Simple super-resolution live-cell imaging based on diffusion-assisted Förster resonance energy transfer

Sangyeon Cho1*, Jaeduck Jang2*, Chaeyeon Song2,3, Heeyoung Lee4, Prabhakar Ganesan1,5, Tae-Young Yoon5, Mahn Won Kim2, Myung Chul Choi3, Hyotcherl Ihee1,5, Won Do Heo4 & YongKeun Park2

1Department of Chemistry, 2Department of Physics, 3Department of Bio and Brain Engineering, 4Department of Biological Sciences, Korea Advanced Institute of Science and Technology, Daejeon 305-701, Republic of Korea, 5Center for Nanomaterials and Chemical Reactions, Institute for Basic Science, Daejeon 305-701, Republic of Korea.

Despite the recent development of several super-resolution fluorescence microscopic techniques, there are still few techniques that can be readily employed in conventional imaging systems. We present a very simple, rapid, general and cost-efficient super-resolution imaging method, which can be directly employed in a simple fluorescent imaging system with general fluorophores. Based on diffusion-assisted Förster resonance energy transfer (FRET), fluorescent donor molecules that label specific target structures can be stochastically quenched by diffusing acceptor molecules, thereby temporally separating otherwise spatially overlapped fluorescence signals and allowing super-resolution imaging. The proposed method provides two- to three-fold-enhancement in spatial resolution, a significant optical sectioning property, and favorable temporal resolution in live-cell imaging. We demonstrate super-resolution live-cell dynamic imaging using general fluorophores in a standard epi-fluorescence microscope with light-emitting diode (LED) illumination. Due to the simplicity of this approach, we expect that the proposed method will prove an attractive option for super-resolution imaging.

Fluorescence imaging of live cells plays a crucial role in the study of biological processes at the cellular and subcellular levels. The diffraction property of light, however, limits the spatial resolution of conventional fluorescence microscopy to ~250 nm and ~600 nm in the lateral and axial directions, respectively (Fig. 1a). A series of super-resolution microscopy techniques have been developed to overcome the diffraction limit1,2, including techniques based on stimulated emission depletion or patterned illumination in order to confine the fluorescence to a sub-diffraction-sized area or volume [such as stimulated emission depletion (STED) microscopy3,4, ground-state depletion (GSD)5, reversible optically linear fluorescence transitions (RESOLFT)6, and saturated structured-illumination microscopy (SSIM)7] or techniques that are based on the repeated on/off switching of fluorescent probes with single-molecule localization [such as photo-activated localization microscopy (PALM)8,9, and stochastic optical reconstruction microscopy (STORM)10,11]. Recently, a breakthrough in super-resolution imaging techniques based on single-molecule localization has been achieved by using standard fluorescent molecules or dyes12,13: direct STORM (dSTORM)14, fluorescence-PALM15, bleaching/blinking-assisted localization microscopy (BaLM), ground-state depletion imaging (GSDIM)16, Bayesian analysis of blinking and bleaching (3B analysis)17, and imaging membrane structures with lipophilic cyanine dyes18.

Although recent super-resolution microscopy techniques can achieve spatial resolutions up to tens of nanometers, these methods still have practical limitations to their direct employment in general biological studies. Most of the existing super-resolution imaging techniques require special imaging systems illuminated with appropriate powers and wavelengths. STED requires high power illumination in order to generate a sharpened excitation spot much smaller than the diffraction-limited focus. The RESOLFT and SSIM methods also generally require the complex alignment of specialized optical system. On the other hand, methods based on single-molecule localization, such as STORM and PALM, are temporally demanding. They generally require photo-switchable probes to activate and excite the probes with a sufficient number of photons for highly accurate localization. They also require a large number of measurements proportional to the number of fluorophores. In addition, they require high signal-to-noise ratios. This prevents the use of conventional epi-fluorescence
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microscopy but limits the total internal reflection (TIRF) excitation geometry. In addition, labeling intracellular structures with efficient reversible photo-switching fluorescence still remains challenging.

