A Fluorogenic Red Fluorescent Protein Heterodimer

Spencer C. Alford, Ahmed S. Abdelfattah, Yidan Ding, and Robert E. Campbell

1Department of Chemistry, University of Alberta, Edmonton, Alberta T6G 2G2, Canada
2Correspondence: robert.e.campbell@ualberta.ca

DOI 10.1016/j.chembiol.2012.01.006

SUMMARY

The expanding repertoire of genetically encoded biosensors constructed from variants of Aequorea victoria green fluorescent protein (GFP) enable the imaging of a variety of intracellular biochemical processes. To facilitate the imaging of multiple biosensors in a single cell, we undertook the development of a dimerization-dependent red fluorescent protein (ddRFP) that provides an alternative strategy for biosensor construction. An extensive process of rational engineering and directed protein evolution led to the discovery of a ddRFP with a K_d of 33 μM and a 10-fold increase in fluorescence upon heterodimer formation. We demonstrate that the dimerization-dependent fluorescence of ddRFP can be used for detection of a protein-protein interaction in vitro, imaging of the reversible Ca^{2+}-dependent association of calmodulin and M13 in live cells, and imaging of caspase-3 activity during apoptosis.

INTRODUCTION

Fluorescent protein (FP)-based biosensors that report on biochemical events in live cells typically operate on one of three principles: the modulation of Förster resonance energy transfer (FRET) efficiency between two FPs of different hues (Miyawaki et al., 1997), the modulation of the fluorescent intensity of a single FP through analyte-induced changes in the chromophore environment (Nakai et al., 2001; Nagai et al., 2001), or the fluorogenic reconstitution of a single FP (also known as complementation) from two polypeptide chains (Ghosh et al., 2000; Kerppola, 2009). Each of these design principles is associated with distinct advantages. For example, FRET-based biosensors benefit from being inherently ratiometric in their response, and single FP-based biosensors often provide better signal-to-noise because of larger intensiometric responses. FP complementation uniquely generates an irreversible response and can be applied for detection of interacting proteins in proteome-wide screens because of negligible background signal (Maglery et al., 2005; Remy and Michnick, 2004).

A shortcoming shared by these three design strategies is that they are most commonly implemented with engineered variants of Aequorea victoria green FP (GFP) and only rarely implemented with Anthozoan-derived red FPs (RFP) (Matz et al., 1999). Although there have been reports of orange FP-RFP FRET pairs (Pilić and Schultz, 2008; Ouyang et al., 2010), single RFP-based Ca^{2+} indicators (Zhao et al., 2011), and complementation of split RFPs (Jach et al., 2006; Fan et al., 2008), such examples are relatively few in number. Among the contributing factors to the discrepancy between the popularity of GFP and RFP variants in biosensing applications is the poor sensitized emission of RFPs when used as FRET acceptors (Ai et al., 2008) and the challenge of engineering circularly permuted RFPs for use in single FP-based biosensors (Carlson et al., 2010).

We reasoned that a new approach, which better harnesses the inherent properties of RFPs, could serve to stimulate development of a broader selection of RFP-based biosensors. One property that differentiates GFP from naturally occurring RFPs is the obligate tetrameric structure of the latter (Baird et al., 2000). For many applications, the tetrameric structure is undesirable, and substantial effort has gone into engineering dimeric and monomeric FPs (Campbell et al., 2002). However, it is also apparent that the oligomeric structure of RFPs helps stabilize the chromophore in a conformation that favors bright fluorescence. This role is apparent when one compares the quantum yields (Φ) of dimer (i.e., dimer2 and dTomato; both 0.69) and monomeric (i.e., mRFP1 and mCherry; 0.25 and 0.22, respectively) variants of Discosoma species RFP (DsRed) (Matz et al., 1999; Campbell et al., 2002; Shaner et al., 2004). This dimerization-dependent brightness suggested to us a new strategy for the creation of RFP-based biosensors.

