A Monomeric Photoconvertible Fluorescent Protein for Imaging of Dynamic Protein Localization

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The use of green-to-red photoconvertible fluorescent proteins (FPs) enables researchers to highlight a subcellular population of a fusion protein of interest and to image its dynamics in live cells. In an effort to enrich the arsenal of photoconvertible FPs and to overcome the limitations imposed by the oligomeric structure of natural photoconvertible FPs, we designed and optimized a new monomeric photoconvertible FP. Using monomeric versions of 
Clavularia sp. cyan FP as template, we employed sequence-alignment-guided design to create a chromophore environment analogous to that shared by known photoconvertible FPs. The designed gene was synthesized and, when expressed in 
Escherichia coli, found to produce green fluorescent colonies that gradually switched to red after exposure to white light. We subjected this first-generation FP [named mClavGR1 (monomeric 
Clavularia-derived green-to-red photoconvertible 1)] to a combination of random and targeted mutageneses and screened libraries for efficient photoconversion using a custom-built system for illuminating a 10-cm Petri plate with 405-nm light. Following more than 15 rounds of library creation and screening, we settled on an optimized version, known as mClavGR2, that has eight mutations relative to mClavGR1. Key improvements of mClavGR2 relative to mClavGR1 include a 1.4-fold brighter red species, 1.8-fold higher photoconversion contrast, and dramatically improved chromophore maturation in 
E. coli. The monomeric status of mClavGR2 has been demonstrated by gel-filtration chromatography and the functional expression of a variety of mClavGR2 chimeras in mammalian cells. Furthermore, we have exploited mClavGR2 to determine the diffusion kinetics of the membrane protein intercellular adhesion molecule 1 both when the membrane is in contact with a T-lymphocyte expressing leukocyte-function-associated antigen 1 and when it is not. These experiments clearly establish that mClavGR2 is well suited for rapid photoconversion of protein subpopulations and subsequent tracking of dynamic changes in localization in living cells.

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Introduction

In recent years, a handful of 
Aequorea green fluorescent protein (GFP) homologues cloned from anthozoan organisms have been reported to undergo irreversible photoconversion from a green fluorescent species to a red fluorescent species upon illumination with light of approximately 400 nm. To date, the naturally occurring photoconvertible proteins that have received the most attention are Kaede from coral 
Trachyphyllia geoffroyi,¹ EosFP from stony coral 
Lobophyllia geoffroyi.
hemprichii,² and Dendra from octocoral Dendro neuropthya sp.³ It has also been demonstrated that a naturally occurring nonphotoconvertible fluorescent protein (FP) can be engineered into a photoconvertible FP. Specifically, the photoconvertible FP known as KikGR was engineered from the green fluorescent variant KikG of the coral Favia favus.⁴ All green-to-red photoconvertible FPs characterized to date share a common His-Tyr-Gly-derived chromophore structure and a common photoconversion mechanism (Fig. 1).⁵ The newly synthesized protein first folds into the characteristic β-barrel structure of the Aequorea GFP superfamily⁶ and then undergoes the steps of posttranslational modification that lead to the formation of a green fluorescent chromophore with a conjugated system identical with that of Aequorea GFP chromophore. The chromophore can exist either in its neutral phenol form or in its anionic phenolate form (Fig. 1). Exactly where the equilibrium between these two forms lies is dependent on the local microenvironment of the chromophore (as determined by amino acid substitutions in close proximity to it) and the pH of the solution. The green-to-red photoconvertible FPs are distinguished from their nonphotoconvertible brethren by the respective consequences of exciting the neutral form, which absorbs most strongly at ~400 nm. In wild-type Aequorea GFP, the excited state of the neutral species undergoes excited-state proton transfer to form the anionic species, which then emits green fluorescence.⁷ In the case of the green-to-red photoconvertible FPs, excitation of the neutral form leads to a break of the polypeptide chain through effective β-elimination of the residue immediately preceding the chromophore-forming His-Tyr-Gly tripeptide.⁸,⁹ This elimination reaction results in the installation of a new double bond between C⁶ and C⁷ of the His residue, placing the side-chain imidazole in conjugation with the remainder of the avGFP-type chromophore. This extended conjugation decreases the energy gap between the highest occupied molecular orbital and the lowest unoccupied molecular orbital, and shifts the emission to the orange-to-red region of the visible spectrum. Photoconversion (i.e., a light-induced change in excitation or emission wavelength maxima) via alternate mechanisms has been observed in other color classes of FP.⁹,¹⁰

FP photoconversion allows researchers to ‘highlight’ a subpopulation of proteins within a cell or tissue through spatially defined illumination with a specific wavelength of light.¹¹ The subsequent dynamics of the highlighted protein can be achieved using FPs that undergo so-called photostabilization (irreversible conversion from a nonfluorescent species to a fluorescent species upon illumination)¹² and photoswitching (reversible conversion between a nonfluorescent species and a fluorescent species upon illumination).¹³ A more recent application of these types of proteins that has generated excitement in the cell biology community is superresolution fluorescence imaging.¹⁴ This application also involves highlighting a subpopulation of protein molecules; however, instead of being in a spatially defined location, they are sparsely distributed throughout the whole sample to be imaged. The sample is then imaged to reveal the precise locations of the point sources of fluorescence, each of which corresponds to a single molecule of an FP. The sample is bleached, and the process is repeated. Many repetitions of this imaging protocol can produce images in which structures, with dimensions of ~50 nm can be resolved.¹⁵

Kaede, the first example of an FP that can undergo irreversible green-to-red photoconversion upon illumination with UV light, was initially described by Ando et al. in 2002.¹¹ Unfortunately, the range of potential applications for Kaede remains limited by the fact that it is an obligate tetramer,¹⁶ and no

![Fig. 1. Chromophore structure and UV-induced modifications of green-to-red photoconvertible FPs.](image-url)

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Residue numbers for the nascent polypeptide are for the mTFP1-derived variants described in this work. Broken lines represent connections to the polypeptide chain of the protein.
monomeric variants have been reported. Unlike monomeric FPs, tetrameric FPs are generally detrimental to the proper trafficking and localization of recombinant fusion proteins.\textsuperscript{19} In producing a monomeric green-to-red photoconvertible FP, the same workers appear to have had more success with engineered variants of the tetrameric KikGR FP, which is substantially brighter and more efficiently photoconverted than Kaede.\textsuperscript{4} A monomeric version of KikGR, known as mKikGR, has recently been reported.\textsuperscript{20}

