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Simultaneous Visualization of Protumorigenic Src and MT1-MMP Activities with Fluorescence Resonance Energy Transfer

Mingxing Ouyang1, He Huang1, Nathan C. Shaner3, Albert G. Remacle4, Sergey A. Shiryaev4, Alex Y. Strongin4, Roger Y. Tsien3, and Yingxiao Wang1,2

Abstract
Both Src kinase and membrane type 1 matrix metalloproteinase (MT1-MMP) play critical roles in cancer invasion and metastasis. It is not clear, however, how the spatiotemporal activation of these two critical enzymes is coordinated in response to an oncogenic epithelial growth factor (EGF) stimulation. Here, we have visualized the activities of Src and MT1-MMP concurrently in a single live cell by combining two fluorescence resonance energy transfer (FRET) pairs with distinct spectra: (a) cyan fluorescent protein (CFP) and yellow FP (YFP), and (b) orange FP (mOrange2) and red FP (mCherry). The new FRET pair, mOrange2 and mCherry, was first characterized in vitro and in cultured mammalian cells. When integrated with the CFP/YFP pair, this new pair allowed the revelation of an immediate, rapid, and relatively dispersed Src activity. In contrast, the MT1-MMP activity displayed a slow increase at the cell periphery, although Src was shown to play a role upstream to MT1-MMP globally. This difference in the activation patterns of MT1-MMP and Src in response to EGF is further confirmed using an optimized MT1-MMP biosensor capable of being rapidly cleaved by MT1-MMP. The results indicate that although Src and MT1-MMP act globally in the same signaling pathway, their activations differ in space and time upon EGF stimulation, possibly mediated by different sets of intermediates at different subcellular locations. Our results also showed the potential of mOrange2/mCherry as a new FRET pair, together with the popular variants of CFP and YFP, for the simultaneous visualization of multiple molecular activities in a single live cell. Cancer Res; 70(6); 2204–12. ©2010 AACR.

Introduction
MT1-MMP is a membrane-anchored enzyme belonging to a matrix metalloproteinase (MMP) family, known to be critical in cancer development by remodeling the extracellular matrix via proteolytic means (1, 2). Because membrane-tethered MMPs can be controlled and concentrated at subcellular locations (3), these membrane-associated MMPs seem to play more significant roles than soluble MMPs during cancer invasion (4). Indeed, MT1-MMP can be detected in a wide range of human cancers in clinical samples (5).

Epithelial growth factor (EGF) signaling and the associated Src activity have been well established to correlate with the invasive potential of a variety of human cancers (6, 7). Recent evidence indicates that Src can regulate MT1-MMP through focal adhesion kinase/endophilin A2 (8) and caveolin (9). Src was also shown to directly phosphorylate MT1-MMP at its cytoplasmic tail and regulate its function (10). However, it remains unclear how the activities of Src and MT1-MMP are coordinated in space and time to affect cancer development.

Genetically encoded biosensors based on fluorescence resonance energy transfer (FRET) have provided powerful tools for the study of molecular signals in live cells (11). To date, the most popular fluorescent protein (FP) pair for FRET is cyan and yellow FPs (CFP and YFP; ref. 12). A variety of genetically encoded biosensors based on CFP and YFP variants have been successfully developed to visualize molecular signals (13). However, with these biosensors, only one type of FRET signal can be visualized in a single live cell. Several studies have used one FP as the common donor or acceptor for two FRET biosensors to simultaneously visualize different molecular signals in the same cell (8, 14, 15). However, these approaches require more sophisticated means to quantify the signals than a simple donor/acceptor emission ratio. Recently, a newly developed FP, Ametrine, was elegantly paired with tdTomato to form a second FRET pair (16). Together with the pair of CFP and YFP variants, mTFP1 and mCitrine,
this new Ametrine/tetTomato pair has allowed the successful visualization of caspase-3 activities in different subcellular compartments. Although each individual FRET pair in this case can be ratiometrically imaged in the presence of the other, it is difficult to simultaneously visualize these two FRET pairs because of the almost identical emission of mAmetrine and mCitrine (16). Recently, a variety of new FPs with different colors have been developed in which mOrange (or mKO as orange FP) and mCherry (as red FP) seem suitable, serving as the donor and acceptor, respectively, for a second FRET pair that is spectrally distinguishable from CFP and YFP variants (12, 17, 18). Either mOrange or mKO has been successfully coupled with mCherry to form a FRET pair with efficiency detectable by fluorescence lifetime imaging microscopy (19). However, the photostability of mOrange is relatively poor (18). The CFP excitation also led to a substantial photoconversion of mKO, which may cause artifacts in cells expressing biosensors containing both CFP/YFP and mKO/mCherry pairs (19). Another two FRET pairs, (a) TagFP and mPlum (20) and (b) T-Sapphire and DsRed (21), have been used together with the CFP/YFP pair for dual FRET imaging. However, mPlum has a low quantum yield and poor brightness whereas DsRed can form artificial tetramers (18). Recently, mOrange2 was developed to have several fold greater photostability than both mOrange and mKO (18). The fusion proteins of α-tubulin and connexin-43 also showed correct subcellular localization when conjugated to mOrange2, but not to mKO (18). Hence, mOrange2 seems better-suited serving as a FRET donor for mCherry.