Our idea was to develop an alternative biosensing system by engineering the oligomeric structure of RFPs to create a low-affinity heterodimer that exhibits bright red fluorescence in the associated state and dim fluorescence in the dissociated state. To engineer such a system, we expected that there would be three major hurdles to overcome: first, developing a heterodimeric RFP; second, engineering high contrast between the associated and dissociated states; and third, decreasing the affinity such that the dimer partners were not fully associated at intracellular concentrations typically used for FP imaging (~1–50 μM) (Martin et al., 2005). This manuscript describes our efforts to overcome these challenges and produce a fluorogenic RFP heterodimer that has proven useful in a series of representative biosensing applications.

RESULTS AND DISCUSSION

Engineering and Characterization of ddRFP

Our strategy for generating a fluorogenic RFP heterodimer was inspired by the well-established practice of introducing interface breaking mutations to produce a monomeric FP from a dimeric precursor (Campbell et al., 2002; Ai et al., 2006). Accordingly, we created a monomeric and dimly fluorescent RFP variant (designated A) by introducing the H162K and A164R substitutions into dTomato (Shaner et al., 2004). We then set out to
“rescue” dimer formation and fluorescence by creating a dTomato-derived partner (designated B) with surface modifications that complemented the modifications to A. To achieve this goal, we turned to the first of the three distinct library screening strategies (Figures 1 and 2; Figure S1 available online) that we implemented during this extensive process of molecular evolution (Table S1). For clarity, we append a subscript number to A and B to designate the generation of a given variant. Thus, dTomato-H162K/A164R is designated as A0.1, and the most extensively optimized version of A is designated A0.4.

To identify a heterodimeric partner for A0.1, we developed a two-step assay that involved an image-based screen for red fluorescence of *Escherichia coli* colonies followed by an electrophoretic mobility shift analysis of protein extracts (Figure 1A). A dual-expression plasmid with two different promoters (P_tac and P_BAD) was used to express A0.1, as a fusion with the 43 kDa *E. coli* maltose binding protein (MBP), and a dTomato-R149X/H162X/Y192X (where X = all 20 amino acids) gene library. These three residues were targeted because of their proximity to H162 and A164 across the AC dimer interface of tetrameric DsRed (Yarbrough et al., 2001). Colonies with the brightest red fluorescence were picked and cultured, and the protein extracts were analyzed by polyacrylamide gel electrophoresis (PAGE) under conditions that preserve high-affinity oligomeric interactions of analyzable proteins (i.e., supplemented with IPTG and L-arabinose). Screening of this library led to the identification of B0.2, which was equivalent to dTomato-E100R/R149L/E160H/H162K, and, by PAGE analysis, predominantly formed a homodimeric complex with A0.2. To further diminish the residual homodimeric character, we removed three residues that contribute to the dTomato interface (F224, L225, and Y226) (Yarbrough et al., 2001) from the C termini of A0.2 and B0.3. The resulting proteins, A0.3 and B0.4, exclusively formed a heterodimer, but exhibited reduced fluorescent brightness.

We next turned to a colony-based assay in which replica plating was used to identify noncovalent AB pairs that were brightly fluorescent when both proteins were expressed, but dimly fluorescent when only one protein was expressed (Figure 1B and Figure S1). Briefly, we used the dual-expression plasmid to express one randomly mutated partner under the isopropyl-β-D-thiogalactopyranoside (IPTG)-inducible P_tac promoter, and one genetically fixed partner under the L-arabinose-inducible P_BAD promoter. For library screening, transformed *E. coli* was plated on media permissive for expression of both partners (i.e., supplemented with IPTG and L-arabinose). The brightest colonies were picked and manually arrayed onto two new plates: one with media for expression of both partners (i.e., supplemented with IPTG and L-arabinose). The brightest colonies were picked and manually arrayed onto two new plates: one with media for expression of both partners (i.e., supplemented with IPTG and L-arabinose).