The two other green-to-red photoconvertible FPs, EosFP and Dendra,\textsuperscript{21,22} have been subjected to protein engineering to convert wild-type tetrarsers into monomers.\textsuperscript{17,21} However, it is apparent that the monomeric version of EosFP retains a tendency to form dimers at higher concentrations.\textsuperscript{17} The monomeric variant of EosFP, known as mEos, was created through the introduction of two point mutations that disrupted the protein–protein interfaces of the tetrameric species.\textsuperscript{2,22} Expression of mEos at temperatures greater than 30 °C is problematic,\textsuperscript{2} but an effectively monomeric tandem dimer variant does express well at 37 °C.\textsuperscript{23} The poor expression of mEos at 37 °C has been overcome with the engineering of mEos2 through targeted substitution of residues with solvent-exposed side chains.\textsuperscript{17} Although mEos2 has been reported to retain some propensity for dimer formation, this property does not appear to have adverse effects on the subcellular targeting of a variety of fusion proteins.\textsuperscript{17}

The growing number of reports on optimized photoconvertible FPs reflects the growing demands on these proteins with respect to their enabling role in some popular cell biology applications. In particular, there is a continued need for bright and monomeric photoconvertible proteins, since these tools can enable the highest-precision superresolution fluorescence imaging.\textsuperscript{17,21} In an effort to create a new monomeric photoconvertible FP with favorable properties, we embarked on a strategy distinct from that previously employed. Rather than start with a tetrameric photoconvertible protein and engineer it to be monomeric, we sought to start with a well-characterized monomeric coral-derived FP and engineer it to be a green-to-red photoconvertible FP. That tetrameric KikG was converted into KikGR through the use of protein design sets a precedent that it should be possible to convert a nonphotoconvertible FP into a photoconvertible FP.\textsuperscript{4} Our starting template is a monomeric version of Clavularia sp. cyan FP (cFP484, a wild-type Clavularia cyan FP) known as mTFP1 (monomeric teal FP 1).\textsuperscript{24}

**Results and Discussion**

**Guided consensus design of a new photoconvertible FP**

The new photoconvertible FP based on the mTFP1 template was designed using a strategy analogous to that previously used to design a consensus FP based on a monomeric Azami green template.\textsuperscript{25} The known photoconvertible FPs, including EosFP,\textsuperscript{3} Dendra2,\textsuperscript{2} KikGR,\textsuperscript{4} and Kaede,\textsuperscript{5} were aligned to find the consensus at each amino acid position (Fig. 2). Amino acids of >50% consensus were maintained in the designed protein. At positions with no clear consensus, the corresponding residue in mTFP1 was used.\textsuperscript{2,3} Residues of mTFP1 that had been substituted during the conversion of the wild-type cFP484 tetramer into a monomer were maintained in the design of the consensus protein. The designed gene was synthesized and, when expressed in *Escherichia coli*, found to exhibit green fluorescence that gradually switched to red after exposure to white light for 1–2 h. This variant was designated mClavGR1 (monomeric *Clavularia*-derived green-to-red photoconvertible 1) (Fig. 2). A complete list of all substitutions present in mClavGR1 and all other variants described in this work is provided in Supplementary Table 1.

**Directed evolution with screening for photoconversion**

Starting from mClavGR1, we undertook 11 rounds of evolution, where each round involved library creation by error-prone PCR\textsuperscript{26} and staggered extension process (StEP) shuffling,\textsuperscript{27} followed by a colony-based screen for those variants with the highest brightness and the highest ratio of red fluorescence after photoconversion to green fluorescence before photoconversion. In each round, approximately 6000 colonies were screened, and approximately 5–15 of the brightest variants were picked. A mixture of the plasmid DNA from these variants was then used as template for the subsequent round of library creation and screening. After the fourth round of evolution, we sequenced the clone with the highest fluorescence intensity in both the green channel (before photoconversion) and the red channel (after photoconversion). This clone, designated mClavGR1+, was equivalent to mClavGR1+T6bI, S34R, K36R, K74R, M113I, L166Q, Y173H (Fig. 2). A genealogy of all mClavGR variants is provided in Fig. 3a. Directed evolution starting from a template mixture of mClavGR1+ and other bright variants was continued for seven more rounds. While we primarily relied on the diversity generated by error-prone PCR and StEP shuffling, we also created libraries in which specific positions that have previously been shown to improve the properties of homologous FPs were randomized (Fig. 3a; Supplementary Table 2). These rationally generated libraries were screened in a manner identical with that used for random libraries.

The brightest variant after the 11th round of directed evolution was equivalent to mClavGR1+T6bI, T41I, D77E, F99Y, K115E, E127V, K139R, A183V (Fig. 3b). Characterization of the oligomeric state of this variant by gel-filtration chromatography revealed that the protein existed as a dimer (data not shown) and thus was designated dClavGR1.6.
Fig. 2. Sequence alignment of mClavGR variants and other proteins relevant to this study. Substitutions in mClavGR1, relative to mTFP1, are represented as black text against a gray background. Substitutions in later mClavGR variants that were identified during the directed evolution are shown as white text against a black background. Residues with side chains directed towards the interior of the β-barrel are enclosed in black boxes. The chromophore-forming residues are numbered 66–68 in this alignment.
where the prefix ‘d’ indicates a dimer. dClavGR1.6 is about 5-fold brighter than mClavGR1 when expressed in *E. coli* and shares only the T6bI substitution with mClavGR1.1. Analysis of the substitutions present in dClavGR1.6, in combination with a structure-based alignment of the X-ray structures of mTFP1 and tetrameric amFP486,24,28 led us to suspect that the E127V substitution was most likely responsible for reconstituting the dimer interface (Fig. 3c). Accordingly, we introduced the single amino acid reversion V127E into dClavGR1.6 by site-directed mutagenesis to produce mClavGR1.8. Although mClavGR1.8 was monomeric, it retained only 50% of the fluorescent brightness of dClavGR1.6 when expressed in *E. coli*. Noting that the V123T substitution had been previously used to convert EosFP into mEos,17 and that position 123 of EosFP aligns with position 127 of mClavGR1.8, we introduce the E127T substitution into mClavGR1.8. The resulting protein, designated mClavGR2 (Figs. 2 and 3b), had regained 60% of the brightness of dClavGR1.6 when expressed in *E. coli* and retained the monomeric character of mClavGR1.8 as determined by gel-filtration chromatography at a concentration of 1 mM (Supplementary Fig. 1). Under the same experimental conditions, Dendra2 was also monomeric, but mEos2 existed as a dimer or higher-order oligomer at concentrations of 0.1 mM and greater. Further efforts to improve the protein by screening libraries in which residues of interest were randomized (Fig. 3a; Supplementary Table 2) did not result in identification of further improved variants.