Recently, a statistical analysis method, super-resolution optical fluctuation imaging (SOFI)\(^\text{19}\), was introduced. Without requiring specialized equipment, SOFI utilizes a correlation of temporal fluctuations between neighboring pixels, which can provide a super-resolution image with a high signal-to-noise ratio\(^\text{20}\) by calculating the cumulant\(^\text{19,21}\). The enhancement of spatial resolution depends on the order of analysis; calculating the \(n\text{th}\) order cumulant provides a factor of \(n\text{th}\) improvement\(^\text{19}\). SOFI has shown its potential to be employed in general optical imaging systems as long as fluorescent signals are randomly fluctuating in time. However, until very recently, SOFI had been only applicable for samples with immuno-labeled quantum dot and organic dyes that have intrinsic blinking characteristics\(^\text{19,22,23}\). Recently, photochromic stochastic optical fluctuation imaging (pcSOFI) has been introduced, which utilizes single-molecule fluctuation using reversibly photochromic labels\(^\text{24}\). Illuminating a reversibly photo-switching fluorescent protein at specific wavelengths, genetically encodable labels can be used for SOFI. However, pcSOFI still face limitations since it requires special reversibly photochromic labels as well as lasers with appropriate wavelengths; these complications still hinder direct applications by many potential users.

Here, we report a simple but powerful technique for super-resolution fluorescence imaging with diffusion-assisted Förster resonance energy transfer (FRET). Fluorescent donor molecules that label target structures can be stochastically quenched in the presence of diffusing acceptor molecules, resulting in the temporal separation of otherwise spatially overlapped fluorescence signals and allowing super-resolution imaging. The proposed technique does not rely on either photo-bleaching events of fluorophores or complex image analysis; thus, it can be readily employed in existing imaging systems.

Since our approach uses general fluorophores including conventional dyes and typical fluorescent proteins in a conventional epi-fluorescence microscope with general illumination – even with light-emitting device (LED) illumination, we refer to it as “direct” SOFI, dSOFI. Our method is based on SOFI reconstruction, and key advantages of SOFI have been shared, namely, its technical simplicity, a broad range of imaging conditions, and a simple imaging process only requiring consecutive images and applying SOFI analysis. The dSOFI method using general fluorophores in a standard epi-fluorescence microscope improves spatial resolution by a factor of two with a markedly improved signal-to-noise ratio and improved optical sectioning capability. We demonstrate the potential of dSOFI under a broad range of imaging conditions, including labeling antibodies, chemically tagged fluorescent dyes, and genetically encoded proteins in dynamic live-cell imaging. We present the sub-diffraction resolution fluorescence imaging of \emph{in vitro} microtubules, actin, and intermediate filaments of live NIH 3T3 cells, as well as their dynamics.

Results

\textbf{dSOFI principle.} The dSOFI method uses general (instead of intrinsically blinking) fluorophores in a conventional epi-fluorescence microscope system. FRET is energy transfer from a donor to an acceptor molecule, with the FRET efficiency depending on the sixth power of the distance between the two\(^\text{25}\). Usually, \(R_0\) (the distance at which the FRET efficiency is 50\%) is approximately 5–10 nm. FRET has been widely used to study single molecular interaction by precise measurement of the distance between two molecules\(^\text{26}\). In dSOFI, we exploit FRET and the diffusion process to induce the stochastic blinking of general fluorophores, the combination of which results in super-resolution fluorescence imaging. Generally, fluorophores do not exhibit significant blinking fluorescence, but the addition of freely diffusive acceptor molecules, which appropriately form a FRET pair with fluorophores, makes the target labeling fluorophore molecules repeatedly blink with fluorescence. Like conventional fluorescence imaging, fluorophores as donor molecules are labeled on the target structures.

In dSOFI, FRET acceptor molecules are introduced and they diffuse freely. FRET occurs only when a diffusive acceptor molecule approaches a donor molecule closely enough and the donor fluorophore transiently stops exhibiting fluorescence. After the acceptor molecule move away, the donor fluorophore resumes exhibiting fluorescence. Thus, this short range (5–10 nm) and random temporal FRET causes uncorrelated fluorescence fluctuation on nearby donor fluorophores. Through the analysis of this uncorrelated fluorescence fluctuation signal, a super-resolution image of target molecules can be achieved (Fig. 1b).

