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Visualizing Dynamics of Cell Signaling *In Vivo* with a Phase Separation-Based Kinase Reporter

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SUMMARY

Visualizing dynamics of kinase activity in living animals is essential for mechanistic understanding of cell and developmental biology. We describe GFP-based kinase reporters that phase-separate upon kinase activation via multivalent protein-protein interactions, forming intensively fluorescent droplets. Called SPARK (separation of phases-based activity reporter of kinase), these reporters have large dynamic range (fluorescence change), high brightness, fast kinetics, and are reversible. The SPARK-based protein kinase A (PKA) reporter reveals oscillatory dynamics of PKA activities upon G protein-coupled receptor activation. The SPARK-based extracellular signal-regulated kinase (ERK) reporter unveils transient dynamics of ERK activity during tracheal metamorphosis in live Drosophila. Because of intensive brightness and simple signal pattern, SPARKs allow easy examination of kinase signaling in living animals in a qualitative way. The modular design of SPARK will facilitate development of reporters of other kinases.

In Brief

A separation of phases-based activity reporter of kinase, dubbed SPARK, is designed via multivalent protein-protein interactions utilizing homo-oligomeric coiled coils. The intensive

AUTHOR CONTRIBUTIONS

DECLARATION OF INTERESTS

The authors declare no competing interests.

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SUPPLEMENTAL INFORMATION

Supplemental Information includes 10 figures and 11 movies and can be found with this article online at https://doi.org/10.1016/j.molcel.2017.12.008.

X.S. conceived the project. X.S. designed the reporters. Q.Z. made the constructs, conducted fluorescence imaging, and characterized the reporters. T.B.K. and H.H. designed the *Drosophila* experiments. S.R.C., R.W., and A.S. designed the zebrafish experiments. Q.Z., H.H., L.Z., R.W., C.I.C., J.T., S.-Q.Z., and A.S. performed the experiments. X.S., Q.Z., and H.H. wrote the manuscript. All authors contributed to the final draft.



brightness and simple signal pattern of SPARK allow robust and direct visualization of kinase signaling in living animals.

INTRODUCTION

Temporal dynamics of signaling can elicit distinct cellular responses (Doupé and Perrimon, 2014; Hynes et al., 2013; Lemmon and Schlessinger, 2010; Purvis and Lahav, 2013), and available evidence shows that, even in the same cell type, different dynamics of the same signaling pathway can produce different outcomes (Marshall, 1995; Santos et al., 2007; Sasagawa et al., 2005; Toettcher et al., 2013). Protein kinases mediate cell signaling in animals, and protein phosphorylation is an essential type of post-translational modification involved in signal transduction (Hunter, 2009; Manning et al., 2002; Pawson and Scott, 2005). Therefore, a mechanistic understanding of animal development and tissue biology will benefit from visualizing dynamics of kinase activity in physiological contexts (Doupé and Perrimon, 2014; Hynes et al., 2013; Sopko and Perrimon, 2013). For example, extracellular signal-regulated kinase (ERK) is highly conserved from single-celled organisms to multicellular animals, and many receptor tyrosine kinases (RTKs) that play central roles in regulating animal development converge on the common ERK pathway (Sopko and Perrimon, 2013). Differential dynamics of ERK signaling is believed to be one of the mechanisms that discriminates different RTKs (Hynes et al., 2013; Purvis and Lahay, 2013; Shilo, 2014; Sopko and Perrimon, 2013; Toettcher et al., 2013).

To visualize dynamic kinase signaling in living cells and animals, genetically encoded fluorescent reporters are ideal because they allow non-invasive dynamic imaging. Examples include the kinase reporters that are based on fluorescence resonance energy transfer (FRET) between variants of the GFP, e.g., a cyan and a yellow fluorescent protein derived from the GFP (Miyawaki and Niino, 2015; Oldach and Zhang, 2014; Rodriguez et al., 2017; Sanford and Palmer, 2017). These FRET reporters are valuable tools for imaging kinase signaling in living cells, including protein kinase A (PKA) (Zhang et al., 2001, 2005) and ERK (Harvey

et al., 2008). While FRET reporters achieve subcellular resolution and have recently unveiled compartmentalized kinase signaling in cultured single cells (Mo et al., 2017), *in vivo* use of the FRET reporters is problematic because of their small fluorescence ratio change of acceptor over donor fluorophores (Kardash et al., 2011). Thus, a robust fluorescent reporter for imaging dynamic kinase activity in living animals is much needed. Such a reporter should have large signal-to-noise ratio and fast kinetics in order to detect when and in which tissues and cells a kinase is activated during development and other *in vivo* settings.