**In vitro characterization of mClavGR2**

As described in Directed Evolution with Screening for Photoconversion, we used an empirical screening process in order to produce a protein that had an optimal combination of brightness and photoconversion efficiency. In order to determine what changes in the spectral properties of the protein

<table>
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<th>Protein</th>
<th>λ_ab (nm)</th>
<th>λ_em (nm)</th>
<th>ε (mM$^{-1}$ cm$^{-1}$)</th>
<th>Φ</th>
<th>Brightness (mM$^{-1}$ cm$^{-1}$)$^a$</th>
<th>pK_a</th>
<th>Photostability (s)$^b$</th>
<th>Φ&lt;sub&gt;PC,GRX&lt;/sub&gt; / Φ&lt;sub&gt;PC,GR1&lt;/sub&gt;$^c$</th>
<th>E&lt;sub&gt;PC,GRX&lt;/sub&gt; / E&lt;sub&gt;PC,GR1&lt;/sub&gt;$^d$</th>
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<td>5.2</td>
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$^a$ Product of ε and Φ. For comparison, the brightness of mCherry is 16 mM$^{-1}$ cm$^{-1}$, and the brightness of EGFP is 34 mM$^{-1}$ cm$^{-1}$.29

$^b$ Time to photobleach from 1000 photons s$^{-1}$ molecule$^{-1}$ to 500 photons s$^{-1}$ molecule$^{-1}$ under wide-field and confocal illumination, respectively.

$^c$ Photoconversion quantum yield (calculated using Eq. (1)) relative to mClavGR1.

$^d$ Photoconversion contrast (calculated using Eq. (2)) relative to mClavGR1.

$^e$ Previously reported values.17,20,30

**Fig. 3.** Overview of the directed evolution of mClavGR. (a) A genealogy of ClavGR variants. The lines ending in filled circles indicate site-directed libraries that were created and screened but did not produce improved variants. An ‘X’ represents a selection of amino acids as specified in Supplementary Table 2. (b) Location of substitutions in mClavGR2 (relative to mClavGR1) that were introduced during the directed evolution process. The X-ray crystal structure of mTFP1 (Protein Data Bank ID 2HQK) is used here to represent mClavGR.24 (c) The dimer of amFP486 (Protein Data Bank ID 2A48) showing the location of valine 129, which is structurally homologous to residue 127 of mClavGR.24,28 As discussed in the text, substitutions at this position can influence the oligomeric state of the protein.
had resulted in the observed improvements in its properties, we undertook a thorough \textit{in vitro} characterization of mClavGR1 and mClavGR2 (Table 1). An overall conclusion from the \textit{in vitro} characterization experiments is that mClavGR2 has spectral properties that are comparable to those of the other members of the class of green-to-red photoconvertible FPs. \cite{17}

Prior to illumination, purified mClavGR2 protein has an excitation maximum at 488 nm and an emission maximum at 504 nm (Fig. 4a). Illumination with the same 405-nm light source used to photoconvert colonies of \textit{E. coli} causes the formation of a new species with primary excitation maximum at 566 nm and emission maximum at 583 nm. As represented in Fig. 1, it is the anionic form of the chromophore that is the primary fluorescent species in both the green fluorescent state and the red fluorescent state. Interestingly, the red fluorescent state has a second strong excitation peak at 355 nm.

Prior to illumination, the absorbance spectrum of mClavGR2 is dominated by two distinct peaks (Fig. 4b): one at 380 nm, which is attributable to the neutral phenol form of the chromophore, and the other at 488 nm, which is attributable to the anionic phenolate form of the chromophore (refer to Fig. 1). Of these two peaks, only the phenolate form also appears in the excitation spectrum (Fig. 4a), indicating that only this form is green fluorescent. The neutral phenol form is not green fluorescent but, as previously established by the action spectra of Kaede,\textsuperscript{1} EosFP,\textsuperscript{2} and Dendra2,\textsuperscript{21} is the form that can undergo photoconversion into the red state when excited (refer to Fig. 1). Indeed, recording of absorbance spectra at regular intervals during otherwise continuous illumination with the 405-nm light source revealed a monotonic increase in the 566-nm absorbance peak and a monotonic decrease in the 380-nm peak (Fig. 4b). The absorbance of the 488-nm peak also undergoes an eventual decrease with continued illumination, illustrating that the phenol and phenolate forms of the green state of mClavGR2 are in equilibrium and subject to the law of mass action.

An interesting property of mClavGR2 is that the 405-nm illumination used to photoconvert the neutral phenol form of the green state can also

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**Fig. 4.** Spectral changes in mClavGR2 upon photoconversion. (a) Excitation (cyan and orange) and emission (green and red) fluorescence spectra of the green state (before photoconversion) and the red state (after photoconversion) of mClavGR2, respectively. (b) Absorbance spectra versus photoconversion time of mClavGR2 in PBS (pH 7.4). The light source used for photoconversion is the 405-nm light-emitting-diode-based illumination chamber used for photoconverting the protein in bacterial colonies. (c) Brief illumination (10 s) with 405-nm light causes a substantial increase in the absorbance of the green fluorescent species.
temporarily drive the phenol/phenolate equilibrium of the green state towards the anionic form. Specifically, we observed that both the absorbance at 488 nm (Fig. 4c) and the overall fluorescence of the phenolate form increase immediately after illumination with 405-nm light. In the subsequent absence of the 405-nm light, the absorbance at 488 nm (and fluorescence at 504 nm) decreases to its original level over several tens of seconds. A similar phenomenon has also been reported to occur with Kaede\textsuperscript{21} and Dendra2.\textsuperscript{19} We speculate that this shift in equilibrium could be the result of photoinduced cis–trans isomerization of the chromophore similar to that observed to occur for the green state of the variant of EosFP known as IrisFP.\textsuperscript{20} An alternative explanation is that illumination shifts the phenol/phenolate equilibrium towards the phenolate state. However, if this were the case, we would expect equilibrium to be reestablished on a timescale of milliseconds as determined for analogous proton transfer processes in Aequorea GFP\textsuperscript{23} and Kaede.\textsuperscript{31} not on the tens of seconds that we observe in mClavGR2. We have not observed any reversible photoswitching or photoconversion of the red state of the chromophore.

