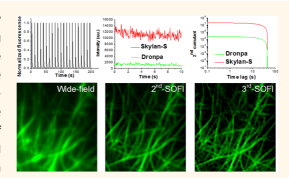


Development of a Reversibly Switchable Fluorescent Protein for Super-Resolution Optical Fluctuation Imaging (SOFI)

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ABSTRACT Reversibly switchable fluorescent proteins (RSFPs) can be effectively used for super-resolution optical fluctuation imaging (SOFI) based on the switching and fluctuation of single molecules. Several properties of RSFPs strongly influence the quality of SOFI images. These properties include (i) the averaged fluorescence intensity in the fluctuation state, (ii) the on/off contrast ratio, (iii) the photostability, and (iv) the oligomerization tendency. The first three properties determine the fluctuation range of the imaged pixels and the SOFI signal, which are of essential importance to the spatial resolution, and the last may lead to artificial aggregation of target proteins. The RSFPs that are currently used for SOFI are low in



averaged fluorescence intensity in the fluctuation state, photostability, and on/off contrast ratio, thereby limiting the range of application of SOFI in biological super-resolution imaging. In this study, we developed a novel monomeric green RSFP termed Skylan-S, which features very high photostability, contrast ratio, and averaged fluorescence intensity in the fluctuation state. Taking advantage of the excellent optical properties of Skylan-S, a 4-fold improvement in the fluctuation range of the imaged pixels and higher SOFI resolution can be obtained compared with Dronpa. Furthermore, superresolution imaging of the actin or tubulin structures and clathrin-coated pits (CCPs) in living U2OS cells labeled with Skylan-S was demonstrated using the SOFI technique. Overall, Skylan-S developed with outstanding photochemical properties is promising for long-time SOFI imaging with high spatial-temporal resolution.

KEYWORDS: reversibly switchable fluorescent protein · super-resolution microscopy · Skylan-S · protein engineering · live-cell imaging · SOFI

Invescence microscopy has become one of the most important biophysical tools for exploring molecular events that occur during subcellular processes in the life sciences. Unfortunately, it has encountered certain challenges in discerning smaller cellular organelles because of the resolution barrier imposed by optical diffraction, which leads to poor imaging of objects smaller than approximately half the wavelength of the probe light. Consequently, a super-resolution technique that is able to overcome the diffraction barrier is highly desirable for further biological research. Various super-resolution techniques have been developed over the past decade, and these techniques can be categorized into three approaches:¹ (1) point spread function modulation, which includes such techniques as stimulated emission depletion (STED),² reversible saturable optical fluorescence transitions (RESOLFT),³ and saturated structured illumination microscopy (SSIM);⁴ (2) single-molecule localization, which includes such techniques as photoactivated localization microscopy (PALM)^{5,6} and stochastic optical reconstruction microscopy (STORM);⁷ and (3) blinking/fluctuation statistics, which includes such techniques as super-resolution optical fluctuation imaging

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(SOFI)^{8,9} and spatial covariance reconstructive (SCORE) super-resolution fluorescence microscopy.¹⁰ Unlike the other techniques which require specialized equipment, SOFI is a purely calculation-based imaging approach that can achieve a 3- to 4-fold improvement in spatial resolution in all three dimensions using a conventional wide-field microscope.^{8,9} SOFI can produce backgroundfree, contrast-enhanced super-resolution images based on the temporal correlation analysis of fluorescence fluctuation/blinking over hundreds of raw images, which is far fewer than those required for PALM/STORM, even under conditions of low signal-to-noise and high background.^{8,9,11} This technique initially used quantum dots and organic fluorophores in a carefully controlled cell-toxic buffer, which hindered its application in most living biological settings.¹² To overcome this limitation, photochromic stochastic optical fluctuation imaging (pcSOFI) has been developed using a reversibly switchable fluorescent protein (RSFP) known as Dronpa; this technique provides a 2- to 3-fold enhancement in spatial resolution, significant background rejection, markedly improved contrast, and favorable temporal resolution in living cells.13