In this article, we have developed genetically encoded FRET biosensors with mOrange2 and mCherry, and showed that these biosensors are capable of visualizing Src and MT1-MMP activities in live cells. The dynamic activation of Src and MT1-MMP at subcellular levels was then concurrently monitored in a single live cell by combining CFP/YFP- and mOrange2/mCherry-based biosensors. The results suggest that the spatiotemporal activation patterns of Src and MT1-MMP were very different, with Src activity being relatively fast and dispersed whereas that of MT1-MMP was slow and highly localized at the cell periphery.
containing a SphI site or a stop codon together with an EcoRI site, respectively. The membrane-targeted Src biosensor was constructed by fusion of prenylation substrate sequence (KKKKKKSKTKCVIM) from KRas to the COOH terminus of Citrine or mCherry. The human MT1-MMP plasmid was a gift from Stephen J. Weiss at University of Michigan at Ann Arbor (25).

Expression of the biosensor proteins and in vitro assays. The biosensor proteins were expressed with N-terminal 6× His tags in Escherichia coli, which were cultured for 16 h at 37°C and purified by nickel chelation chromatography as previously described (ref. 23; Supplementary Fig. S1A). Emission ratio of mOrange2/mCherry (564 nm/604 nm) with excitation wavelength at 515 ± 10 nm was measured by a fluorescence plate reader (TECAN, Sapphire II). The cleavage assay for MT1-MMP biosensor was conducted at 37°C by incubating the recombinant catalytic domain of human MT1-MMP (MT1-CAT; 2 μg/mL) with 1 μmol/L of biosensor protein in a MT1-MMP proteolysis assay buffer [50 mmol/L HEPES (pH 6.8), 10 mmol/L CaCl₂, 0.5 mmol/L MgCl₂, 50 μmol/L ZnCl₂, and 0.005% Brij-35; ref. 26]. To compare the cleavage efficiency of different substrates, MT1-CAT was incubated for 3 h at 37°C with the protein substrates at a 1:25 to 1:1,600 molar ratio with human plasma α1-antitrypsin (AAT; Calbiochem), MBP-37, and the Nl(ECFP/YPet) biosensor.

Cell culture and transfection. Cell culture reagents were purchased from Life Technologies. HeLa cells were purchased from American Type Culture Collection and cultured in DMEM supplemented with 10% fetal bovine serum, 2 mmol/L of L-glutamine, 100 units/mL of penicillin, 100 μg/mL of streptomycin, and 1 mmol/L of sodium pyruvate in a humidified 95% air/5% CO₂ incubator at 37°C before imaging experiments. The cells were washed twice with HBSS, exposed to acidic conditions for 1.5 min [50 mmol/L glycine (pH 4.0), and 100 mmol/L NaCl] to remove the bound TIMP-2, and neutralized with 0.5 mol/L of HEPES (pH 7.5) and 100 mmol/L of NaCl. The cells were then washed with a CO₂-independent medium for three times before the FRET responses of the biosensor were monitored.