Although both the green state and the red state of mClavGR2 are approximately five times brighter than those of mClavGR1 when expressed in E. coli, its inherent brightness (i.e., the product of ε and Φ) has improved only by a modest 7% and 42% for the green and red states, respectively (Table 1). This indicates that the improvement of brightness seen in E. coli is very likely due to a more complete maturation or a more efficient folding of the FP. mClavGR2 further benefits from a 20% increase in the quantum yield of photoconversion (Φ\textsubscript{PC}) relative to mClavGR1. The combination of the improved inherent brightness of the red state and the improved Φ\textsubscript{PC} provides an overall 80% improvement in photoconversion contrast (E\textsubscript{PC}). E\textsubscript{PC} is a relative measure of the overall intensity of red fluorescence that can be generated from a fixed concentration of protein molecules.

**Structure-based comparison of mClavGR2 and other green-to-red photoconvertible FPs**

Of the previously reported green-to-red photoconvertible FPs, mClavGR2 has the highest amino acid percentage identity with mEos2 (Fig. 2; Supplementary Tables 3 and 4).\textsuperscript{17} However, if one considers only those residues with side chains directed towards the interior of the β-barrel, mClavGR2 is most similar to mKikGR, with a relatively conservative amino acid difference of only 7 (versus 8 for Kaede, 9 for mEos2, and 13 for Dendra2) (Supplementary Tables 5 and 6). Of the green-to-red FPs reported to date, the two that are most similar with respect to the identity of residues directed towards the interior of the β-barrel are mEos2 and Kaede, with a difference of only 4. The two that are most divergent are Dendra2 and mKikGR, with a difference of 17. As mentioned above, the similar photophysical characteristics of the members of this class of FPs are attributed to the fact that they are all very similar with respect to their interior residues and chromophore environments.

One of the most distinctive amino acid differences in close proximity to the chromophore is the residue at position 73. In mClavGR2 and Dendra2 (numbered as residue 69), this residue is Thr; in mEos2 and Kaede, it is Ala; and in mKikGR, it is Val (Fig. 2; Supplementary Table 5). It has previously been suggested that hydrogen bonding between the side chain of Thr at this position and the guanidinium group of Arg70 (Arg66 in Dendra2) may hold the guanidinium farther from the chromophore and thus limit charge stabilization.\textsuperscript{21} This diminished degree of charge stabilization in Dendra2 has been used to explain both the blueshifted emission and the higher apparent pK\textsubscript{a} of the chromophore.\textsuperscript{21} We attempted to improve the properties of mClavGR1 by introducing substitutions at position 73. When we generated a library of all possible substitutions at position 73 and screened it in colonies to identify the variants that exhibited the highest green fluorescence before photoconversion and the highest red fluorescence after photoconversion, we found that the brightest colonies retained Thr at position 73. We also created the Thr73Ala and Thr73Val variants individually and characterized them. When expressed in E. coli, the green state of the Thr73Ala variant was approximately equivalent in brightness to the Thr73 progenitor, but the red (photoconverted) state was substantially dimmer. For the Thr73Val variant, both the green state and the red state were substantially dimmer than the original Thr73 variant.

There are only two residues of mClavGR2 that are directed towards the interior of the β-barrel and have amino acid identities that are not found in any of the other green-to-red photoconvertible FPs (Supplementary Table 5). These residues are Ile44 (Met, Ala, and Val in others) and Val183 (Ala and Ser in others). The valine at position 183 is the result of an amino acid substitution (Ala183Val) identified through screening of randomly mutated libraries for improved brightness and photoconversion. This is located near one end of the ninth β-strand of the protein and is relatively distant from the chromophore. Given its distant location, it is unlikely that Val183 has a direct influence on the properties of the chromophore and more likely contributes to improved brightness by increasing the folding efficiency of the protein. In contrast, Ile44 is in very close proximity to the chromophore and likely interacts with the side chain of His66, which is an integral part of the red chromophore. We generated a saturation library at Ile44 and again found that the best-performing variant remained to be Ile. We also created the Ile44Met and Ile44Val variants individually. We found that the Met substitution produced a variant with diminished photoconversion efficiency and that the Val substitution produced a variant with dimmer fluorescence in both the green state and the red state.
Phenol/phenolate equilibrium and photoconversion

The pH-dependent absorption spectra of the green and red states of mClavGR2 are shown in Supplementary Fig. 2. With apparent pK_a values of 8.0 and 7.3, respectively, both states of the chromophore exhibit substantial pH sensitivity in the physiological pH range (Table 1). It is notable that mClavGR2 has the highest pK_a reported for any of the green-to-red photoconvertible FPs. Due to our extensive empirical optimization, we expect the pK_a value for the green species to be near an optimal value with respect to photoconversion efficiency. That is, with this pK_a value, a substantial portion of the protein exists in the neutral phenol form at pH 7.4 and, thus, the overall rate of photoconversion is increased. A similar relationship between higher pK_a and enhanced photoconversion efficiency has been noted for both KikGR and Dendra2. In the case of KikGR, directed evolution for brighter red fluorescence following photoconversion resulted in a shift of the green-state pK_a from an initial value of 4.2 to a final value of 7.8.4 This change dramatically shifted the phenol/phenolate equilibrium towards the neutral form at pH 7.4. In the case of Dendra2 (pK_a = 7.1), it was noted that the 20-fold improvement in photoconversion efficiency relative to EosFP (pK_a = 5.8) is fully explained by the 20-fold higher concentration of the phenol form under physiological conditions.21 Presumably, the phenol/phenolate equilibrium in mClavGR2 could be shifted even further to the neutral form in variants with an even higher pK_a. In this case, the intensity of the green fluorescence would have been further diminished. Accordingly, the optimal pK_a of the green state represents a compromise between the desirable but conflicting demands of high photoconversion efficiency and high brightness of the green state.