Several properties of RSFPs strongly influence the quality of SOFI images. These properties include (i) the averaged fluorescence intensity in the fluctuation state for each pixel among sequential frames and (ii) the on/off contrast ratio. These two properties affect the dynamic ranges of the fluorescence fluctuation of the imaged pixels, which, in turn, determine the spatial resolution of the resulting SOFI images. On the one hand, for each pixel, a higher dynamic range of fluorescence fluctuation will produce a higher SOFI signal. On the other hand, a higher dynamic range of the fluorescence fluctuation of one pixel will cause the adjacent pixels to be more strongly affected and produce higher cross-cumulants, meaning that less heterogeneity in the molecular intensity of the adjacent pixels can be obtained. Such heterogeneity is extremely important for SOFI calculations at higher orders, predominantly because SOFI at order n raises the heterogeneities in molecular intensity to the power of n; at orders higher than 4, dimmer and/or weakly fluctuating molecules become lost in the background, and the imaged structures lose connection.¹¹ Thus, a high SOFI signal resulting from a high dynamic range of fluctuation will produce a continuous structure by equalizing the blinking heterogeneities. (iii) The third property that affects SOFI quality is the photostability of the RSFPs. SOFI is based on the temporal correlation analysis of fluorescence fluctuations over a sequence of hundreds of raw images. These fluctuations must be sufficiently slow compared with the acquisition time (tens of milliseconds in practice) to be resolvable,¹³ and they must be sufficiently persistent to be observable over the full measurement duration. Therefore, this technique requires RSFPs to be fairly stable and not to become bleached before the reconstruction of the super-resolution images can be achieved. (iv) The fourth property that affects SOFI quality is the oligomerization tendency of the RSFPs. Dimerization is an undesirable characteristic of fluorescent proteins and may lead to artificial aggregation of the target proteins. At present, several RSFPs have been reported for SOFI super-resolution imaging.¹⁴ However, these RSFPs suffer from low averaged fluorescence intensity in the fluctuation state, photostability, and on/off contrast ratios or produce low SOFI signals, which limits the range of application of SOFI in biological superresolution imaging.

In previous studies, we developed several unique RSFPs, referred to as the mGeos family. These RSFPs were developed based on an irreversible green-to-red photoconverter protein, mEos2, by mutating the first position of the tripeptide (HYG) of the chromophore (His62) of mEos2.¹⁵ We then developed two real monomeric photoconvertible proteins by examining the residue-residue interactions at two interfaces and mutating the three key residues that may participate in oligomerization, named mEos3.1 and mEos3.2, which are suitable for PALM imaging.¹⁶ In the current study, we used the same mutation strategy that was used in the development of mGeos and performed saturation mutations at the first position of the tripeptide in the chromophore (His62) of mEos3.1. We found that many of the mutants yielded green species with no conversion ability. As in the case of mGeos, the His62 mutants of mEos3.1 can be switched on by 405 nm light and switched off by 488 nm light over many cycles, and they constitute a unique series of green RSFP variants (data not shown). There is one mutant (mEos3.1 H62S) that exhibits very high averaged fluorescence intensity in the fluctuation state, photostability, and on/off contrast ratio, suggesting that it is highly suitable for application in SOFI imaging (Figure S1). We named this mutant Skylan-S (Sky lantern for SOFI) and investigated its photochemical properties and its performance in SOFI imaging.

RESULTS AND DISCUSSION

To evaluate the properties of Skylan-S, we expressed Skylan-S proteins in bacteria and determined the bulk spectroscopic properties of purified Skylan-S; we also tested Dronpa as a control, as its SOFI performance has been previously established. The excitation, emission, and absorption spectra of Skylan-S do not differ significantly from those of Dronpa, as shown in Figures 1a and S2; however, Skylan-S exhibits higher brightness (115% and 134% of the brightnesses of Dronpa and mEos3.1, respectively), as shown in Table 1. To determine the contrast ratio, we measured the fluorescence intensity of E. coli cells expressing Skylan-S under sequential illumination with 405 nm light (switching on) and 488 nm light (switching off). As shown in



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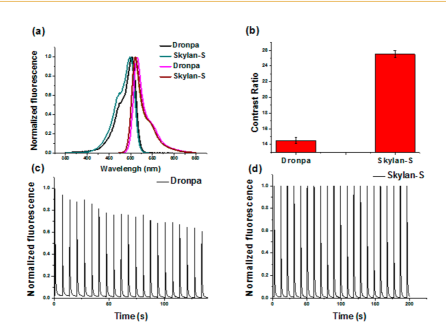


Figure 1. (a) Excitation and emission spectra (measured at the excitation maximum). The excitation and fluorescence emission spectral profiles of Skylan-S and Dronpa displayed similar maxima at approximately 490 and 510 nm, respectively. (b) Contrast ratios of Dronpa and Skylan-S. (c,d) Photoswitching kinetics of Dronpa and Skylan-S, respectively, measured in *E. coli. E. coli* cells expressing Dronpa and Skylan-S were continuously excited with 488 nm illumination (0.24 mW), and pulsed 0.01 s bursts of illumination with 405 nm light at a laser power of 0.2 mW were used to photoactivate the fluorescent proteins. The fluorescence was recorded continuously for 21 cycles.