Another example of genetically encoded kinase reporters is the phosphorylation-dependent GFP translocation-based kinase reporters, which shuttle between cytoplasm and nucleus of cultured cells upon kinase activation (Regot et al., 2014). Most of these reporters achieve large fluorescence ratio change between cytoplasm and nucleus compared to FRET reporters. For instance, the ERK reporter has 2- to 3-fold fluorescence ratio change upon ERK activation. The PKA reporter (PKA-KTR), in contrast, only achieves ~30% ratio change upon PKA activation (Regot et al., 2014). Robust detection of kinase activity in living animals may require larger dynamic range (fluorescence change) and simpler signal pattern, because *in vivo* imaging is known to be challenging due to tissue autofluorescence and light scattering, rapid cell movement, and shape changes (Kardash et al., 2011).

DESIGN

Since biological performance of kinase reporters is determined by their physical or chemical principles, to overcome the challenges for *in vivo* imaging, we seek to design and demonstrate phase separation-based kinase reporters. The rationale is that fluorescence change of a kinase reporter is determined by kinase activity-dependent changes to the brightness of a digital fluorescence image, which not only depends on intrinsic brightness (= quantum yield × extinction coefficient) of the fluorophore but also is proportional to the number of imaged fluorophores per pixel (Waters, 2009). FRET kinase reporters report changes of intrinsic brightness of donor and/or acceptor fluorophores via FRET (Fairclough and Cantor, 1978). A different approach is to design reporters that change the number of imaged fluorophores per pixel. We thus decided to design a different class of fluorescent reporters that, upon kinase activation, phase-separate and form highly concentrated droplets.

Our design was inspired by a recent work showing that multivalent interactions drive protein phase separation and formation of protein droplets (Li et al., 2012). Therefore, we rationally designed multivalent protein-protein interaction (PPI)-driven phase separation-based kinase reporters; (1) to introduce multivalency, we utilized *de novo* designed homo-oligomeric coiled coils; (2) to sense kinase activity, we introduced kinase activity-dependent PPI—a consensus kinase substrate sequence (a phosphopeptide) and a phosphopeptide-binding domain; and (3) to obtain fluorescence signal, we incorporated EGFP into the reporter. We chose EGFP as the fluorophore because its fluorescence requires no cofactor except molecular oxygen and because its use in living cells and animals has been validated in many contexts (Tsien, 1998, 2009).

The phase separation-based kinase reporter is called SPARK (separation of phases-based activity reporter of kinase). Upon kinase activation, the kinase substrate peptide is phosphorylated and then interacts with phosphopeptide-binding domain; the resulting

multivalent PPI leads to highly concentrated $(10\times)$ EGFP droplets. The intensive brightness and punctal signal pattern enable SPARK to robustly detect kinase activity *in vivo*. SPARK has ~20-s response time and is reversible, with no perturbation of kinase signaling and no apparent toxicity in transgenic animals. The modular design of SPARK allows straightforward design of reporters of other kinases by swapping the kinase substrate sequence and the phosphopeptide-binding domain.