While the pK_a of the green state of mClavGR2 may be close to optimal, it would be preferable to have a lower pK_a for the red state of mClavGR2. A lower pK_a is beneficial for two reasons: first, there are fewer potential artifacts due to local changes in pH; and, second, the protein is brighter due to a greater fraction of the protein existing in the red fluorescent phenolate form, as opposed to the nonfluorescent phenol form. In mTFP1 (pK_a = 4.3), the imidazole side chain of His163 is hydrogen bonded to the phenolate moiety of the chromophore.24,25 In mClavGR2, this residue has been substituted with a Met, which should be much less effective at stabilizing the anionic form and almost certainly contributes to the dramatically increased pK_a. In an effort to decrease the pK_a of the red state of mClavGR2, we generated the mClavGR2 Met163His variant, even though it seemed likely that this substitution would decrease the pK_a of both the green state and the red state. Not unexpectedly, the green state of this variant was found to exist entirely in the anionic phenolate form at physiological pH, and photoconversion thus proceeded with very poor efficiency.

Live cell imaging of various mClavGR2 fusion proteins

Given the results of our characterization experiments, it is apparent that there are some potential advantages and some potential disadvantages of mClavGR2 relative to other members of the class of green-to-red photoconvertible FPs with respect to applications in the imaging of dynamic protein localization. Most notable among the advantages of mClavGR2 are its monomeric structure, its highly optimized and relatively rapid folding efficiency (Supplementary Fig. 3), and its high photoconversion efficiency due to the high pK_a of the green state. The primary disadvantage is the high pK_a of the red state (Table 1; Supplementary Fig. 2), which causes the protein to have approximately half of its maximal fluorescent brightness under physiological pH conditions. It should be noted that the red state of mClavGR2, despite having only approximately half of its maximal brightness, retains brightness slightly greater than the red state of the popular red FP known as mCherry at pH 7.4 (Table 1).35

Another advantage of mClavGR2 that became apparent while we were performing photobleaching experiments with protein expressed in live cells is the improved photo-stability of the red state of the protein under confocal illumination conditions relative to other green-to-red photoconvertible FPs. The time to bleach the red state to 50% of the initial intensity for mClavGR2 (3644 s; see Table 1 and Supplementary Fig. 4) represents a substantial improvement over mEos2 (2700 s) and Dendra2 (2420 s).17 Oddly, this advantage does not extend to wide-field illumination conditions where the time to bleach to 50% of the initial intensity for mClavGR2 (175 s) is somewhat lower than those reported for mEos2 (323 s) and Dendra2 (378 s). It should be noted that the red state of mKikGR exhibits relatively rapid photobleaching under both wide-field (21 s) and confocal (530 s) illumination conditions. The time to bleach the green state of mClavGR2 to 50% of its initial intensity using confocal illumination (233 s) is only slightly faster than those of both mEos2 (240 s) and Dendra2 (260 s), but substantially slower than that of mKikGR (80 s). For wide-field photobleaching, mClavGR2 (17 s) is more similar to mKikGR (14 s) than it is to the more photostable mEos2 (42 s) and Dendra2 (45 s) variants.

In an effort to determine if mClavGR2 is potentially useful for the imaging of dynamic protein localization, we have assembled mammalian expression plasmids for a variety of mClavGR2 constructs, used these plasmids to transfect cultured mammalian cells, and acquired fluorescence images of the resulting cells both before photoconversion and after photoconversion. Following expression of mClavGR2 in HeLa cells as an unfused protein, we were able to photoconvert the protein in a localized region and to image the subsequent diffusion of the red-state protein as it evenly dispersed throughout the cytoplasm over a timescale of approximately 30 s.
Fig. 5. Photoconversion of unfused mClavGR2 in live cells. (a) Green fluorescence image from two cells expressing mClavGR2 prior to photoconversion. (b) The same cells as in (a) imaged using a filter set for red fluorescence prior to photoconversion. The three areas of interest are indicated with a circle, diamond, and square. The area indicated with a square corresponds to the location where the cell is illuminated with a 408-nm laser from \(t = -2\) to 0 s. (c) Mean red fluorescence intensities in each of the areas of interest marked in (b). (d) A series of images showing the free diffusion of photoconverted mClavGR2. Indicated times represent time elapsed since the end of the 408-nm laser illumination.

This experiment established that mClavGR2 did not exhibit any anomalous localization when expressed alone in cells and was capable of free and rapid diffusion in the cytoplasm.

Encouraged by these promising results, we proceeded to construct fusion constructs in order to test whether mClavGR2 could be fused to different protein partners and targeting sequences without causing mistargeting or possible misfolding of either protein. In Fig. 6, we show images of the green state of mClavGR2 in 16 commonly used fusion constructs. These localization patterns for all fusion constructs were judged to be indistinguishable from the localization patterns of the analogous Aequorea GFP constructs imaged under identical conditions in the same laboratory. The favorable performance of mClavGR2 in these constructs serves as ancillary evidence of its monomeric state. Notably, the fine filament structure of the actin cytoskeleton is clearly preserved with the mClavGR2-β-actin fusion expressed in HeLa cells (Fig. 6a). For histone 2B (H2B), connexin 43 (Cx43), and Rab5a fusion constructs, we performed localized photoconversion and subsequent imaging of the dynamic localization of both green fluorescence and red fluorescence. Two-color image sequences for these experiments are provided as Supplementary Figs. 5–7, and time-lapse movies are available online as Supplementary Movies 1–3. These experiments established that mClavGR2, when fused to other proteins, causes minimal perturbation of localization and trafficking dynamics. Furthermore, the green and red states of mClavGR2 have sufficient brightness and photostability for long-term imaging experiments.