TABLE 1. Characteristics of Dronpa, mEos2, mEos3.1, and Skylan-S:

	abs.	emi.		€-max ^b		MW ^c	oligomerization ^d
	(nm)	(nm)	QY ^b	$(M^{-1} cm^{-1})$	brightness	(kDa)	[<i>K</i> _d]
Dronpa ^a	503	522	0.68	125000	85	23.0	monomer
mEos2 (G)	504	515	0.84	56000	47	42.8	weak dimer
mEos3.1 (G)	505	513	0.83	88400	73	29.4	monomer
Skylan-S	499	511	0.64	152408	98	23.8	monomer

^{*a*} Literature values in refs 15 and 25. The quantum yield (QY) and extinction coefficient (ϵ) of Skylan-S were measured in PBS (pH = 7.4) for their on-states. ^{*b*} The quantum yield (QY) and extinction coefficient (ϵ) were measured in PBS (pH = 7.4) for the on-states of the proteins. Abs: maximum absorption; Emi: maximum emission; QY: quantum yield; ϵ -max: extinction coefficient. The brightness was calculated as extinction coefficient \times quantum yield/1000. Please note that the brightness here is corresponding and near to the peak intensity before photobleaching to a low fluorescence plateau state with fluctuation, and is not the averaged fluorescence intensity in the fluctuation state that is used to reconstruct a SOFI image. ^c The molecular weight (MW) was measured via sedimentation equilibrium. ^{*d*} K_d was measured via nonlinear least-squares fitting using the Microcal Origin software package. "Monomer" indicates a very high K_d that was beyond the limits of our instrumentation.

Figure 1b, Skylan-S has a much higher contrast ratio than Dronpa. A comparison of the 405/488 nm-illuminationinduced photoswitching properties of Skylan-S and Dronpa was performed by conducting *in vivo* photoswitching assays of the two fluorescent proteins expressed in *E. coli* cells. Dronpa exhibited a level of photodecay of approximately 35% after 21 switching cycles, whereas Skylan-S retained nearly 100% photostability without any photodecay after the same switching cycles (Figure 1c,d). The extremely high photostability of Skylan-S is expected to allow for sufficient switching cycles for SOFI and observable fluctuations throughout the entire acquisition time. The switching speed properties were calculated from Figure 1c,d and are presented in Supplementary Table 1. To determine more specific properties of Skylan-S related to SOFI imaging, time-resolved fluorescence spectroscopy was performed by using the time-correlated single photon counting method. The fluorescent decay time was measured to be 3.94 ns (Figure S3). The relaxation half-time of Skylan-S was 139 min (Figure S4), shorter than Dronpa (840 min).¹⁵ The absorption spectra of purified Skylan-S at on-state and off-state are shown in Figure S5.

Dimerization of fluorescent proteins may cause undesired aggregation of the target proteins. As we mutated only the amino acid inside the mEos3.1 protein, Skylan-S was expected to present the same oligomeric character as mEos3.1. Indeed, our gel filtration analysis, ultracentrifuge analysis and sedimentation velocity analysis of purified Skylan-S indicated that Skylan-S behaved as a true monomer (Table 1 and Figures S6 and S7). To further investigate the performance of Skylan-S as a fusion tag in living cells, we observed the localizations of Skylan-S fusion proteins transiently expressed in U2OS cells using confocal microscopy. As shown in Figure 2, Skylan-S can be used to accurately label β -actin, caveolin, clathrin, and microtubule. Next, we investigated a calcium channel, ORAI1, which is expressed in the plasma membrane and is sensitive to the oligomeric character of fluorescent protein tags.^{16,17} We found that Skylan-S exhibited a similar pattern to that of mEos3.1 without any

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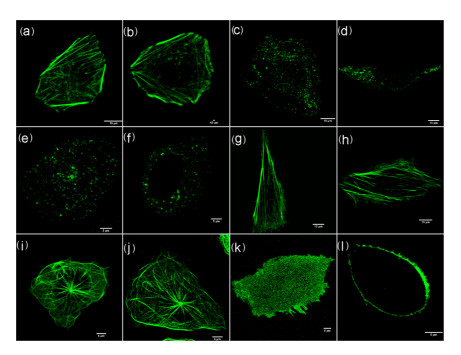


Figure 2. Confocal images of U2OS cells expressing Skylan-S-labeled (a,b) β -actin, (c,d) caveolin, (e,f) clathrin, (g,h) Lifeact, (i,j) MAP4, and (k,l) ORAI1: (k) section near the plasma membrane and (l) middle section.

artificial aggregation (Figure 2k,l). These results suggest that Skylan-S is an excellent fusion tag for labeling proteins in living cells.