![Figure 1. Overview of the Screening Procedures Used to Identify Heterodimeric and Fluorogenic RFPs](image_url)

(A) Electrophoretic mobility shift screen for heterodimeric pairs of RFPs. 
(B) Replica-plating screen for fluorogenic and heterodimeric pairs of RFPs. 
(C) Tandem heterodimer proteolysis-based screen. See also Figure S1 and Table S1.
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**Figure 2. Characterization of Homo- and Heterodimeric Structure by Gel Electrophoresis**

The contrast of each whole image has been adjusted to emphasize weak bands.

(A) Validation of the SDS-PAGE electrophoretic mobility shift screen for pairs of protein variants that exhibit heterodimeric character. Proteins were loaded without boiling and were detected by imaging the red fluorescence. Lane 1, uninduced culture; lane 2, dTomato; lane 3, MBP-dTomato; lane 4, coexpressed dTomato and MBP-dTomato. Asterisk (*) indicates a species resulting from proteolysis of one dTomato-MBP linker.

(B) Red fluorescence image of a gel used for PAGE analysis of four representative variants analyzed during screening (numbered 1–4).

(C) Coomassie-stained gel of purified A and B variants. Samples were boiled in sample buffer prior to electrophoresis. The double asterisk (**) indicates the 19 kDa product of chromophore hydrolysis routinely observed for RFPs (Gross et al., 2000). The 11 kDa fragment was not observed.

FP partners exhibited a 5-fold increase in fluorescence upon dimerization both in E. coli (Figure 4A) and in vitro (Figure 4B), and a $K_d$ of 1.1 $\mu$M (Figure 3B).

To further improve the fluorogenic contrast of ddRFP-A$_{0.5}$B$_{0.5}$, we developed a third screening procedure that involved the use of proteolyzable tandem heterodimers (Figure 1C) composed of A and B joined by a 23-residue cleavable linker. Libraries of randomly mutated heterodimer genes were expressed in E. coli, and the brightest colonies were picked and cultured. The fluorescence of protein extracts was measured before and after treatment with trypsin, and clones with the greatest contrast were carried on to subsequent rounds. Application of this strategy for several rounds provided only moderate improvements in contrast, so we next attempted to rationally engineer improved contrast.

Because the fluorescence contrast of ddRFP is limited by the residual fluorescence of A in the absence of B (Figure 4B), we reasoned that manipulating the intraprotein interactions of the chromophore in partner A could lead to variants with improved contrast. Accordingly, we introduced the S146A substitution into A$_{0.4}$ to produce A$_{0.5}$, which had one less hydrogen bond donor in close proximity to the phenolate oxygen of the A chromophore (Yarbrough et al., 2001). From a library in which positions E144 and A145 of A$_{0.5}$ were randomized, we picked the clones that were most dimly fluorescent when expressed in the absence of B. The resulting A$_{0.6, \text{pool}}$ was fused in tandem to a B$_{0.5}$-derived gene library in which three residues (Y192, Y194, and H222) in close proximity to residues 144–146 of the dimer partner were randomized (Yarbrough et al., 2001). To decrease the $K_d$ of the heterodimer, we also reverted E100R and installed R153E in the B partner (Figure 3A) such that two favorable Arg-Glu electrostatic interactions were replaced with unfavorable Glu-Glu contacts. The library was screened as tandem heterodimers, and clones with substantially improved contrast were identified. The A and B partners of the clone that exhibited the highest contrast were designated A$_1$ and B$_1$.