To further test the utility of mClavGR2 and to fully exploit the advantages provided by its monomeric structure, we investigated whether mClavGR2 could be used to monitor the dynamics of a transmembrane adhesion protein, intercellular adhesion molecule 1 (ICAM-1). We specifically asked whether we could quantitatively assess changes in the lateral mobility of ICAM-1 on the surface of a mammalian cell while engaged in cell–cell contact with a T-lymphocyte (Jurkat) cell expressing the ICAM-1 receptor leukocyte-function-associated antigen 1 (LFA-1). ICAM-1 lateral mobility has been shown to be mediated by cortactin, filamin B, and adhesion receptor microdomains.36–38 Fluorescent images of HeLa cells transiently transfected with a plasmid encoding ICAM-1-mClavGR2 revealed an unambiguous pattern of surface FP localization (Fig. 7a) consistent with the expected localization of ICAM-1 in the plasma membrane. Incubation of these adherent cells with a suspension of Jurkat cells resulted in the formation of cell–cell contacts where a Jurkat cell had settled onto the plasma membrane of a HeLa cell. Bright-field imaging was used to identify Jurkat cells in apparent physical contact with transfected HeLa cells (Fig. 7b). For each apparent contact, the existence of specific ICAM-1/LFA-1-mediated contact was confirmed by locating a bright cluster of fluorescence at the contact point suggested by the bright-field images (Fig. 7c and d). Localized photoconversion at the point of contact was performed, and a time series of red fluorescence images was acquired to trace the diffusion of ICAM-1 initially found at the contact zone (Fig. 7e). Identical experiments were performed at arbitrary locations on the plasma membrane of transfected HeLa cells in the absence of Jurkat cells. Postcollection analysis was used to extract the fluorescence intensity at the contact point as a function of time (Fig. 8a).
Our analysis of ICAM-1 diffusion revealed that the diffusion half times increased from 6.6±0.4 s for proteins not involved in cell-cell contacts to 22.9±3.7 s for proteins involved in cell-cell contacts (Fig. 8b). This result suggests an almost 4-fold reduction in the rate of diffusion of ICAM-1 found within a contact zone versus ICAM-1 found elsewhere on the cell surface. Regardless of whether or not ICAM-1 was in adhesive contact, approximately 25% (24±4% or 26±2%, respectively) of the protein was effectively immobile over the 100-s timescale of the experiment (Fig. 8b). These results are consistent with previous reports indicating that ICAM-1 mobility is reduced by cross-linking of the receptor, although cross-linking here is induced by the presentation of receptors as part of the lymphocyte cell surface. These results unambiguously demonstrate that photoconversion of mClavGR2-containing constructs will be a powerful tool for dissecting the mechanisms that mediate cellular adhesion by allowing long-term tracing of receptors and lateral diffusion measurements.

**Summary**

We have developed and characterized a new green-to-red photoconvertible FP variant known as...
mClavGR2. Unlike other members of this class of FP, mClavGR2 was engineered from a well-characterized and monomeric FP progenitor. Our results demonstrate that mClavGR2 has favorable spectral properties, retains the monomeric structure of its progenitor, and, in the constructs tested to date, does not interfere with the correct localization of a genetically fused protein partner. We anticipate this new addition to the toolbox of engineered FPs to be of great utility in imaging fast protein dynamics in live cells. Experiments aiming to determine whether the advantages of mClavGR2 translate into improved performance in superresolution imaging applications have been initiated.16

Materials and Methods

General methods and materials

All synthetic DNA oligonucleotides for cloning and library construction were purchased from Integrated DNA Technologies (Coralville, IA). PCR products and products of restriction digests were purified using the QIA

Fig. 7. Photoconversion of ICAM-1-mClavGR2 on a HeLa cell in contact with a Jurkat cell expressing LFA-1. (a) Green fluorescence image of a HeLa cell in contact with a Jurkat cell. The arrow indicates the location of the Jurkat cell as shown in (b). Brighter fluorescence is observed at the cell–cell contact point due to the interaction of LFA-1 on the surface of the Jurkat cell with ICAM-1-mClavGR2 on the HeLa cell. The white circle at the cell–cell contact is the area to be photoconverted. Diagonal lines marked with a diamond and a circle correspond to the reconstructed x–z cross sections represented in (c) and (d). (b) Bright-field image of the same field of view as (a) with focus adjusted to show the Jurkat cell (white arrow). (c) Cross-sectional view of cells shown in (a) at the line indicated with a circle. (d) Cross-sectional view of the cell shown in (a) at the line indicated with a diamond. (e) Red fluorescence image of the same field of view as (a) prior to photoconversion. (f) Time-lapse images of the red fluorescence following photoconversion.

Fig. 8. Diffusion of ICAM-1-mClavGR2 on the plasma membrane. (a) Red fluorescence intensity at the location of photoconversion versus time after photoconversion for HeLa cells either at the point of adhesion with Jurkat cells (circles) or at an arbitrary point on the plasma membrane in the absence of Jurkat cells (squares). Data represent the average results from 8 and 22 independent measurements, respectively. Lines represent the best fit of the data using Eq. (3). (b) Summary of diffusion times (t_{1/2}) as obtained by fitting of the data provided in (a). The t_{1/2} values for the two data sets are significantly different (p<0.01). The ‘adhesion’ label refers to whether or not the site of photoconversion was within an adhesion contact, as identified in bright-field and fluorescence images.
gel extraction kit (Qiagen) in accordance with the manufacturer’s protocols. Restriction enzymes were purchased from New England Biolabs. The cDNA sequences for all mClavGR variants were confirmed by dye terminator cycle sequencing using the DyeCycle ET Kit (Amersham Biosciences) or the BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems). Sequencing reactions were analyzed at the University of Alberta Molecular Biology Service Unit or the Florida State University Department of Biological Science DNA Sequencing Facility.

**Plasmid library creation and site-directed mutagenesis**

For production of the designed cDNA encoding mClavGR1, 14 forward oligonucleotides and 14 reverse oligonucleotides with 51 bases each were synthesized and assembled using circular assembly amplification.

A forward primer with an XhoI restriction site and a reverse primer with an EcoRI restriction site were used to amplify the cDNA, which was subsequently digested and ligated into a similarly digested pBAD/His B vector (Invitrogen). Randomly mutated libraries were created by error-prone PCR, where dNTPs mixed with a deficient component were used as template mix. The mixture was subjected to 100 cycles (94 °C for 45 s, then 55 °C for 5 s), followed by an extra extension step at 72 °C for 5 min. Regardless of library assembly method (i.e., error-prone PCR, saturation mutagenesis, or STEP PCR), gene libraries were inserted into pBAD/His B between the XhoI site and the EcoRI site, as described above. Electrophoretic E. coli strain DH10B (Invitrogen) was transformed and plated on LB agar plates supplemented with ampicillin (0.1 mg/ml) and t-arabinose (0.2%). Plates were incubated for 14 h at 37 °C prior to screening.