For SOFI imaging, robust fluctuation of each imaged pixel is necessary. As mentioned in the introduction, higher fluctuation will produce a higher SOFI signal and a more continuously imaged structure. In every pixel at each imaging time point, the measured intensity is the sum of the intensities contributed by several adjacent RSFP molecules. Thus, the fluctuation is predominantly determined by the photochemical properties of the RSFPs, such as averaged fluorescence intensity in the fluctuation state and contrast ratio. Therefore, we examined the pixel fluctuations using fixed cells expressing Skylan-S-labeled Lifeact under various excitation (488 nm) laser powers and various exposure times. We found that Slylan-S exhibited excellent fluorescence fluctuation, and the pixel fluctuation is strongly dependent on both the intensity of the excitation laser power and the exposure time. Higher excitation intensity will produce a higher dynamic range of fluctuation but also more photobleaching of the Skylan-S (Figure S8). We chose a power of 2.8 μ W for the 488 nm laser and an exposure time of 40 ms for SOFI imaging using Skylan-S in our homemade TIRF setup. To make a fair comparison, we also optimized conditions and chose excitation power of 6.4 μ W for the 488 nm laser with 40 ms exposure time for Dronpa to get as high fluctuation as possible with acceptable photobleaching (Figure S9). Figures3a,b reveals the subtle differences among six consecutive images that arise from the fluctuations of the probes under the optimized imaging conditions. As seen in Figure 3c, in the fluctuation state, the fluorescence intensity of the green pixel in Figure 3a of Dronpa varied between 500 and 2000, whereas the fluorescence intensity in the red pixel of Skylan-S spanned a range of 8000–14000. Then we compared the averaged fluorescence intensity in the fluctuation states of Skylan-S and Dronpa, the red pixel with Skylan-S (11118.1) is much higher than that with Dronpa (1271.9). The higher fluctuation and averaged fluorescence intensity in the fluctuation state of Skylan-S produces a higher second-order cumulant value than that provided by Dronpa at each time point (Figures 3e, S10, and S11), where the second-order autocorrelation function $G_2(r,\tau)$ is given by^{8,9}

$$G_2(r,\tau) = \sum_k U^2(r-r_k)\varepsilon_k^2 \langle \delta s_k(t+\tau) \delta s_k(t) \rangle$$

where ε_k is the constant molecular brightness, which is related to the averaged fluorescence intensity in the fluctuation state for each pixel, s_k is a time-dependent fluctuation, $\langle \delta s_k(t + \tau) \delta s_k(t) \rangle$ denotes the time-averaged correlation value, and r_k and $U^2(r - r_k)$ represent the probe position and the PSF of the system, respectively. Notably, a higher averaged fluorescence intensity in the fluctuation state can yield a higher SNR and better resolution even with lower fluctuation (simulated data, Figure S11).

To quantitatively compare the fluctuation of the two probes, the standard deviation (SD) of the intensity observed from each pixel indicated by an asterisk in Figure 3a,b was derived from 10 s of image acquisition. As shown in Figure 3c,d, the fluctuation SD of Skylan-S was 1062.4, which was approximately 4-fold higher than that of Dronpa (SD = 241.1). These results indicate that Skylan-S provides an approximately 4-fold greater

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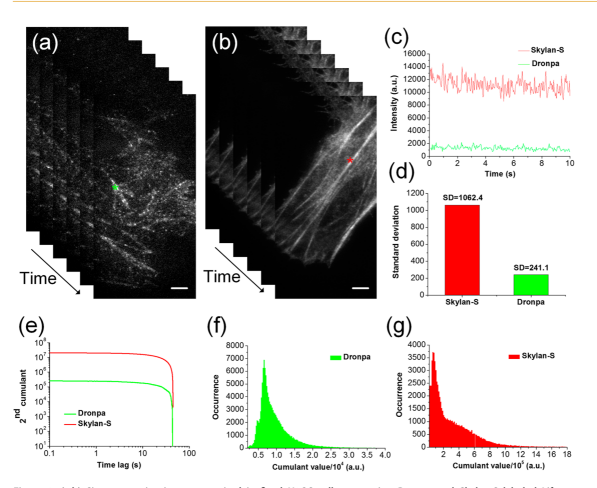