Results

Characterization of mOrange2 and mCherry as a novel FRET pair in vitro. Among the recently developed FPs (18), mOrange2 and mCherry may form a second FRET pair that is spectrally distinguishable from CFP and YFP variants. We have previously developed specific FRET biosensors for monitoring the activities of Src and MT1-MMP separately (22, 23). To visualize Src and MT1-MMP activities concurrently in the same cells, we generated a novel MT1-MMP biosensor by replacing the efficient ECFP and YPet pair (27) in the original biosensor with mOrange2 and mCherry (Fig. 1A). It is expected that active MT1-MMP would cleave the biosensor substrate peptide CPKESCNLFVLKD, thus separating mOrange2 and mCherry to result in a FRET decrease that could be tracked by an increase in the emission ratio of mOrange2/mCherry. Indeed, following the incubation of the purified biosensor (Supplementary Fig. S1A) with the catalytic domain of MT1-MMP (MT1-CAT), a decrease in mCherry emission was detected along with a concomitant increase in mOrange2 emission, indicative of a FRET decrease (Fig. 1B). The mOrange2/mCherry emission ratio gradually and consistently increased upon the incubation of MT1-CAT (Fig. 1C), confirming the effect of MT1-CAT on the biosensor. Analysis by gel electrophoresis further verified that the biosensor was completely cleaved upon the MT1-CAT incubation for 8 hours (Supplementary Fig. S1B).

Characterization of mOrange2 and mCherry as a FRET pair in mammalian cells. The mOrange2/mCherry-based MT1-MMP biosensor was also targeted to anchor outside of the plasma membrane in mammalian cells by a platelet-derived growth factor receptor transmembrane domain (22). This membrane-targeted biosensor was transfected together with the wild-type MT1-MMP or its control vector into HeLa cells, which express minimal levels of endogenous MT1-MMP (26). Strong fluorescence signals could be observed in both mOrange2 and mCherry channels with the mOrange2 excitation (Supplementary Fig. S2), suggesting an efficient FRET between these two FPs. EGF induced an increase in the emission ratio of mOrange2/mCherry, and hence, a FRET decrease of the biosensor in HeLa cells transfected with the wild-type MT1-MMP (Fig. 2A and C), but not with the control vector (Fig. 2B and C). Hence, these results indicate that the MT1-MMP FRET biosensor based on the mOrange2/mCherry pair could report MT1-MMP activity in vitro and in mammalian cells.
To further examine whether mOrange2 and mCherry are an efficient FRET pair for other kinds of biosensors, a Src FRET biosensor developed previously (23) was modified by replacing its ECFP and Citrine with mOrange2 and mCherry, respectively. A prenylation substrate sequence (KKKKKKSKTKCVIM) from KRas was then fused to the COOH terminus of mCherry to anchor the Src biosensor at the inner surface of the plasma membrane (KRas-Src biosensor; Fig. 1A; ref. 22). EGF induced an apparent increase in the mOrange2/mCherry emission ratio of this KRas-Src biosensor (Fig. 2D). This result confirmed that mOrange2 and mCherry could act as an efficient FRET pair for the development of different biosensors.

To further examine whether mOrange2 and mCherry are an efficient FRET pair for other kinds of biosensors, a Src FRET biosensor developed previously (23) was modified by replacing its ECFP and Citrine with mOrange2 and mCherry, respectively. A prenylation substrate sequence (KKKKKKSKTKCVIM) from KRas was then fused to the COOH terminus of mCherry to anchor the Src biosensor at the inner surface of the plasma membrane (KRas-Src biosensor; Fig. 1A; ref. 22). EGF induced an apparent increase in the mOrange2/mCherry emission ratio of this KRas-Src biosensor (Fig. 2D). This result confirmed that mOrange2 and mCherry could act as an efficient FRET pair for the development of different biosensors.