RESULTS AND DISCUSSION

Design of Rapamycin-Inducible Protein Phase Separation

To develop phase separation-based kinase reporters, we first sought to design a general platform for inducible protein phase separation via multivalent PPI. We chose the well-characterized FKBP and Frb as an inducible PPI system since their interaction can be robustly induced by rapamycin (Banaszynski et al., 2005). To design multivalent reporters, it is not ideal to use multiple repeat protein domains, which not only will make the reporter protein extremely large but also may be problematic for making transgenic animals (e.g., potential DNA recombination). Thus, instead of using multiple repeats, we turned to *de novo*-designed homo-oligomeric coiled coils. We named these coiled coils as HO-Tag (homo-oligomeric tag). Most *de novo*-designed coiled coils are short peptides, ~30 amino acids (Woolfson et al., 2015), and are, therefore, ideal tags to introduce multivalency. We chose seven coiled coils previously characterized in protein *de novo* design studies (Grigoryan et al., 2011; Huang et al., 2014; Thomson et al., 2014). These coiled coils are here named from HOTag1 to HOTag7 (Figure S1A). To achieve high multivalency, we chose coiled coils that have high stoichiometry, forming tetramer, pentamer, hexamer, and heptamer.

Characterization of these HOTags using the FKBP and Frb PPI pair in HEK293 cells showed that HOTag3 and HOTag6 were most robust in driving rapamycin-inducible protein droplet formation over a wide range of protein concentrations (see the Star Methods and Figures S1A–S1D, S2, and S3). HOTag3 and HOTag6 were highly soluble when fused to FKBP and EGFP fusion (Figures 1A and 1B). To examine kinetics of the EGFP phase separation upon rapamycin-induced PPI in living cells, we conducted time-lapse fluorescence imaging, which revealed that fluorescent droplets formed within 2 min upon rapamycin addition (Figure 1C; Movie S1). Since this includes the time necessary for the rapamycin to diffuse through the cell membrane, the kinetics of droplet formation are expected to be faster (see below). The fluorescent droplets could be qualitatively and easily identified because of their brightness and signal pattern. They could also be quantified and plotted as a fluorescent histogram at pixel level, which showed high brightness and large signal (Figure 1C). Since the hexameric HOTag3 and the tetrameric HOTag6 (Figure S1E) can robustly drive protein phase separation upon protein interaction, we chose HOTag3/6 as the pair for use in kinase reporters.

Protein phase separation and formation of droplets can be attributed to two main factors: PPI and multivalency (Banani et al., 2017; Hyman et al., 2014; Li et al., 2012). In our system, rapamycin induced PPI between FKBP and Frb whereas multivalency was introduced by the HOTags. Rapamycin and HOTags induced multivalent PPIs that drove protein phase separation and formation of EGFP droplets (Figure 1D, EGFP is not shown); the addition of rapamycin induced protein separation into two phases: a dilute phase and a concentrated phase. The high-concentration phase was the protein condensate or droplet (Figure 1D). At equilibrium state, the chemical potential of the dilute and condensed phases were equivalent for both proteins and solvent. Our findings show that the paired HOTag3 and HOTag6 are excellent multivalent tags capable of driving protein phase separation upon PPI over a large range of protein concentrations. We suggest that this HOTag pair will be a valuable tool for many applications, including the induction of protein phase separation for biological investigation of function and for characterizing the consequences of protein phase separation (Banani et al., 2017).

Design of Phase Separation-Based Activity Reporter of Protein Kinase A

To determine whether HOTag3 and HOTag6 can be used to design a SPARK, we generated a construct for detecting PKA activity. PKA is an important mediator of G protein-coupled receptor (GPCR) signaling (Taylor et al., 2012), but the signal of current PKA reporters, including both FRET- and translocation-based reporters, is small, with ~30% ratio change of the fluorescence (Regot et al., 2014; Zhang et al., 2005).

To design a SPARK-based PKA reporter (named PKA-SPARK), we utilized a consensus peptide sequence that is specifically phosphorylated by PKA (Durocher et al., 2000). To introduce PKA activity-dependent PPI, we chose forkhead-associated domain 1 (FHA1), which is a phosphothreonine-binding domain (Durocher et al., 2000). Both motifs have been successfully used in the FRET-based PKA reporters (Zhang et al., 2005). We fused the PKA substrate sequence to EGFP followed by HOTag3, and we fused FHA1 to HOTag6. These two fusion sequences were connected by a self-cleaving 2A sequence (Figure 2A). The rationale is that substrate and the FHA1 induce PKA activity-dependent PPI, and the two HOTags introduce multivalency; upon PKA activation, multivalent PPIs are induced, leading to phase separation and formation of EGFP droplets (Figure 2A).