Purified A$_1$ and B$_1$ exhibit a 10-fold increase in fluorescence upon dimerization (Figure 4C) and a $K_d$ of 33 $\mu$M (Figure 3B). As with ddRFP-A$_{0.4}$B$_{0.5}$, interaction with B$_1$ results in an instantaneous increase in the fluorescent brightness of A$_1$; A$_1$ alone has $\Phi = 0.026$ and an extinction coefficient ($\epsilon$) = 11,800 M$^{-1}$ cm$^{-1}$ at pH 7.4, and these values increase to 0.074 and 48,300 M$^{-1}$ cm$^{-1}$, respectively, upon formation of a heterodimer with B$_1$. No visible wavelength absorbance or fluorescence could be detected for B$_1$, even after several weeks at 4°C. Furthermore, boiling of purified B$_1$ resulted in none of the backbone hydrolysis that is characteristic of DsRed-type proteins (including A$_1$) with mature chromophores (Figure 2C) (Gross et al., 2000). We observed no limits to the solubility of A$_1$ and B$_1$; expression in E. coli gave yields of 10–80 mg/l. Both A$_1$ and the A$_1$B$_1$ complex exhibit relatively high fluorescence pK$_{as}$s (9.0 and 7.4, respectively), and, thus, their fluorescence is sensitive to physiologically relevant changes in pH (Figures S3A–S3C).

Size exclusion chromatography confirmed that both A$_1$ and B$_1$ are monomeric in isolation but form a heterodimer when mixed (Figure S3D). Interestingly, we noted that A$_1$B$_1$ undergoes a reversible reaction with I-mercaptoethanol (possibly a nucleophilic addition to the chromophore; Dong et al., 2008) that results in loss of the red fluorescent state and the formation of a new

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Figure 4. Characterization of ddRFPs
(A) Fluorescence image of E. coli expressing various proteins discussed in the text. A<sub>0.4</sub>B<sub>0.5</sub> and A·B<sub>1</sub> are tandem constructs. (B) Absorbance and fluorescence emission spectra for A·B<sub>1</sub> (dashed line; 10 µM), B<sub>1</sub> (gray line; 10 µM), and the equimolar mixture of A<sub>0.4</sub> and B<sub>0.5</sub> (black line; 10 µM each). (C) Absorbance and fluorescence spectra of A<sub>1</sub> (gray line; 20 µM) and tandem ddRFP-A<sub>1</sub>B<sub>1</sub> (black line; 20 µM) before (solid line) and after (dashed line) treatment with trypsin. B<sub>1</sub> has baseline absorbance and fluorescence. See also Figure S3.

species that absorbs at 402 nm and emits at 466 nm (Figures S3E and S3F).

The convoluted evolutionary pathway that led to ddRFP-A<sub>1</sub>B<sub>1</sub> reflects the practical challenge of simultaneously optimizing at least five distinct, and possibly conflicting, properties: heterodimeric structure, minimal homodimeric structure, high-contrast fluorogenesis, high brightness in the associated state, and high K<sub>d</sub>. Accordingly, ddRFP-A<sub>1</sub>B<sub>1</sub> represents a compromise, but it is the best compromise that we have identified to date.

In Vitro and Live Cell Applications of ddRFP
We next investigated the utility of ddRFP-A<sub>1</sub>B<sub>1</sub> in representative in vitro and live cell imaging applications. To determine whether ddRFP-A<sub>1</sub>B<sub>1</sub> could be used to detect protein-protein interactions in vitro, we turned to the rapamycin-dependent interaction of FK506-binding protein (FKBP) and the FKBP-rapamycin-binding domain (FRB) (Chen et al., 1995). We fused A<sub>1</sub> and B<sub>1</sub> to the C termini of FRB and FKBP with a 43-residue unstructured linker to produce FRB·A<sub>1</sub> and FKBP·B<sub>1</sub>. These fusions were expressed in E. coli, purified, and mixed at a concentration (0.5 µM each) that is well below the K<sub>d</sub> for ddRFP-A<sub>1</sub>B<sub>1</sub>. As shown in Figure 5A, the addition of rapamycin gave a dose-dependent increase in fluorescence. This result indicates that ddRFP-A<sub>1</sub>B<sub>1</sub> is useful for detection of protein-protein interactions with dissociation constants well below the ddRFP-A<sub>1</sub>B<sub>1</sub> K<sub>d</sub>.