Figure 3. (a,b) Six consecutive images acquired in fixed U2OS cells expressing Dronpa and Skylan-S labeled Lifeact, respectively. The scale bar represents 5 μ m. (c) Intensity traces of Dronpa and Skylan-S for the two pixels indicated by green and red asterisks in panels a and b. (d) A histogram of the standard deviation of panel c. (e) The 2nd-order cumulants of the fluctuation statistics of Skylan-S and Dronpa. (f,g) Cumulant distributions (recorded with 32 bit grayscale values) of all imaged pixels in panels a and b. All cumulant values were calculated at the second order.

range of fluctuation and dramatically higher averaged fluorescence intensity in the fluctuation state with respect to its Dronpa counterpart (Figure 3c,d). Then, the cumulant curve (Figure 3e) and absolute second cumculant value distribution (recorded with 32 bit grayscale values) of each pixel were analyzed for both Dronpa and Skylan-S, and the results revealed that Skylan-S yielded a two-order of magnitude larger dynamic range of cumulant values than did Dronpa (Figure 3f,g), leading to richer grayscale and consequently much finer details in the image.¹⁸ Furthermore, although Dronpa can give nice reconstructed results shown in Figure S12a, Skylan-S can give much richer structural information suitable for both strong and weak fluorescent signals (Figure S12c,d).

To demonstrate the labeling capability of Skylan-S in living cells and its application in SOFI, the tubulin structure of living U2OS cells was observed using Skylan-S-labeled MAP4. Figure 4 clearly reveals the striking enhancement in resolution and signal-to-noise ratio achieved after second-order SOFI analysis and the even greater improvement achieved via the third-order cumulants. The resolution was improved approximately 2-fold after third-order SOFI analysis.

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Because of the very high averaged fluorescence intensity in the fluctuation state and the excellent photostability of Skylan-S, it was not necessary to perform post-piecewise analysis in blocks of frames, which is often required to compensate for the photobleaching of organic dyes or other fluorescent proteins.¹⁹

Next, we further tested the imaging capabilities of Skylan-S in SOFI using Skylan-S-labeled clathrin-coated pits (CCPs) in living U2OS cells. CCPs are important components of endocytic machinery and responsible for the endocytosis of many cell-surface receptors or their cargos; they are typically 150 to 200 nm in diameter. As expected, the SOFI images offered substantially improved resolution compared with conventional images. Under a conventional TIRF microscope, the Skylan-S-labeled CCPs exhibited a diffractionlimited cluster structure with no discernible morphological details (Figure 5a-c). By contrast, as shown in Figure 5d-f, second-order SOFI analysis revealed detailed structures by virtue of the resolution improvement. When third-order SOFI analysis was applied, even better resolution of below 100 nm could be achieved (Figure 5g-i). Doughnut-shaped structures of approximately 200 nm in diameter (with an inner-

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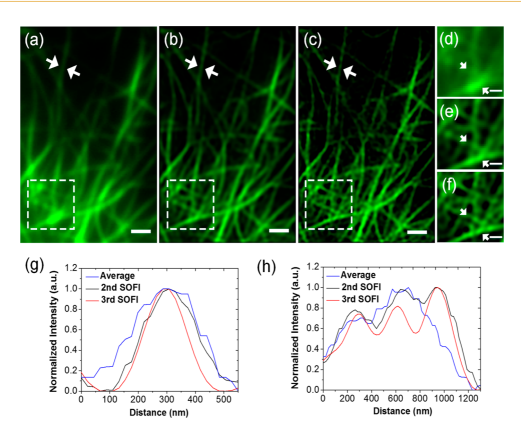


Figure 4. Fluorescence and corresponding SOFI images of living U2OS cells expressing Skylan-S fused with MAP4. (a) Average of original 800 fluorescence frames, and the exposure time was 30 ms per frame. (b) Second and (c) third-order SOFI images obtained through the analysis of the same 800 frames. (d–f) Zoomed-in views of the boxed regions in panels a–c, respectively. (g) Intensity profiles of cross sections taken along the two white arrowheads indicated in panels a–c. (h) Intensity profiles of cross sections taken along the two white arrowheads indicated in panels d–f. The scale bars represent 1 μ m (a–c) and 500 nm (d–f).

circle diameter of approximately 100 nm) could be clearly differentiated. Moreover, the size distribution of the CCPs that was obtained via third-order SOFI analysis was in quantitative agreement with the results obtained using electron microscopy (EM).²⁰ To further compare the capability of Dronpa and Skylan-S to resolve ultrastructure in long-time live-cell SOFI imaging, we labeled clathrin with these two fluorescent proteins. As shown in Figure S13, defined structure of CCPs could not be resolved by Dronpa-labeled CCP due to its extremely low signal-to-noise ratio (SNR) under ultrafast frame rate (333 frames/second). On the contrary, mature CCPs were resolved as a ring within 1.5 s using Skylan-S. More importantly, Dronpa showed apparent photobleaching among sequential time points. However, Skylan-S is stable enough during a 60 s time window (20,000 frames, 333 frames/second) to reconstruct 40 SOFI images (Figure S13). These results proved that Skylan-S could be used for long-time live-cell SOFI imaging at higher temporal resolution to resolve ultrastructure, whereas Dronpa is not suitable because of the low SNR and apparent photobleaching.