To test this construct, we added the β -adrenergic agonist isoprenaline to HEK293 cells that endogenously express β -adrenergic receptors (β ARs). β AR activation induces guanosine diphosphate (GDP) for guanosine triphosphate (GTP) exchange in the heterotrimeric G protein Gs, and GTP-bound Gas activates adenylyl cyclase (AC), which converts ATP into cyclic AMP (cAMP) that activates PKA. Upon the addition of the β AR agonist, fluorescent droplets formed in the reporter-expressing HEK293 cells (Figure 2B). In contrast, preincubation with the PKA inhibitor H89 abolished isoprenaline-induced droplet formation (Figure 2B). These results suggest that PKA activation induced EGFP phase separation. To confirm that the phase separation results from phosphorylation of PKA-SPARK at the expected threonine residue, we changed the threonine to alanine in the PKA substrate sequence (LRRATLVD). This mutant PKA-SPARK did not form fluorescent droplets in cells stimulated with isoprenaline. These results suggest that phosphorylation of the single

threonine is responsible for the PKA activity-dependent phase separation of the reporter. Lastly, removal of either HOTag3 or HOTag6 or both abolished droplet formation upon isoprenaline stimulation, indicating that both HOTags are required for droplet formation. Our data thus suggest that PKA activity-triggered multivalent interaction between FHA1-HOTag6 and the PKA substrate peptide-EGFP-HOTag3 leads to EGFP phase separation.

We next asked whether phase separation of the reporter is reversible, i.e., whether the EGFP droplets disassemble or dissolve when PKA is inactivated. We pre-incubated cells with isoprenaline to induce phase separation, then removed isoprenaline and added H89 to inactivate PKA. The EGFP droplets disassembled rapidly (Figure 2C), which is consistent with studies that used a FRET PKA reporter (AKAR2) and reported that H89 inactivates isoprenaline-induced PKA activation (Zhang et al., 2005). As a control, the EGFP droplets were stable over time in the cells with removal of isoprenaline but addition of medium without H89 (Figure 2C). The reversibility of the EGFP droplets suggests that the droplets are accessible to protein phosphatases, which is consistent with previous studies that protein droplets not only contain scaffold proteins responsible for droplet formation but also contain client proteins, which interact with scaffold proteins but are not required for the droplet formation (Banani et al., 2016, 2017). To examine whether PKA-SPARK could quantitatively measure the droplets' disassembly over time after PKA inhibition, we defined SPARK signal as the sum of fluorescent droplets' pixel intensity divided by sum of cells' pixel intensity (Figure 2C). Upon the addition of H89, the SPARK signal decreased over time. In contrast, for the control the SPARK signal remained constant over time (Figure 2C). These results suggest that SPARK signal quantitatively described droplets' disassembly upon PKA inhibition.

We also conducted time-lapse fluorescence imaging of PKA-SPARK at 10 s/frame. Fluorescent droplets formed after the addition of isoprenaline, and the SPARK signal increased over time (Figure 2D). PKA activation upon the addition of isoprenaline was confirmed by AKAR2 (Figure 2E). We detected small droplets of PKA-SPARK as early as at 20 s upon the addition of isoprenaline (Figure 2F). The quick activation of PKA is consistent with rapid activation of the BAR-cAMP-PKA pathway. The small droplets grew quickly and coalesced into large and intensively fluorescent droplets, which were up to 10fold brighter than the cellular fluorescence before PKA activation (Figure 2F). In contrast, PKA-KTR showed much smaller fluorescence change (Figure 2G). Therefore, PKA-SPARK achieved up to a 10-fold fluorescence increase, which was advantageous to AKAR2 and PKA-KTR, which had ~10% (Figure 2E) and ~30% (Figure 2G) fluorescence ratio change, respectively. Our results demonstrate that PKA-SPARK achieves a large dynamic range, high brightness, and fast kinetics for reporting kinase activity. Furthermore, PKA-SPARK could be used with other fluorescent proteins for cellular imaging, such as histone 2B (H2B)-fused monomeric infrared fluorescent protein (mIFP) (Yu et al., 2015) and a red fluorescent protein mCherry (Shaner et al., 2004) in the cells (Figure S4A; Movie S2). We also verified that PKA-SPARK could detect PKA activity over a wide range of reporter expression (Figure S4B). Lastly the concentration of PKA-SPARK (2–10 μ M) appeared to have no effect on the kinetics of droplet assembly (Figure S4C) and disassembly (Figure S4D).