Visualization of Src and MT1-MMP activities concurrently in a single live cell. Src activation is related to the activity of MT1-MMP in regulating cancer development (8, 9). However, it remains unclear on the dynamic interrelationship between Src and MT1-MMP at subcellular levels, which is particularly important given that the function of Src is largely dependent on its subcellular location/environment, e.g., Src inhibits RhoA at the focal adhesion sites (28) but activates RhoA at podosomes (29). We first investigated the spatiotemporal distribution of Src and MT1-MMP by cotransfection of EGFP-conjugated Src and mCherry-conjugated MT1-MMP. As shown in Supplementary Fig. S3, EGF induced a significant colocalization of Src-GFP and MT1-MMP-mCherry at the cell periphery, especially at the lamellipodium-like regions. This result is consistent with previous reports that active Src can be transported to the plasma membrane to act on downstream signaling molecules (30, 31). We then combined the membrane-targeted and CFP/YFP-based KRas-Src biosensor together with the mOrange2/mCherry-based MT1-MMP biosensor so that the MT1-MMP activity at the outer surface and the Src activity at the inner surface of the plasma membrane could be monitored concurrently with high spatiotemporal resolution (Fig. 1A). The KRas-Src biosensor, MT1-MMP biosensor, and MT1-MMP were cotransfected into the same HeLa cells. EGF induced an immediate, fast, and relatively dispersed increase of Src activity which reached the peak within a few minutes. In contrast, MT1-MMP activity increased slowly and was concentrated at the cell periphery upon EGF stimulation in the same cell (Fig. 3A). The quantification of the FRET response at a local region proximal to the cell periphery further confirmed a fast and transient activation of Src but a slow and gradual activation of MT1-MMP in response to EGF (Fig. 3B). A similar phenomenon with a fast Src and slow MT1-MMP activation in response to EGF was also observed when the KRas-Src biosensor was switched to the mOrange2/mCherry pair and the MT1-MMP biosensor to the ECFP/YPet pair (data not shown). These results suggest that the different spatiotemporal responses of Src and MT1-MMP observed are not due to the difference in FRET pairs.

Figure 2. The characterization of mOrange2 and mCherry as a FRET pair in mammalian cells. A and B, representative mOrange2/mCherry emission ratio images of the MT1-MMP biosensor before and after EGF stimulation in HeLa cells expressing MT1-MMP (A) or a blank vector (B). C, representative time courses of normalized mOrange2/mCherry emission ratio of the MT1-MMP biosensor in HeLa cells expressing MT1-MMP (▴) or a blank vector (●). D, representative images (left) and time course of normalized mOrange2/mCherry emission ratio (right) in HeLa cells expressing the Src biosensor before and after EGF stimulation. Bars, 30 μm.
Because Src activity is related to the activity of MT1-MMP (8, 9), the different time courses of Src and MT1-MMP suggest that Src may play a role upstream of MT1-MMP in response to EGF. Indeed, the inhibition of Src by PP1 abolished the EGF-induced FRET response of the MT1-MMP biosensor in HeLa cells expressing exogenous MT1-MMP (Fig. 3C). In contrast, although the absence of exogenous MT1-MMP expression eliminated the response of the MT1-MMP biosensor in HeLa cells, EGF still induced a significant FRET response of the Src biosensor (data not shown). These results suggest that Src activity acts upstream of MT1-MMP activation in response to EGF.

The development of a MT1-MMP biosensor with a fast cleavage speed. It remains possible that the slow response of the MT1-MMP FRET biosensor upon EGF stimulation is attributed to the slow cleavage of the biosensor by active MT1-MMP, but not the slow activation of MT1-MMP itself. To address this question, we first examined the cleavage efficiency of the MT1-MMP biosensor (labeled as “NL” with the original substrate peptide sequence CPKESCNLFLVKLD). The cleavage rate of this biosensor upon incubation with MT1-CAT was compared with two typical MT1-MMP substrate molecules, AAT and MBP-J37. The purified substrate biosensors/proteins and MT1-CAT were incubated together for 3 hours with the enzyme/substrate ratios ranging from 1:25 to 1:1,600. As shown in Fig. 4A, a 50% cleavage of AAT and MBP-J37 could be accomplished at an enzyme/substrate ratio between 1:200 and 1:400, whereas the 50% cleavage of the NL biosensor was achieved at a ratio between 1:25 and 1:50. As a control, GM6001, a potent inhibitor of MMPs, inhibited MT1-CAT activity and blocked the cleavage of different substrates. Therefore, the NL biosensor is not an efficient substrate for MT1-MMP, suggesting a significant potential in improving the biosensor sensitivity by modifying the substrate peptide. A fast-cleavage biosensor could also allow the investigation of the activation speed of MT1-MMP in response to EGF with high accuracy.