CONCLUSIONS

SOFI is a super-resolution imaging technique that uses the temporal correlations of the fluctuations of

imaged pixels to generate microscopic images with better resolution than is allowed by the diffraction limit. The fluctuations are produced by the flickering of fluorescent labels. RSFPs can be used for live-cell SOFI imaging, primarily because they can be easily fused with target proteins through genetic manipulation and can be photoactivated over many cycles. Compared with PALM/STORM imaging, SOFI offers several advantages in its suitability for low-SNR acquisition and its relatively lax requirements on labeling density.¹¹ In practice, the image quality provided by SOFI strongly relies on the averaged fluorescence intensity in the fluctuation state, photostability, and contrast ratio of the fluorophore. Although several RSFPs have been previously reported for application in SOFI, their tetrameric form is not suitable for the imaging of living cells.¹⁴ In this study, we developed a novel monomeric green RSFP, Skylan-S, which is optimized for SOFI and features very high photostability, contrast ratio, and averaged fluorescence intensity in the fluctuation state. As demonstrated in the previous paper by Peter Dedecker et al.,¹³ the fluorescence of Dronpa decreases greatly upon high power 488 nm laser illumination and reaches a plateau at very low level. Because of the rapid photobleaching, only the fluctuation of the low fluorescence at the plateau state

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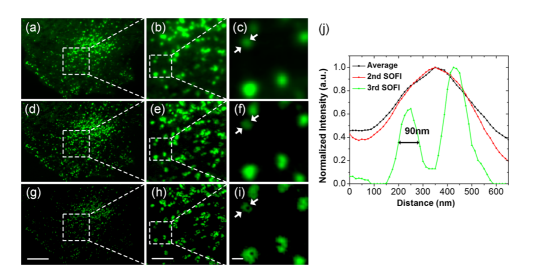


Figure 5. SOFI images of clathrin-coated pits (CCPs) in live U2OS cells. (a–c) TIRFM images of CCPs of living U2OS cells fused with Skylan-S under its optimal condition. (d–f) Second-order SOFI images. (g–i) Third-order SOFI images. (b,e,h) Zoomed-in views of the boxed regions in panels a, d, and g, respectively. (j) Intensity profiles of cross sections taken along the white arrows indicated in panels c, f, and i. The scale bars represent (a,d,g) 10 μ m, (b,e,h) 3 μ m, and (c,f,i) 500 nm.

can be used for SOFI reconstruction. In Table 1, the brightness was calculated as extinction coefficient \times quantum yield/1000 at the on-state of the proteins, corresponding to the peak intensity.¹³ So there is no very big difference in brightness (fluorescence intensity) between Skylan-S and Dronpa when they are both excited to on-states (the increase in brightness is 15% in Table 1). However, for SOFI imaging, the properties of fluorescent proteins at the plateau (averaged fluorescence intensity and fluctuation) are extremely important for SOFI reconstruction, which is greatly different between Skylan-S and Dronpa. At the plateau, there are continuous activation and deactivation of individual molecules, which is an equilibration of the light-induced on and off switching. We attribute dynamic changes of the number of activated molecules (or deactivated molecules), as well as contrast ratio of the on/off-states, to the dynamic range of fluctuation. We speculate that more individual molecules of Skylan-S compared with Dronpa are involved in the equilibration of lightinduced on and off switching, and higher contrast ratio of Skylan-S than Dronpa is helpful to generate a wider dynamic range of fluctuation. We also attribute the photostability to switching cycles. Skylan-S is more immune to photobleaching than Dronpa, resulting in more switching cycles that make it more suitable for long-time SOFI imaging. Using Skylan-S, a 4-fold improvement in the fluctuation range of imaged pixels can be obtained (Figure 3d). The morphological structures of tubulin and clathrin-coated pits (CCPs) in U2OS live cells could be easily obtained via the SOFI technique using Skylan-S. Moreover, in addition to large fluctuation, Skylan-S also demonstrates attractive singlemolecule statistics under high power 488 nm irradiation. We have performed PALM localization microscopy with Skylan-S, as shown in Figure S14. High power excitation can drive the Skylan-S molecules to switch between onand off-states, enabling the single-molecule localization. Comparing with the wide-field counterparts, PALM of Skylan-S can yield much higher resolution (\sim 20 nm) and consequently better visualization of the fine details of the actin structure. In the near future, RSFPs that can produce rapid fluctuation but stable average fluorescence signal will be required for further improving the temporal resolution of SOFI,²¹ and it will be necessary to develop a stable red RSFP for dual-color SOFI imaging.²² In conjugation with other techniques such as JT-SOFI and spinning disk confocal, high spatiotemporal super-resolution for 3D live cell imaging can also be anticipated.23,24