To determine whether ddRFP-A<sub>1</sub>B<sub>1</sub> could be employed to detect reversible protein-protein interactions in live cells, we borrowed the so-called “split-cameleon” strategy in which the Ca<sup>2+</sup>-induced interaction of calmodulin (CaM) and the M13 peptide brings two FPs into close proximity (Miyawaki et al., 1999). Accordingly, mammalian expression plasmids were prepared in which A<sub>1</sub> was fused to the N terminus of CaM (A<sub>1</sub>-CaM) and B<sub>1</sub> was fused to the C terminus of the M13 peptide (M13-B<sub>1</sub>). HeLa cells were cotransfected with both plasmids and imaged at 24 hr after transfection. Upon histamine treatment, we observed oscillations in the red fluorescence intensity (Figure 5B) that were consistent with the expected changes in intracellular Ca<sup>2+</sup> concentration. Other FP-based Ca<sup>2+</sup> indicators based on FRET (Miyawaki et al., 1999) and single FP Ca<sup>2+</sup> biosensors (Zhao et al., 2011) exhibit Ca<sup>2+</sup> spikes of duration similar to those observed with A<sub>1</sub>-CaM and M13-B<sub>1</sub>, indicating that, at least on the timescale of seconds, heterodimer association and dissociation is not a rate-limiting step. In situ calibration of the dynamic range was accomplished by treating cells with ionomycin/EGTA to deplete Ca<sup>2+</sup> and ionomycin/Ca<sup>2+</sup> to saturate CaM. These experiments revealed that the dynamic range for A<sub>1</sub>-CaM plus M13-B<sub>1</sub> in cells was 5.7 ± 1.3 (N = 16). This dynamic range is approximately half of the in vitro range for A<sub>1</sub> and B<sub>1</sub> alone, indicating that the partners are approximately 50% associated in the absence of Ca<sup>2+</sup> and at the intracellular protein concentrations of these experiments. Accordingly, we estimate that the intracellular concentrations of A<sub>1</sub>-CaM and M13-B<sub>1</sub> in those cells bright enough to be imaged, is approximately 30 µM.

For a third representative application of ddRFP-A<sub>1</sub>B<sub>1</sub>, we envisioned a strategy for protease activity sensing in which A<sub>1</sub> and B<sub>1</sub> were fused in tandem with the well-characterized Asp-Glu-Val-Asp (DEVD) caspase-3 substrate sequence in the linker region (Xu et al., 1998). As in our trypsinolysis assay, cleavage of the substrate should result in a loss of fluorescence as the partners dissociate, provided the concentration is well below the K<sub>d</sub>. HeLa cells were transfected with a plasmid harboring the DEVD tandem heterodimer, treated with tumor necrosis factor-α (TNF-α), and imaged through time. We reliably observed a decrease in red fluorescence intensity (dynamic range of 8.6 ± 4.3; N = 9), just prior to the cell shrinkage, and...
blebbing associated with the end stages of apoptosis (Figure 5C). The rate of the transition from the bright to dim state was comparable to the rate previously observed for a FRET-based biosensor of caspase-3 activity (Ai et al., 2008), indicating that heterodimer association and dissociation is not a rate-limiting step on the relatively slow timescale (minutes) of this experiment. Overall, these results confirm that the affinity of A₁ and B₁ is sufficiently low to allow protease detection and imply that the tandem heterodimer is sufficiently bright for imaging at intracellular concentrations well below the \( K_d \) of 33 \( \mu \text{M} \).

Homology Modeling of ddRFP-A₁B₁
To gain a better understanding of how the 35 amino acid substitutions in ddRFP-A₁B₁ (relative to dTomato) contribute to its heterodimeric character and fluorogenesis, we created a homology model of ddRFP-A₁B₁ (Figure 6). Inspection of this model reveals that A₁ and B₁ have six and four substitutions, respectively, at positions with side chains directed toward the interior of the β-barrel. In A₁, just three of these residues are in close proximity to the chromophore: the conservative I161L substitution and the nonconservative S146A and K163G substitutions in ddRFP-A₁B₁ are shown in stick and representation (Yarbrough et al., 2001). Figure was generated with PyMOL (http://www.pymol.org). See also Figure S4.