\textbf{In vitro microtubule imaging.} To demonstrate the principle of dSOFI and compare its performance with that of conventional microscopy, we performed \emph{in vitro} measurement of microtubules stained with fluorescent amino-methyl-coumarin-acetate (AMCA) dye, which had 350/450 nm excitation/emission wavelengths.
Photoactive yellow protein (PYP) was used as an acceptor molecule to form a FRET pair with AMCA. A series of fluorescence microscopic images were measured with a conventional epi-fluorescence microscope (Fig. 1c) and then analyzed with the SOFI algorithm. The reconstructed dSOFI image is shown in Fig. 2a. The dSOFI method improved the resolution two-fold compared with that of a conventional wide-field fluorescent image. After applying a deconvolution algorithm to the DSOFI image, this method achieved a spatial resolution down to a full-width half-maximum (FWHM) of 32.5 nm (Fig. 2b), which is an almost eight-fold improvement compared to the performance of diffraction-limited conventional fluorescence microscopy. The dSOFI images obtained in this study clearly reveal individual microtubule structures, which are not resolved in the conventional wide-field fluorescence image.

Stochastic quenching by diffusing acceptor molecules. To demonstrate that the resolution enhancement in dSOFI primarily results from FRET-induced fluorescence blinking and not from the photo-bleaching effect, we analyzed temporal fluorescence signals from sequential raw fluorescence images. If fluorescence molecules labeled at target structures blink enough to localize the target structures within the diffraction spot size, the temporal fluorescence signals from the structure must be differentiable from signals from nearby points, which can be quantified by calculating correlation or covariance. We analyzed the fluorescence signals of a single microtubule fiber labeled with AMCA (Fig. 3a) in the absence and (Fig. 3b) in the presence of corresponding FRET molecule PYPs, respectively. Both cases show comparable spatial resolution since no SIFT analysis was performed in Figure 3b. Then, we calculate the cross-correlation between various spatial positions:

\[
\langle I \ast I \rangle (\Delta r) = \int I(r, t) I(r + \Delta r, t + \tau) \, d\tau.
\]

The normalized cross temporal correlations between pixels on the white dashed lines denoted in Figs. 3a–b are shown in Figs. 3c–d, respectively. The widths of cross-correlation normal to the main diagonal line show how much temporal fluorescence signals are correlated; only uncorrelated temporal signals can be distinguished by SOFI reconstruction. Thus, the widths of cross-correlation normal to the main diagonal line correspond to the spatial resolving power.

In the absence of appropriate FRET acceptor pairs, the target molecules did not exhibit significant blinking fluorescence, and fluorescence signals on the microtubule structure showed strong spatial correlation between nearby pixels along off-diagonal directions with the diffraction spot size (250 nm ~ 5 pixels, Fig. 3c). This indicates a strong correlation between pixels within a diffraction spot size; thus, there is no significant resolution enhancement. However, in the presence of FRET acceptor molecules, fluorescence signals on the microtubule structure presented little spatial correlation,
indicating that blinking caused by stochastic FRET events can improve spatial resolving power when combined with SOFI analysis (Fig. 3d).

**Acceptor concentration on dSOFI analysis.** To study the effects of acceptor concentration on dSOFI analysis, we performed *in vitro* microtubule imaging experiments with various concentrations of PYP. At a fixed donor concentration of fluorescent amino-methyl-coumarin-acetate (AMCA) dye which stains microtubules (See Methods), we performed experiments with a diverse range of PYP concentrations (0 μM, 0.5 μM, 2.5 μM, 5 μM, 7.5 μM, 10 μM, 12.5 μM, 15 μM, 25 μM, and 45 μM). The results show that there is an optimal range of acceptor concentration (7.5–12.5 μM) in which dSOFI analysis effectively works with good repeatability. Too high or too low acceptor concentration easily frustrates the blinking of donor molecules to the extent that SOFI reconstruction does not work effectively. Under a low concentration regime, most donor molecules will emit fluorescence; however, a few FRET events will occur at single-molecule levels. On the other hand, with a high concentration of acceptor molecules, most donor molecules will exhibit ensemble FRET events; thus, induced blinking is not sufficient for dSOFI analysis. At an optimal concentration of acceptor molecules, diffusion-assisted FRET will cause appropriate small ensemble of blinking, which can be used for SOFI reconstruction.