We have hence developed an optimized MT1-MMP biosensor containing a selective and efficient peptide sequence (CRPAHLRD9G) capable of being rapidly and specifically...
cleaved by MT1-MMP, but not by gelatinases MMP-2 or MMP-9. The selection of this sequence was the result of our extensive substrate phage cleavage studies performed at the Center for Proteolytic Pathways in the Burnham Institute for Medical Research to identify the cleavage preferences of the individual MMPs including MT1-MMP.5

This optimized MT1-MMP substrate peptide (CRPAHRLRDG) flanked by GGS on both sides (labeled as "AHLR") was applied to replace the NL substrate peptide (CPKESCNLFLKLD) in the MT1-MMP biosensor. Following incubation with MT1-CAT, a rapid and significant decrease in YPet of the new AHLR biosensor was detected along with a concomitant increase in ECFP, suggesting an efficient FRET loss (Supplementary Fig. S4A). Gel electrophoresis further revealed that the biosensor could be completely cleaved by MT1-CAT (Supplementary Fig. S4B), confirming the high sensitivity of the AHLR biosensor. When the two purified biosensors (1 μmol/L) were incubated with MT1-CAT (2 μg/mL), the initial slopes of the emission ratio time courses for the NL and AHLR biosensors were 0.017 and 0.182, respectively (Fig. 4B). Therefore, the optimized AHLR biosensor is 10-fold more sensitive than the original NL biosensor. The FRET response of this AHLR biosensor in vitro is in fact faster than that of the Src biosensor measured by kinase assay as reported before (32).

The activation speed of the optimized AHLR biosensor in mammalian cells. We further compared the cleavage speeds of the two NL and AHLR biosensors in HeLa cells transfected with MT1-MMP. The basal activity of MT1-MMP was preinhibited by GM6001 for 12 hours such that the newly cycled and membrane-integrated biosensors were kept intact to result in a low emission ratio (ECFP/YPet) at the plasma membrane. After GM6001 washout, the biosensor proteins were exposed to the restored basal MT1-MMP activity on the cell surface, which resulted in an increase of FRET ratio. As shown in Fig. 5A, the emission ratio of the AHLR biosensor increased rapidly upon GM6001 washout, much faster than that of NL biosensor. A similar result could be observed when the MT1-MMP activity was preinhibited by a specific MT1-MMP inhibitor, TIMP-2 (Fig. 5B). These results indicate that the AHLR biosensor could be cleaved by MT1-MMP at a much faster speed than the NL biosensor in mammalian cells.

We then applied EGF to stimulate the AHLR biosensor in HeLa cells expressing MT1-MMP. As shown in Fig. 5C, the FRET response of the AHLR biosensor was still slow and comparable to that of the NL biosensor upon EGF stimulation. This EGF-induced response of the AHLR biosensor was significantly slower and weaker than that of the inhibitor washout, possibly suggesting a relatively high basal level of MT1-MMP activity, which is further inducible by EGF with moderate magnitude. The pretreatment of cells with the MT1-MMP inhibitor TIMP-2 completely blocked this EGF-induced response of the AHLR biosensor, suggesting the specificity of the AHLR biosensor (data not shown). These results confirmed that, upon EGF stimulation, the slow response of the biosensor is due to the slow activation of MT1-MMP but not to a slow cleavage rate of the biosensor by MT1-MMP. Therefore, the distinct spatiotemporal responses of the Src and MT1-MMP biosensors upon EGF stimulation indeed reflect the different activation patterns of Src and MT1-MMP upon EGF stimulation.

Discussion

Both Src and MT1-MMP have been shown to play pivotal roles in regulating cancer development and metastatic invasion. However, the coordination between these two critical molecules in space and time remains unclear. With mOrange2 and mCherry serving as a new FRET pair besides the popular CFP/YFP pair, we have combined two FRET biosensors with distinct colors for the concurrent visualization of Src and MT1-MMP activities in a single live cell. The results revealed distinctive spatiotemporal activation patterns of Src and MT1-MMP in response to EGF, with the Src activation occurring fast and dispersed whereas that of MT1-MMP being slow and concentrated at the cell periphery.