The homology model of ddRFP-A₁B₁ also reveals that shape complementarity and electrostatic interactions across the dimer interface (Figure S4C) have been substantially modified relative to dTomato (which preserves the interface of DsRed) (Figure S4D). Indeed, of the 19 substitutions in A₁ and B₁ with side chains directed to the exterior of the β-barrel, 15 are located in the interface. The homology model suggests that a principal determinant of the heterodimeric character is a new `bump-and-hole` interaction across the dimer interface. In the X-ray crystal structure of DsRed, the large side chain of R149 is positioned close to the small side chain of A164 on the dimer partner (Figure S4B) (Yarbrough et al., 2001). In ddRFP-A₁B₁, the A164R substitution of A₁ is a `bump` that was introduced to destabilize the dTomato homodimer as a result of steric clashes and

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**Figure 5. Fluorescence Detection of In Vitro Chemically Induced Dimerization and Imaging ddRFP-A₁B₁ in Live Cells**

(A) Fluorescence of various combinations of FRB-A₁ (0.5 mM), FKBP-B₁ (0.5 mM), rapamycin, and dimethyl sulfoxide (DMSO) vehicle control. Fluorescence intensity is the mean integrated emission intensities for three independent experiments (± standard deviation) and corrected for background (i.e., assay buffer alone).

(B) Imaging of Ca²⁺ dynamics with A₁-CaM and M13-B₁ in HeLa cells. Transfected HeLa cells were treated with histamine, followed by EGTA/ ion/ Ca²⁺/ CaM vehicle control. Fluorescence intensity is the mean integrated emission intensities for three independent experiments (± standard deviation) and corrected for background (i.e., assay buffer alone).

(C) Imaging of caspase-3 activity with a ddRFP-A₁B₁ tandem heterodimer in HeLa cells. Transfected HeLa cells were treated with TNF-α (t = 0) to initiate apoptosis, and red fluorescence was imaged as a function of time.

**Figure 6. Structural Model of ddRFP-A₁B₁ Based on the AC Dimer of DsRed**
The amino acid side chains for all mutations in ddRFP-A₁B₁ are shown in stick representation (Yarbrough et al., 2001). Figure was generated with PyMOL (http://www.pymol.org). See also Figure S4.

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**Table 1:**

<table>
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<th>Rap (μM)</th>
<th>DMSO</th>
<th>FRB-A₁</th>
<th>FKBP-B₁</th>
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electrostatic repulsion with R149 of B1. In our model, the “hole” on the surface of B1 is formed by reorientation of the side chain of R149M of B1 such that it is directed away from A164R of A1 and is filling an adjacent cavity created by Y194C of B1 (Figure S4A).

Two factors contribute to the fluorogenesis of ddRFP-A1B1: pKα modulation and Φ modulation. At physiological pH and in the absence of B1, the fully formed chromophore of A1 exists primarily in the protonated and nonfluorescent state because of its pKα of 9.0. Upon interaction with B1, the chromophore environment of A1 is modified such that the pKα is lowered to 7.4 and the equilibrium shifts toward the anionic fluorescent state. A similar mechanism has been proposed for a system in which a single domain antibody (nanobody) modulates the fluorescence of GFP (Kirchofer et al., 2010). An additional contribution to fluorogenesis is the increase in Φ from 0.026 in A1 to 0.074 in the A1B1 complex. We speculate that, in free A1, the chromophore spends more time in noncoplanar conformations that favor excited-state deactivation by processes other than fluorescence (Shu et al., 2006). Upon formation of the A1B1 complex, a modified chromophore environment stabilizes a coplanar conformation of the chromophore, and Φ increases. Further structural studies should shed light on the molecular details of the interface interactions that contribute to fluorogenesis.