To further demonstrate the principle of dSOFI with control groups, we measured polystyrene beads with a diameter of 10 μm coated with AMCA dye (See Methods) under various imaging conditions. First, conventional wide-field images of the beads show diffraction-limited spatial resolution irrespective of the presence or absence of acceptor molecules, PYP (Fig. 4a). When SOFI reconstruction is applied to the images of the beads in the presence of PYP, it produces super-resolution images of the beads. However, in the absence of PYP, SOFI reconstruction does noting, and it still gives diffraction-limited images.

**Optical sectioning in dSOFI.** To evaluate the optical sectioning capability of dSOFI, we measured the depth-of-field by imaging polystyrene beads with a diameter of 10 μm coated with AMCA dye. The wide-field fluorescence images of the beads with and without PYP protein as an acceptor molecule are diffraction-limited with large oscillatory patterns caused by optical diffraction (Fig. 4a). When the dSOFI method was applied, the edge of the bead was visualized with a very shape profile. This result demonstrates the optical sectioning capability of dSOFI as well as enhanced spatial resolution it achieves. To quantitatively analyze the depth-of-field, the intensity profiles measured by wide-field imaging and dSOFI are shown as functions of the lateral position (Fig. 4b). In the wide-field image, strong intensity signals were seen inside the radius of the bead; fluorescent dyes at defocus planes were seen due to long depth-of-field. However, the dSOFI image shows a very sharp profile at the edge, which demonstrates that the depth-of-field is significantly reduced. The axial positions of the bead corresponding to the position with 50% fluorescent intensity are 2.2 μm and 1 μm for the wide-field image and dSOFI image, respectively.

**Super-resolution live-cell imaging of F-actin and α-tubulins.** To demonstrate the capability of dSOFI in super-resolution live-cell imaging, we visualized two major cytoskeleton structures in live cells (see Methods section). Filamentous actin (F-actin) and α-tubulins in NIH 3T3 cells were visualized with Lifeact (F-actin marker) and α-tubulin labeled with cerulean [a cyan fluorescence protein (CFP) variant]. Citrine [a yellow fluorescence protein (YFP) variant] is expressed in the cytoplasm as a FRET pair to cerulean.

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Note that acceptor molecules can also be introduced into cell cytoplasm via various alternative routes, including micro-injection, electroporation, endocytosis, and so on. However, here we employed a double-expression method to achieve the physiologically relevant concentration of both acceptor and donor molecules. Both the cerulean and citrine were genetically expressed *in vivo*, and super-resolution imaging has been effectively demonstrated, suggesting that dSOFI still works for the physiological conditions of live cells.

More importantly, dSOFI allows us to visualize the dynamics of living cells over a long period and at extremely high spatial resolving power, since dSOFI techniques do not require the fixation of biological samples. F-actin structures labeled with cerulean in a live NIH 3T3 cell were imaged with dSOFI every 5 min during a period of 15 min (Fig. 6). The dynamics of actin polymerization and cytoskeleton movements were captured in extremely specific detail (green and blue arrows in Fig. 6). Since the analysis of blinking fluorescence signal with the SOFI algorithm provides optical sectioning capability as well as enhanced resolving power, the dynamics of F-actin fluctuation in the axial direction can be observed.

**Discussion**

In conclusion, we present a novel super-resolution microscopic technique, dSOFI, which is easily transferable to conventional fluorescent microscopy imaging systems. The dSOFI method represents the first
attempt to achieve super-resolution imaging in live cells using 
FRET\textsuperscript{28}; dSOFI exploits diffusion-assisted FRET to achieve stochas-
tically fluctuating fluorescence from standard fluorophores – fluo-
rescent proteins and dyes. This new technique achieves dynamical 
super-resolution imaging in live cells. dSOFI is a very simple, general 
and cost-efficient super-resolution imaging method, which can be 
used directly in existing fluorescent imaging systems with general 
fluorophores. Using this method, we demonstrated super-resolution 
live-cell imaging with standard non-photoswitchable fluorophores 
under a conventional epi-fluorescent microscope equipped with 
LED illumination. Simply by adding diffusive acceptor molecules 
near target structures, anyone can produce super-resolution molecu-
lar imaging with existing instrumentation.