**Plasmid library screening**

E. coli colonies expressing the mClavGR libraries were grown on 10-cm Petri dishes. In order to screen libraries for variants that were brighter and exhibited more efficient photoconversion, we developed a method for photoconverting FPs expressed in colonies of E. coli. Our method involves a three-step process. In the first step, a custom imaging system equipped with filter sets (Chroma) for 470/40 nm excitation with 510/20 nm emission (i.e., for green fluorescence) and filter sets for 560/40 nm excitation with 630/60 nm emission (i.e., for red fluorescence) is used to acquire both green and red fluorescence images of the Petri dish. In the second step, the dish is placed in a ‘photoconversion chamber,’ where it is evenly illuminated with light from six 9 x 11 arrays of 405-nm light-emitting diodes (OptoDiode Corporation, Newbury Park, CA). Following 10-20 min of illumination, the dish is imaged again using the same filters used in the first step. The digital images are loaded into Image Pro plus software (Media Cybernetics, Bethesda, MD), where they are aligned and processed using a custom macro. The output of the macro is a list of the mean intensities for each colony in each of the four images. These data are imported into Microsoft Excel, where they are plotted as green fluorescence before photoconversion versus red fluorescence after photoconversion. Colonies that exhibited the highest intensities in both channels were picked and used as templates for the following round of library creation.

**Protein purification and characterization**

Protein purification was carried out as previously described, and the buffer was changed into phosphate-buffered saline (PBS; pH 7.4). Samples of mClavGR2, Dendra2, and mEos2 used for determination of oligomeric structure were first subjected to an additional step of purification on a HiLoad 16/60 Superdex 75 pg column (GE Healthcare). Molar extinction coefficients (ε) of the green states were measured by alkali denaturation method and then used as reference to measure ε for the red states.

Fluorescence quantum yields (Φ) were determined using fluorescein in 10 mM NaOH (Φ = 0.95) and Rhodamine 6G in ethanol (Φ = 0.94) as standards. For determination of pH dependence, purified protein in PBS was diluted 1:50 into a series of pH-adjusted citrate saline (pH ≤ 8) or sodium phosphate (pH > 8) buffers in a 96-well black clear-bottom plate (Corning). Fluorescence was measured using a Safire2 plate reader (Tecan).

The oligomeric structures of mClavGR variants, mEos2, and Dendra2 were determined by gel-filtration chromatography with a HiLoad 16/60 Superdex 75 pg gel-filtration column on an AKTA basic liquid chromatography system (GE Healthcare). Samples of dimeric dTomato and monomeric mCherry proteins were expressed and purified in the same way as mClavGR variants and used as size standards. Purified mClavGR variants were mixed with dTomato or mCherry, and the resulting elution profiles were monitored at 488 nm for mClavGR variants, at 555 nm for dTomato, and at 585 nm for mCherry.

To measure the maturation profiles of mClavGR1, mClavGR2, mEos2, and Dendra2 (Clontech), we cultured E. coli transformed with pBAD/His B plasmids bearing the encoding cDNA overnight in glucose-supplemented LB media to prevent gene expression. The culture was diluted to an OD<sub>600</sub> of 0.6, purged with argon for 20 min, sealed with a rubber septum, and incubated for another 1 h to allow thorough consumption of the residual oxygen. Arabinose (0.25%) was then added via a syringe with needle to induce expression of the FPs. After 4.5 h of incubation with shaking at 37 °C, the cultures were transferred to an ice bath for 10 min and maintained at 4 °C, unless indicated otherwise. Cells were centrifuged at 8000 rpm for 2 min, and cell pellets were lysed using vacuum-degassed B-PER II (Fierce) and incubated at room temperature for 12 min. The lysate was then centrifuged at 12,500 rpm for 5 min, and the supernatant was diluted five times into PBS (pH 7.4). Fluorescence maturation was monitored at 37 °C using a Safire2 plate reader (Tecan).

To determine the relative quantum yields for photoconversion, we measured each purified variant’s absorbance at 487 nm to determine the concentration. The proteins were then diluted 10-fold into PBS, and their red fluorescence was recorded. The photoconversion quantum yield relative
to mClavGR1 \((\Phi_{PC,GRX}/\Phi_{PC,GR1})\) was calculated using Eq. (1):

\[
\frac{\Phi_{PC,GRX}}{\Phi_{PC,GR1}} = \frac{C_{red,GRX}}{C_{green,GRX}} \cdot \frac{C_{green,GR1}}{C_{red,GR1}} = \frac{A_{565,GRX}}{A_{487,GRX}} \cdot \frac{A_{487,GR1}}{A_{565,GR1}}
\]

In Eq. (1), \(A_{565}\), \(A_{487}\), \(c_{487}\), and \(c_{565}\) are the absorbance and extinction coefficients at 487 nm (before photconversion) and 565 nm (after photconversion), respectively. GR1 refers to mClavGR1, while GRX refers to one of the other variants. This relative photconversion quantum yield gives the frequency of generation of the red species of the evolved variants relative to the parent mClavGR1 when illuminated by the screening system. Relative photconversion efficiency \((E_{PC,GRX}/E_{PC,GR1})\) was calculated using Eq. (2), again with mClavGR1 as reference:

\[
\frac{E_{PC,GRX}}{E_{PC,GR1}} = \frac{I_{GRX}}{c_{GRX}} \cdot \frac{I_{GR1}}{c_{GR1}}
\]

In Eq. (2), \(c\) is the concentration of the purified protein solution, and \(I\) is the red fluorescence intensity following photconversion. This relative photconversion efficiency can differ from the photconversion yield due to additional factors such as the ratio of neutral form to anionic form at the beginning and the brightness of the red state. All absorption measurements were acquired on a DU-800 UV–visible spectrophotometer (Beckman). All fluorescence spectra were recorded on a QuantMaster spectrofluorimeter (Photon Technology International) and corrected for detector response.