While quantitative analysis can be performed, we emphasize that the advantage of PKA-SPARK is that its intensive fluorescence and simple signal pattern allow easy and straightforward examination of PKA signaling in a qualitative way, based on raw images with no requirement of quantitative data analysis (Movie S3). This is in contrast to the previous PKA reporters, including AKAR2 and PKA-KTR, in which the small fluorescence change makes it difficult to identify PKA activity without quantitative data analysis. Therefore, the SPARK technology represents another class of fluorescent reporters for direct visualization of kinase signaling.

SPARK Visualizes Dynamic PKA Activity upon GPCR Activation

To determine whether PKA-SPARK could reveal the dynamics of PKA activity, we activated PKA via two types of GPCRs: β ARs and adenosine receptors (ARs) (Chen et al., 2013; Higley and Sabatini, 2010; Lebon et al., 2011; Ohta et al., 2006). We added isoprenaline to activate endogenous β ARs in HEK293 cells. EGFP droplets formed and persisted for at least 3 hr, suggesting sustained PKA activity (Figure S5; Movie S4). Pre-incubation of cells with the β AR antagonist carvedilol abolished the sustained PKA-SPARK signals induced by isoprenaline (Figure S6; Movie S5).

To activate endogenous ARs in HEK293 cells, we added adenosine, but unlike the prolonged PKA activation via βAR, we detected oscillatory dynamics of PKA-SPARK. Some cells displayed multiple cycles of PKA-SPARK phase separation (Figures 3A and 3B; Movies S6 and S7). The first pulse of PKA-SPARK droplets lasted for about an hour for most cells. Subsequent pulses had relatively short duration, mostly 10–30 min. But shorter (as low as 3 min) and longer (up to 1 hr) periods were also observed (Figure 3B). The AR antagonist caffeine abolished the PKA-SPARK signals (Figure S7; Movie S8) (Fredholm et al., 1999; Jazayeri et al., 2017). While our data unveil interesting oscillatory PKA activities when HEK293 cells are stimulated with adenosine, detailed mechanisms of the oscillatory behavior will require further investigations.

SPARK Can Be Used to Design an Activity Reporter for ERK

The modular design of SPARK should lend itself to use for other kinases. To determine whether the SPARK design generalizes, we chose the highly conserved kinase ERK, which regulates numerous biological processes during animal development, including cell proliferation, differentiation, and migration (Chen and Krasnow, 2014; Sopko and Perrimon, 2013). To design a SPARK-based ERK reporter (ERK-SPARK), we utilized the ERK activity-sensing motifs that were used for the FRET-based ERK reporters (Harvey et al., 2008; Komatsu et al., 2011). One of the ERK activity-sensing motifs is a phosphopeptide sequence from Cdc25C, followed by an ERK-specific docking site that determines substrate specificity (Lee et al., 2004; Sheridan et al., 2008). The companion fragment is a WW domain that binds the phosphorylated substrate. To construct the ERK-SPARK, we fused the ERK substrate sequence to EGFP followed by HOTag3 and the WW domain to EGFP followed by HOTag6 (Figure 4A). The two fusions were connected by a self-cleaving 2A sequence. The expectation is that substrate motif and the WW domain induce ERK activity-dependent PPI such that, upon ERK activation, the HOTags and the sensing motifs together

induce multivalent PPI of the reporters, which then phase-separate and form fluorescent droplets.