METHODS.

Protein Expression and Purification. The mEos2, mEos3.1, Dronpa, and Skylan-S proteins were expressed in the *E. coli* strain BL21 (DE3) and purified using a Ni-NTA His-Bind resin (Qiagen) followed by a gel filtration step using the Superdex 200 column (GE Healthcare). For further analysis, the purified proteins were concentrated via ultrafiltration and diluted in phosphate buffered saline (PBS).

Measurement of Spectral Properties. The proteins were first diluted in PBS (pH 7.4) to limit their UV absorption at a concentration of 0.1 μ g/ μ L to allow their quantum yields to

be accurately measured. Then, the absorption and excitation/ emission spectra were immediately recorded using an Agilent 8453 UV/V spectrophotometer and an Edinburgh Instruments FLS920 spectrometer, respectively. For the acquisition of emission spectra, Skylan-S was irradiated with 503 nm light. To determine the fluorescence excitation spectra, the fluorescence was recorded at 513 nm. The fluorescence quantum yields and the molar extinction coefficients at the absorption maxima were determined relative to the reported value for mEos2.

 $\begin{array}{c} \textbf{Cell Culture, Transfection, and Fixation.} \\ \textbf{U2OS cells were cultured} \\ \textbf{in McCoy's 5A Medium Modified (Gibco) supplemented with} \end{array}$

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10% fetal bovine serum and were maintained at 37 °C and 5% CO_2 in a humidified incubator. The cells were then transiently transfected using Lipofectamine 2000 (Invitrogen) following the manufacturer's protocol. Five hours after transfection, either the cells were trypsinized and plated at lower density on clean coverslips (Fisher Scientific) to induce spreading for another 24 h in complete medium without phenol red for live-cell imaging or the cells were fixed with 4% (w/v) *para*-formalde-hyde and 0.2% glutaraldehyde in PBS for 15 min at 37 °C, washed three times with PBS, and stored in PBS until fixed-cell imaging.

Analysis of the Switching Kinetics of Skylan-S. To analyze the switching kinetics of Skylan-S in E. coli, we used a homemade TIRF microscope to acquire images. Skylan-S proteins were expressed in the E. coli strain BL21 (DE3), and the E. coli cells were cultured in LB liquid medium at 37 °C for amplification. Then, the medium was changed to PBS (pH = 7.4). To keep the E. coli cells immobilized, cleaned glass coverslips were sequentially coated with 10 µL of Skylan-S-expressing E. coli in PBS, 10 μ L of 10% acrylamide, 1 μ L of TEMED, and 1 μ L of 10% AP and were then covered with another glass coverslip for 30 min at room temperature. E. coli cells expressing Dronpa and Skylan-S were continuously excited with 488 nm light (0.24 mW), and pulsed 0.01 s bursts of illumination with a 405 nm laser at a power of 0.2 mW were used to photoactivate the fluorescent proteins. The fluorescence was recorded continuously for 21 cycles. We repeated on/off switching until we observed considerable photobleaching of Dronpa. Throughout all experiments, the scanning speed, gain, amplification, and offset value were held constant. The ROI value was identified and read out using ImageJ software (NIH) and analyzed using OriginPro 8.

Measurement of Contrast Ratio. The contrast ratio was the mean value measured based on n cycles (n > 3) in Figure 1c,d. For each cycle, the ratio was calculated as the maximum fluorescence value (switching on) divided by the minimum fluorescence value (switching off).