In addition, the dSOFI technique provides extremely fast imaging 
acquisition and analysis, especially compared to techniques based 
on the localization of individual fluorescent molecules\textsuperscript{10,29}. Only 300 raw 
fluorescent image measurements, typically captured at video frame 
rate of 10–50 Hz, are required to reconstruct one super-resolution 
dSOFI image as shown in Fig. 1d. Considering the typical diffusion 
properties of fluorescent molecules, comparable resolution improve-
ment is expected to be achieved up to an acquisition speed of 0.1 ms 
using the proposed method. Depending on the acquisition speed of 
the camera, the fast dynamics of biological systems can be visualized 
at super-resolution. In addition, to analyze a region of around 25 × 
25 μm, processing takes ~3.5 s on a typical desktop computer (Intel 
Core i5-2430M CPU, 2.40 GHz).

As demonstrated here for a fluorescent dye – protein pair and a 
fluorescent protein – protein pair, the principle of dSOFI should work 
for any FRET pairing molecules. Conventional fluorescent micro-
scope imaging systems can easily adopt this system. The simple 
addition of an acceptor that can form a FRET pair with the target 
labeling fluorophore, and the analysis of the blinking fluorescence

**Figure 5** | Live-cell dynamic imaging using dSOFI. (a) Actin-ceruleine expressed in live NIH 3T3 cell imaged with conventional wide-field fluorescence microscopy (left) and dSOFI (middle). Cytoplasmic citrine acts as a FRET acceptor. (i, ii) Magnifications of the white box regions. (b) Tubulin-
cerulean expressed in live NIH 3T3 cell imaged with conventional wide-field fluorescent microscopy (left) and dSOFI (middle). (iii, iv) Magnifications of 
the white box regions.

**Figure 6** | Long-time duration dynamic images of actin-ceruleine expressed live NIH 3T3 cells. (a) imaged with wide-field fluorescence microscopy. 
(b) imaged with dSOFI. Magnified images of the box regions measured every 5 min up to 15 min.
signal can produce a super-resolution image. No special photo-activatable molecules or complex optical systems are required. As demonstrated in this study, even LED illumination can be used since the proposed technique does not require high-intensity illumination. However, the proposed method also bears some practical limitations. For example, the use of a FRET pair to obtain a single-channel image may complicate sample preparation and make multi-color imaging difficult. Nonetheless, the proposed approach still has appealing advantages, namely, the simplicity of a method that only requires a FRET pair to induce the blinking of labeling molecules as well as the fact that potential users can utilize existing imaging systems.

In summary, we demonstrated the viability of this principle through the imaging of cytoskeleton structures in vitro and as expressed in live cells. Our new technique is sufficiently broad and general to allow for potential application in cell biology, biophysics, biochemistry, biotechnology, and biomedicine, wherever super-resolution molecular imaging can have an impact. All users of microscopes can use their own super-resolution fluorescence microscopes to apply the present technique. Thus, dSOFI will immediately find direct and versatile applications in diverse fields such as cell biology, biophysics, and biochemistry; therefore, we believe that our new method represents a vast improvement on existing super-resolution imaging technologies, especially in live-cell studies.

Methods
Microtubule preparation. AMCA-labeled microtubules were assembled at 35°C for 20 min from 0.91 μM (0.1 mg/ml) tubulin (from Cytoskeleton, cat. # TL440M) in PEM50 buffer (50 mM Pipes, 1 mM EGTA, 1 mM MgCl2, pH 6.8) in the presence of 1 mM GTP and 5 wt% glycerol and then stabilized by 0.91 μM taxol (2 mM taxol in DMSO was added in a stepwise manner)53. Purified PYP was synthesized based on Ref. 31. PYP was concentrated to 5 μM with PEM50 buffer. We mixed the microtubule solution and 5 μM PYP solution in 1:1 volume ratio.

Fluorescence bead preparation. 50 μl of aliphatic amine latex beads with a diameter of 4.5 μm (Invitrogen, Molecular probes, Inc. A37370) from the stock solution (2 × 10^10 beads/ml) was diluted to a volume of 1 ml in distilled water. The bead solution was centrifuged at 3600 rpm for 15 minutes and re-suspended in a 90 μl PBS (pH 7.4). 50 μg of Alexa350 (Invitrogen, Alexa Fluor® 350 NHS esters, A10168) NHS esters and 10 μl of sodium bicarbonate are then added to the bead solution and incubated at room temperature in a tube rotator for 30 minutes. Then, Alexa350/bead solution was subjected to three centrifugations (3600 rpm for 15 minutes) and re-suspension steps (PBS, pH 7.4) to remove any free Alexa350 NHS esters.