To characterize ERK-SPARK, we added epidermal growth factor (EGF) to HEK293 cells. EGF binds the EGF receptor, resulting in ERK activation. As expected, EGFP droplets formed upon the addition of EGF (Figure 4B). In contrast, cells expressing a reporter with a mutated CDC25C peptide fragment with a threonine-to-alanine substitution that was not expected to be phosphorylated by ERK did not exhibit the formation of EGFP droplets after EGF treatment (Figure 4B). Because the ERK-docking site was previously shown to be critical for ERK recognition and binding to substrate (Lee et al., 2004; Sheridan et al., 2008), we also generated a reporter lacking the 4-amino acid-docking site. As expected, this mutated reporter did not show phase separation upon the addition of EGF (Figure 4B). Lastly, removal of either HOTag3 or HOTag6 or both abolished droplet formation upon EGF stimulation, indicating that both HOTags are required for droplet formation.

We also conducted time-lapse imaging and analyzed SPARK signal, which revealed transient ERK activation upon EGF addition (Figure 4C). The level of ERK-SPARK signal was correlated with the level of dually phosphorylated ERK (ppERK) measured by western blot (Figure 4C; Figure S8A), suggesting that ERK-SPARK can quantitatively measure ERK activity level. The transient ERK activation in HEK293 cells upon EGF stimulation was also confirmed by the FRET ERK reporter EKAR (Figure S8B).

To examine specificity of ERK-SPARK, we pre-incubated the cells with the ERK inhibitor PD 0325901, which abolished phase separation of the reporter upon the addition of EGF (Figure 4D). Pre-incubation with inhibitors for other MAPK family kinases, specifically the JNK inhibitor SB 203580 or the p38 inhibitor VIII, did not abolish the formation of EGFP droplets upon EGF addition (Figure 4D). These results suggest that ERK-SPARK specifically reports ERK activity under the conditions used.

Several GPCRs have been reported to activate ERK signaling (DeFea et al., 2000; Lefkowitz and Shenoy, 2005; Luttrell et al., 2001). To investigate whether ERK-SPARK might detect GPCR-induced ERK activity, we added the peptide SFLLRN to HEK293 cells. This hexapeptide is a full agonist for protease-activated receptor-1 (PAR1) (Coughlin, 2000; Vu et al., 1991a, 1991b). Time-lapse imaging revealed abundant EGFP droplets upon addition of the agonist; the signal peaked at around 6–9 min and dissolved between 15 and 20 min (Figure 4E; Movie S9). These results suggest that ERK-SPARK detects ERK activity via endogenous PAR1 in HEK293 cells. Furthermore, the PAR1-induced ERK activity appears to be transient, lasting about 10 min. Therefore, ERK-SPARK will be a useful tool for imaging ERK signaling that is activated via various pathways, including RTKs and GPCRs.

SPARK Reports ERK Activity in Animal Tissues without Perturbing Signaling or Development

As FRET-based kinase reporters are problematic for animal imaging due to their weak signal, we decided to determine whether the SPARK-based kinase reporters are applicable to animal imaging. First, we created an ERK-SPARK transgenic line of *Drosophila*, and we expressed the reporter in the air sac primordium (ASP) of the *Drosophila* tracheal system. In

the ASP, fibroblast growth factor (FGF) is an essential mitogen and FGF receptor (FGFR)induced ERK signaling has been well established (Sato and Kornberg, 2002; Shilo, 2014). Indeed, we detected EGFP droplets in the ASP (Figure 5A). To confirm that the observed EGFP droplets resulted from a phosphorylation event, we expressed an ERK-unresponsive mutant of the reporter in which the threonine was changed to alanine. Although cells in the ASP had green fluorescence, no EGFP droplets were detected, suggesting that formation of the EGFP droplets in the ASP is dependent on ERK phosphorylation of the reporter.

Next, we expressed a dominant-negative form of FGFR (FGFR^{DN}) in the trachea (Reichman-Fried and Shilo, 1995); it abolished EGFP droplets. In contrast, with a brief pulse of elevated expression of FGF (via heat shock in an animal fly carrying an HS-FGF transgene) (Sato and Kornberg, 2002), the ERK-SPARK signal became more abundant and the EGFP droplets appeared throughout the organ. These data thus suggest that the observed ERK activation is induced by FGF signaling, consistent with previous studies (Sato and Kornberg, 2002).