To create the mClavGR2 fusion expression vectors, we amplified the gene-encoding mClavGR2 with a 5′ primer with an Nh el site and a 3′ primer with an XhoI site. The purified PCR products were then digested and ligated into pEGFP-actin and pEYFP-NLS (Clontech), whose FP-coding gene has been previously removed by the same restriction enzymes. To create the mClavGR2 vector for cytoplasmic expression, we inserted the mClavGR2 gene between the XhoI restriction site and the EcoRI restriction site of a modified pcDNA3 vector (Invitrogen) in which the multiple cloning site had been modified to contain XhoI and EcoRI restriction sites in the same reading frame as the identical sites in pBAD/His B. The full-length ICAM-1 (Addgene plasmid 8632) and mClavGR2 cDNA were incorporated into the modified pcDNA3 vector mentioned above, with a 24-amino-acid linker between them. The linker’s sequence is KALSAAAAGGGSGGGCGGCGGSEY. All other mClavGR2 fusion expression vectors were constructed using C1 and N1 (Clontech style) cloning vectors, as previously described. The mClavGR2 cDNA was amplified with a 5′ primer encoding an AgeI site and a 3′ primer encoding either a BspEI (C1) site or an NotI (N1) site. The purified and digested PCR products were ligated into similarly digested pEGFP-C1 and pEYFP-N1 expression vector backbones. To generate fusion vectors, we digested the appropriate mClavGR2 cloning vector and an EGFP fusion vector, either sequentially or doubly, with the appropriate enzymes and ligated them together after gel purification. Thus, to prepare mClavGR2 N-terminal fusions, we performed the following digests: human nonmuscle α-actinin, EcoRI and NcoI (fusion cDNA source; Tom Keller, The Florida State University); rat α-1-Cx43, EcoRI and BamHI (Matthias Falk, Lehigh University); human H2B, BamH I and NotI (George Patterson, National Institutes of Health); N-terminal 81 amino acids of human β-1,4-galactosyltransferase, BamHI and NotI (Golgi; Clontech); human keratin 18, EcoRI and NotI (Open Biosystems, Huntsville, AL); human zyxin, BamH I and NotI (Clare Waterman, National Institutes of Health); lifeact, BamH I and NotI (Integrated DNA Technologies); human VE cadherin (Origene, Rockville, MD); chicken c-src (Marilyn Resh, Sloan-Kettering, New York). To prepare mClavGR2 C-terminal fusions, we performed the following digests: human β-actin, Nhel and BglII (Clontech); human lamin B1, Nhel and BglII (George Patterson, National Institutes of Health); human flierlinin, Agel and BglII (Evrogen); mouse
CAF1, Nhel and BglII (Akash Gunjan, The Florida State University); mouse microtubule binding domain of MAP4, Nhel and BglII (Richard Cyr, Penn State University); endoplasmic reticulum, Agel and BspEI (Integrated DNA Technologies); c-Ha-Ras (CAAX), Agel and BspEI (membrane; Clontech). DNA for mammalian transfection was prepared using the Plasmid Maxi kit (Qiagen). Morphological features in all fusion constructs were confirmed by imaging fixed cell preparations on coverslips using a Nikon 80i upright microscope with ET-GFP and/or ET-DsRed filter sets (Chroma) coupled to a Hamamatsu Orca ER camera.

**Cells and transfection**

HeLa cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Invitrogen) supplemented with 10% heat-inactivated fetal calf serum (FCS; Invitrogen) and 2 mM GlutaMax (Invitrogen) at 37 °C and 5% CO2. Jurkat cells (clone E6.1) were maintained in RPMI 1640 (Invitrogen) supplemented with 10% FCS. Plasmid DNA for transfection was prepared by Qiagen Plasmid Midi Kit (Qiagen) in accordance with the manufacturer’s instructions. Lipofectamine 2000 (Invitrogen) was used as transfection reagent. Cells for transfection and imaging were cultured on 35-mm glass-bottom culture dishes. The transfection protocol was essentially performed as recommended by the manufacturer, but used 3 μg of plasmid DNA, 7.5 μl of transfection reagent, and 2.5 h of incubation time (incubation of the cell with the transfection reagent). The medium was changed into Hepes-buffered Hank’s buffered salt solution (Invitrogen) before imaging. Imaging was performed 24–36 h after transfection. To examine the localization of fusions and dynamics in live cell imaging and photobleaching measurements, we grew HeLa or LLC-PK1 (pig kidney epithelial; ATCC, Manassas, VA) cells in a mixture of DMEM and Ham’s F-12 (50:50) containing 12% Cosmic calf serum (Georgia Biowest, Manassas, VA) cells in a mixture of DMEM and Ham’s F-12 (50:50) containing 12% Cosmic calf serum (HyClone) to 70% confluence and transfected them with 1 μg of the appropriate vector DNA using Effectene (Qiagen) in accordance with the manufacturer’s instructions. The medium was changed after 24 h, and cells exhibiting normal morphology were imaged after 36 h.

**Live cell imaging**

Fluorescence microscopy of mClavGR2 diffusion in the cytoplasm was performed using an inverted Nikon Eclipse Ti microscope equipped with a QuantEM 512SC electron-multiplying CCD (Photometrics), a 75-W xenon lamp for epifluorescence illumination, a 63× immersion objective (Carl Zeiss), an electron-multiplying CCD (Hamamatsu), and a 20-mW 405-nm laser (Melles Griot). Temperature was maintained at 37 °C during acquisition. An exposure of approximately 40 ms at 15% laser power (405 nm) was used to photoconvert mClavGR2 within the defined area of interest. Four images of each of the green and red channels were acquired before photoconversion, followed by 10 s of acquisition at maximum speed, followed by 2 min at 3 s per frame for 2 min. The average intensity of the four images acquired before photoconversion was subtracted from the background-corrected red fluorescence intensity, which was then normalized to the first image after photoconversion. Fluorescence as a function of time (t(t)) was fitted with the equation:

\[
F(t) = A + B \exp\left(-t / \tau\right)
\]

where A and B correspond to the fractional contributions of immobile and mobile proteins, respectively, and \( \tau \) is the time constant for the decay of fluorescence intensity. The diffusion half time \( \left(t_{1/2}\right) \) is 0.69\( \tau \).

**Supplementary Data**

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2010.06.056

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**Competing Financial Interests Statement.** New FPs that originate from the Campbell lab and are described in this manuscript are covered by a US patent application owned by the University of Alberta. Allele Biotechnology is the Licensed distributor of plasmids containing genes encoding these FPs.
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