Outlook for ddRFP-A1B1 as a Biosensing Strategy
With the advent of ddRFP-A1B1, a new entry has been added to the short list of strategies (i.e., FRET, insertion in a single FP, and FP complementation) that can be used for creation of FP-based biosensors. We have demonstrated the practicality of ddRFP-A1B1, by using it to detect protein-protein interactions, image a reversible Ca2+-dependent protein-protein interaction, and image protease activity. The primary advantage of ddRFP-A1B1 relative to other strategies is that it provides a general means of creating red intensiometric biosensors with a reversible response. Potential disadvantages include the pH sensitivity and relative low brightness of ddRFP-A1B1, though we note that the brightness is comparable to the commercially available DsRed-monomer (Strongin et al., 2007). Relative to the three well-established biosensing strategies, dimerization of A1 and B1 to form ddRFP-A1B1 is most closely analogous to FP complementation (Ghosh et al., 2000; Kerppola, 2009). A key difference between single FP complementation and ddRFP-A1B1 heterodimerization is that, although the fragments of a split FP are nonfluorescent, A1 retains 10% of the brightness of the complex. This residual fluorescence is likely to complicate efforts to use ddRFP-A1B1 in high-throughput screens for identification of interacting proteins (Jackrel et al., 2010).

Future efforts to improve ddRFP-A1B1 should provide variants with decreased affinity and increased contrast. Until such variants become available, the most appropriate applications of ddRFP-A1B1 are as an alternative to intermolecular FRET and for protease activity sensing with tandem heterodimers. As we have done with the split cameleon-type Ca2+-bound biosensor and the caspase-3 biosensor, it should be relatively straightforward to convert existing FRET-based biosensors into ddRFP-A1B1-based biosensors. In these contexts, ddRFP-A1B1 will complement the existing repertoire of FP-based biosensing strategies, because it enables creation of spectrally distinct biosensors with an intensiometric and reversible red fluorescent signal.

SIGNIFICANCE

By employing a series of three protein library screening strategies, we have engineered ddRFP-A1B1, a fluorogenic DsRed-derived heterodimer with an associated state that is 10-fold brighter than the dissociated state. The interaction has been engineered to have a Kd of 33 μM, and, therefore, the heterodimers can exist primarily in the dissociated state at cytoplasmic concentrations sufficient for live cell imaging. Homology modeling has revealed that this heterodimeric character results from a new “bump-and-hole” interaction that is not present in the homodimeric parent protein. The utility of ddRFP-A1B1 has been demonstrated in three applications, including detection of a protein-protein interaction in vitro, imaging of a Ca2+-dependent protein-protein interaction in live cells, and imaging of caspase-3-dependent proteolysis during apoptosis.

EXPERIMENTAL PROCEDURES

Library Screening by Gel Shift
A custom bacterial dual-expression plasmid, containing a Xhol/HindIII polylinker under control of Pmlec and an EcoRl/BglII polylinker under control of PBAD, was used for the rescue-of-binding assay. A silent mutation was introduced into the gene encoding maltose-binding protein (MBP) to remove an internal BglII site. The fusion of MBP and dTomato H162K/A164R with an intervening KpnI site was ligated into the PBAD polylinker. Random mutations at sites R149, H162, and Y192 were introduced into a second dTomato gene by overlap extension PCR (Bessette et al., 2003), and the gene product was ligated into the Pmlec polylinker. Colonies of E. coli were screened with 535/50 nm excitation and visualization through 600 nm long-pass goggles. Bright colonies were picked into Luria-Bertani (LB) media supplemented with 100 μg/ml ampicillin, 0.1 mM IPTG, and 0.02% L-arabinose and were incubated overnight. Proteins were purified using Nickel-NTA, electrophoresed by SDS-PAGE, and analyzed for in-gel fluorescence using digital imaging with appropriate filters (excitation 535/50 nm; emission 630/60 nm).

