mL HBS buffer (20 mM HEPES, 107 mM NaCl, 6 mM KCl, 1.2 mM MgSO₄, 2 mM CaCl₂, 11.5 mM glucose, pH 7.4) was added to a 2.5 mL RBC suspension. After centrifugation, the supernatant was removed, and the sedimented RBCs were resuspended in 50 mM HEPES buffer (pH 7.2, 100 mM NaCl, 0.17 mM TFA, 20% D₂O (v/v)) and subjected to NMR or MRI measurements at 25°C.