Relaxation Half-Time Measurement. The protocol was used as described before.¹⁵ Purified protein of Skylan-S was first photoswitched to its off-states using a 60 mW 488 nm LED. Then, the relaxation curves from off-state into thermal equilibrium state were obtained using a Varioskan Flash spectral scanning multimode reader (Thermo Scientific) with 96-well optical bottom plates (Nunc) for approximately 20 h at 25 °C. A weak blue light (490 nm) was used to excite the sample, and the fluorescence of the sample at 513 nm was recorded every 4 min during the period. Then, the determination of relaxation half-time for Skylan-S was based on single exponential function fitting of the curve.

Absorption Spectra of Skylan-S in On/Off-States. The purified Skylan-S protein was diluted in PBS (pH = 7.4) and an appropriate power of 488 nm LED was used to partially photoswitch the protein to the off-state. The absorption spectra were immediately recorded using an Agilent 8453 UV/V spectrophotometer in the on-state and off-state, respectively.

Time-Resolved Fluorescence Spectroscopy. Time-resolved fluorescence spectroscopy was performed by using the time-correlated single photon counting method.²⁶ Fluorescence from Skylan-S molecules (pH 7.4 in PBS) was registered using an Edinburgh Instruments FLS980 spectrometer at room temperature. The excitation and emission wavelength were set to be 473.4 and 513 nm, respectively.

Analysis of Oligomerization. For oligomerization analysis, sizeexclusion chromatography was first performed using a Superdex 200 column and an Akta purifier system (GE Healthcare) to confirm the oligomeric states of Skylan-S. Before chromatography, the proteins were concentrated to 3–4 mg/mL. The flow rate was set to 0.5 mL/min. Protein absorption was monitored at 280 nm. All measurements were performed at 16 °C. Then, the protein was subjected to analytical ultracentrifuge assays. Sedimentation equilibrium experiments were performed using a Beckman Optima XL-1 analytical ultracentrifuge at 20 °C. Purified proteins at 8 to 20 μ M were loaded into 6-channel centrifugation cells and normalized to the corresponding dialysis buffer (PBS, pH 7.4). The samples were sequentially centrifuged at 10000, 15000, and 20000 rpm. The data were analyzed via nonlinear least-squares analysis using the Microcal Origin software package supplied by Beckman. The solvent density, partial specific volume, and calculated molecular weight used in the analysis were determined using SEDNTERP v. 1.01. To perform sedimentation velocity analytical ultracentrifugation, we purified proteins at concentrations ranging from 280 to 320 μ g and then loaded them into centrifugation cells and normalized them to the corresponding dialysis buffer (PBS, pH = 7.4). The theoretical molecular weight of the monomer Skylan-S with the $6 \times$ His tag was approximately 23.8 kDa.

Imaging System and Acquisition. SOFI imaging of Skylan-S was performed as previously described.^{15,16} A commercial optical microscopy system (Olympus IX-71; Olympus, Japan) with two high-NA oil objectives (100×, NA 1.49 and 150×, 1.45 NA Olympus PLAN APO with internal $1.6 \times$ magnification lenses in the microscope system) were used for imaging acquisition. A 405 nm laser and a 488 nm laser (Coherent, Santa Clara, CA, USA) were used to excite the Skylan-S with which the target sample was labeled. A long-pass filter (500-550 nm) was placed before an electron-multiplying charge-coupled device (EMCCD) camera (Andor iXon DV-897 BV). The maximum power near the back pupil of the objective was 8 mW for the 405 nm laser (LASOS) and 2.5 mW for the 488 nm laser (Cobolt Dual Calpos). For longtime live-cell SOFI imaging, clathrin were labeled with Skylan-S and Dronpa, and sequential frames of Dronpa-CCP and Skylan-S-CCP were acquired using TIRF microscope (150 \times , 1.45 NA objective and internal 1.6× magnification lenses in the microscope system) under the same experimental conditions, except for optimized 488 laser powers for each protein to get high fluctuation. Each sequential 500 frames were used to reconstruct a SOFI image at different time points (Figure S13).

Data Processing. To correct for drift during imaging, we first analyzed the raw data using a subpixel drift-correction algorithm based on discrete Fourier transforms and nonlinear optimization implemented in self-reorganizing code in MATLAB 2013a (Mathworks Inc., USA). Then, SOFI processing was implemented by calculating the spatiotemporal cross-cumulants between adjacent pixels. After balancing, the nonlinear brightness response in SOFI images can be linearized and the spatial resolution can be further improved linearly with the cumulant order. In this Article, second-, third-, or fourth-order cross-cumulant balanced SOFI (bSOFI) was applied to 500–1000 frames using the shortest accessible lag time, as shown in Figures 3–5.^{8,9,11,27,28}

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

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Supporting Information Available: Supplementary figures and figure legends. This material is available free of charge via the Internet at http://pubs.acs.org.

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