Library Screening for Fluorogenesis
Error-prone PCR was performed using Taq polymerase as previously described (Campbell et al., 2002). E. coli was transformed with gene libraries in the Pmlec and PBAD polylinker sites of the dual-expression plasmid. Bright colonies were picked and replica plated in a regular grid pattern on one plate supplemented with 1 mM IPTG and 0.2% L-arabinose and on one plate supplemented with 1 mM IPTG and 10 mM glucose. Following overnight incubation at 37°C, the red fluorescence of both plates was imaged, and the ratios of intensities on the L-arabinose versus glucose plates for each colony were determined using image processing macros. Clones with the highest ratios were pooled and used in the next round.

Library Screening with Tandem Heterodimers
Tandem heterodimers were initially constructed by a three-part ligation strategy, which provided a chimera of the form A1-linker-B in Xhol/HindIII sites of pBAD/His B, where the linker was a 23-residue sequence that included a KpnI site. Bright colonies were picked and cultured in LB with ampicillin and L-arabinose (0.02%) overnight. Crude protein extracts were treated with trypsin at approximately 10 μg/ml for 30 min, and emission spectra were acquired using a 96-well microplate reader. Contrast ratios were calculated as the integrated emission peak area of the nontrypsinized extract divided by the peak area of the trypsinized extract.
Protein Production and Characterization

To obtain purified proteins, the genes for tandem ddRFP variants and monomeric partners were subcloned into pBAD/His B and were used to transform E. coli. Cultures at an optical density of 0.5–0.7 were induced with 0.02% L-arabinose and allowed to grow for a further 12–16 hr. Soluble proteins were purified from cleared lysates by Nickel-NTA and were dialyzed into 5 mM Tris-Cl and 100 mM NaCl (pH 7.5). pH sensitivity measurements were performed by incubating purified proteins in buffers of desired pH and by acquiring emission spectra with a 96-well microplate reader. Spectra represented in this manuscript were recorded with a DU-800 UV-visible spectrophotometer (Beckman) or a QuantMaster spectrofluorimeter (Photon Technology International, Inc). The amino acid sequence of His-tagged recombinant proteins was determined by automated Edman degradation.

FKBP-FRB Interaction

The genes encoding FRB and FKBP were fused to A, and B, respectively, to generate FRB-A, and FKBP-B, fusions. For both constructs, the linker sequence corresponded to residues 95 to 135 of phage λ repressor (accession NP_040628). The fusion proteins were expressed from pBAD/His B and soluble proteins were purified as described for the tandem ddRFP variants. Protein concentrations were determined by the BCA method.

Live Cell Imaging

The A, and B, copies in the DEVD tandem heterodimer were linked by the sequence GHTGTHSTHSHHSHSTHSHSEVDA. The gene sequence encoding the nuclear exclusion sequence (LAALKLAGLDIGS) was appended to the 3′ end of the gene (Wen et al., 1999). For the split cameleon-type biosensor, the genes encoding A, and CaM, as well as M13 and B, were joined by the sequence GHTGTHSTHSHHSHSTHSHSEVDA. The genes encoding the tandem heterodimers and both the CaM and M13 fusions were ligated into the HindIII sites of pcDNA3.1(+). HeLa cells were transfected using the FuGene transfection reagent (Roche). Transfected cells were imaged in HEPES-buffered Hank’s balanced salt solution and were consecutively treated with histamine (25 μM), ETGTA (3 mM) with ionomycin (5 μM), and 10 mM CaCl₂ with 5 μM ionomycin. For caspase-3 imaging, apoptosis was initiated by treatment with 100 ng/ml TNF-α.

ACKNOWLEDGMENTS

We thank the University of Alberta MBSU for technical assistance and Andreas Ibraheem for the dual-expression plasmid. Work in the Campbell laboratory is made possible by grants from CIHR and NSERC. S.C.A. is supported by Ph.D. scholarships from NSERC and Alberta Ingenuity. R.E.C. holds a Tier II Canada Research Chair in Biochemical Chemistry.

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