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Manuscript in preparation.
Cancer invasion is largely dependent on integrin-mediated cell adhesive interactions with the surrounding extracellular matrix (33, 34). These integrin-associated activities can be linked to MMP expression, activation, and proteolytic activity (25) in part via Src kinase-dependent processes (35, 36). Upon integrin activation, the focal adhesion kinase becomes tyrosine phosphorylated at site 397 and recruits Src to phosphorylate various signaling molecules, including focal adhesion kinase itself, p130Cas, caveolin, and endophilin A2. These events further promote cell surface MT1-MMP activity and invasion (8, 9). Consistently, Src has been shown to be related to MT1-MMP in a variety of cell systems (8–10), which is also in line with our results that the inhibition of Src blocked the FRET response of the MT1-MMP biosensor (Fig. 3C). However, it remains unclear how, in live cells, Src activity controls the MT1-MMP activity at subcellular levels. Our dual FRET imaging results indicate that the activation of Src and MT1-MMP differ both in space and time (Fig. 3A–B). Our optimized MT1-MMP biosensor capable of fast cleavage further confirmed that the EGF-induced MT1-MMP activation occurs slowly (Fig. 5C–D), in contrast to the fast activation of Src. This immediate activation of Src and the significant delay of MT1-MMP activation in the same subcellular location suggests that the effect of Src on MT1-MMP may require other molecular intermediates/mechanisms in addition to direct interactions, e.g., the direct phosphorylation of MT1-MMP cytoplasmic tail by Src as previously reported (10). The direct phosphorylation of MT1-MMP by Src is expected to occur much faster than the observed delay between these two signals. Indeed, this view of indirect interaction between Src and MT1-MMP is consistent with other reports that Src may indirectly regulate MT1-MMP through focal adhesion kinase/endophilin A2 (8) or caveolin (9). Src also regulates podosomes, in which MT1-MMP resides and plays an active role (37–39). It is hence possible that Src affects podosome formation in modulating the MT1-MMP activation (40). Further studies are warranted to elucidate the molecular mechanism by which Src regulates MT1-MMP activity in space and time in response to EGF.

It has become increasingly clear that signaling molecules inside cells are not functional in isolation. Instead, different signaling pathways interact with each other in a nonlinear fashion and form a coordinated network. Recent evidence further suggests that signaling networks or molecular hierarchies are largely dependent on subcellular localization, possibly due to the different molecular mediators at different subcellular locations. For example, Src induces the p190Rho-GAP activation and subsequently inhibits RhoA at the focal adhesion sites (28), whereas Src activates RhoA at podosomes (29). RhoA also couples with its downstream molecule ROCK at the cell rear and a contractile region behind lamellipodium, but colocalizes with another substrate molecule mDia at the leading edge of a migrating cell (41, 42). Hence, the development and integration of multiple FRET biosensors, e.g., using
mOrange2/mCherry and CFP/YFP pairs, could provide powerful tools to visualize molecular hierarchies at subcellular levels in live cells. The information obtained should significantly advance our in-depth and systematic understanding of the spatiotemporal molecular network coordinating cellular functions, including those crucial for cancer invasion and metastasis.

Recently, multiple new FPs have been developed with different colors (17, 18). This has provided opportunities for new FRET pairs which are spectrally distinguishable from the popular CFP/YFP or BFP/GFP pairs. As we have shown here, mOrange2 and mCherry could act as a new FRET pair to monitor active signaling events in vitro and in mammalian cells. However, there is a certain overlap between the excitation spectra of mOrange2 and mCherry (17, 18), i.e., the excitation of donor mOrange2 can directly excite the acceptor mCherry and cause nonspecific cross-talk. This artifact may have contributed to the lower dynamic range of the mOrange2/mCherry-based MT1-MMP biosensor (40% in vitro as shown in Fig. 1C) in comparison to the ECFP/YPet-based biosensor (570% in vitro; ref. 22). Fluorescence lifetime imaging microscopy technology could help to solve this problem because the emission lifetime measurement of the donor mOrange2 is alone sufficient to deduce the FRET efficiency, without the need to measure the emission lifetime of the acceptor mCherry (43). It is envisioned that the integration of fluorescence lifetime imaging microscopy and FRET will become increasingly accepted as a platform for FRET imaging, which is also an ongoing project in our laboratory.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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