Cell preparation: cell culture and transfection. NIH 3T3 cells were purchased from American Type Culture Collection (ATCC). Cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM, PAA) supplemented with 10% fetal bovine serum (FBS, Gibco) and 1% antibiotics – anti mycotics. NIH 3T3 cells were maintained in 10% CO2 at 37°C. NIH 3T3 cells were plated on glass bottom 96-well plates (Metworks Inc.) and plated on poly-D-lysine-coated glass bottom 96-well plates. Transfection was performed after one day after plating, by using lipofectamine 2000 (Invitrogen) with 500 ng of DNA according to manufacturer’s protocol. Transfected cells were incubated for 24 hrs before imaging.

Cell preparation: DNA constructs. Cerulean-Tubulin constructs were made by subcloning tubulin, Human tubulin alpha 1B, from the pAxGFP-Tubulin (Clontech) plasmids into Cerulean-C1 using Xhol and BamH1 restriction enzymes. Lefactory fragment was generated by annealing method with following primers: sense 5'-AATTCTGGTGTCGCAGATTTGATCAAGAAATTCGAAAGCATCTCAAAGG-AAGAAG-3' and antisense 5'-GACTTCCTCTTTCTGAGTCTGTTCGAA-TTCTGTGACAAATCGGACACCA-3'. Lefactory fragment was cloned into Cerulean-C1 using EcoRI and BamH1 restriction enzymes. Citrine-C1 plasmids that were used in this study were previously described45.

Imaging system and acquisition. A conventional optical microscopy system (IX-71, Olympus) equipped with an objective lens (US1200×, Olympus) was used for SIFT. An UV LED (M38LS2-C1, Thorlabs, 270–430 mW) or a white-light LED (MCWHL2-C1, Thorlabs, 500–700 mW) was used to replace the donor molecules that labeled the target samples. For the AMCA-PYP FRET pair, the UV LED was used to excite a dichroic mirror (MD416, Thorlabs) and an emission filter (FB460-10, Thorlabs) mounted in a fluorescence filter cube (IX-RAF, Olympus) were used to acquire fluorescence image. For the Cerulean-C1 FRET pair, the white-light LED (MCWHL2-C1, Thorlabs) source was used for illumination. An excitation filter (FF01-438/24, Semrock), an emission filter (FF01-483/32, Semrock), and a dichroic mirror (FF458-Di01, Semrock) were used to acquire fluorescence image. The illumination powers at the sample plane for AMCA and CFP are 9 W/cm^2 and 1.3 W/cm^2, respectively. For an imaging system, we employed a high magnification objective lens (US12100×, Olympus) with 1.30 NA and a sCMOS camera (NEO, Andor) with a 6.5 μm pixel size. The camera was directly adapted to a ×1 magnification port of IX-71, and one pixel size of the detector corresponds to 65 nm at the sample plane. We typically acquired 300–1000 consecutive images with 10–50 frames per second.

Data processing. To retrieve a super resolution image from the measured fluorescent images, we analyze the data with a SOFI algorithm using a custom-made code in Matlab (Mathworks Inc., Natick, MA). Before analysis, raw images were digitally magnified by factor of 2. We calculated the 2nd or 4th-order auto-cumulant SOFI (2nd or 4th AC-SOFI) by using the shortest accessible lag time (the frame integration time) and then deconvolved the image by using a two-dimensional Gaussian function, whose full-width half-maximum is a size of diffraction limit of the used optical imaging system, employing deconvlucy (Lucy-Richardson) command44.
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Author contributions
S.C. and J.J. designed experiments, performed research, analyzed the data and wrote the paper; C.S., H.L., P.G., T.-Y.Y., M.W.K. and M.C.C. performed research and analyzed the data; H.I., W.D.H. and Y.P. designed experiments, analyzed the data, and wrote the paper.

Additional information
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