Second, we expressed ERK-SPARK in the *Drosophila* wing imaginal disc, a tissue that requires epidermal growth factor receptor (EGFR)-dependent signaling (Sturtevant et al., 1994; Gabay et al., 1997). To examine ERK-SPARK in this tissue, we expressed both ERK-SPARK and the ERK-unresponsive mutant ($T \rightarrow A$ mutation). The imaginal disc that expressed ERK-SPARK had abundant fluorescent droplets (Figure 5B; Movie S10). In contrast, discs that expressed the mutant ERK-SPARK had no droplets. Most of the fluorescent droplets were at the dorsoventral boundary and distal hinge of the wing pouch, a pattern that overlaps with the regions that stain with antibody directed against doubly phosphorylated ERK (dpERK) (Gabay et al., 1997). To determine whether expression of the reporter ERK-SPARK perturbed ERK signaling, we first compared dpERK staining in discs that expressed ERK-SPARK with control discs, and we did not detect an apparent difference (Figure S9A).

Lastly, to examine if the expression of ERK-SPARK affects animal development, we compared the adult wings that are developed from wing imaginal discs. Flies expressing ERK-SPARK developed normal wings that were indistinguishable from wings of control flies (Figure 5C). This suggests that ERK-SPARK does not perturb wing development and is consistent with the dpERK staining that showed no apparent difference between the control and the ERK-SPARK-expressing flies. We further examined viability of the transgenic fly, which was not affected when ERK-SPARK was expressed in the trachea or in the whole animal (Figure S9B). These results suggest that ERK-SPARK is a valid indicator of ERK activity in animal tissues.

To demonstrate PKA-SPARK, we expressed the reporter in the wing imaginal discs, and we observed that cells in the pouch region of the wing discs had evenly distributed green fluorescence (Figure S9C), suggesting expression of the reporter. We then expressed a constitutively active form of the alpha subunit of stimulatory G protein (Ga.s*), which activates AC for cAMP production and results in PKA activation (Quan et al., 1991; Wolfgang et al., 1996). EGFP droplets were detected in the wing discs subsequent to ectopic

expression of Ga.s* (Figure S9D). We conclude that PKA-SPARK enables visualization of PKA activity in the *Drosophila* tissue.

SPARK Can Be Used in Live Imaging to Visualize the Dynamics of ERK Signaling In Vivo

Live imaging of kinase activities in animals such as *Drosophila* is technically challenging due to the small dynamic range of the previous reporters and optical issues, such as tissue autofluorescence and light scattering as well as rapid cell movements and shape changes (Kardash et al., 2011). Since the SPARK reporters have a large dynamic range and a simple signal pattern that is readily detected, they may enable imaging of dynamic kinase activity in developing animals. To determine whether ERK-SPARK could be used in live imaging to visualize dynamics of ERK signaling during animal development (Shilo, 2014; Sopko and Perrimon, 2013), we focused on the tracheal system, which is accessible to live imaging and given that FGFR/ERK signaling plays a critical role in tracheal development during metamorphosis at the pupal stage (Chen and Krasnow, 2014), but the dynamics of ERK signaling are unknown. To visualize ERK signaling, we expressed ERK-SPARK in the tracheal cells by crossing the UAS-ERK-SPARK line with the *btl*-GAL4 line.

Time-lapse fluorescence imaging of the trachea between 6 and 11 hr after puparium formation (APF) revealed transient ERK signaling (Figures 6A and 6B). First, EGFP green fluorescence was observed along the dorsal trunk, indicating that ERK-SPARK is expressed in the tracheal cells. Few EGFP droplets were detected between 6 and 7.5 hr APF. Abundant EGFP droplets appeared between 7.5 and 9.5 hr APF, indicating that ERK was activated in the tracheal cells during this period. In the following 2 hr, few EGFP droplets were detected, suggesting that ERK was inactivated. Our imaging results thus indicate that ERK activity was transiently activated in the trachea between 6 and 11 hr APF (Figure 6C; Movie S11). Our work thus demonstrates that the SPARK kinase reporter can be used to visualize dynamic kinase signaling in a live animal during development. SPARK represents another class of fluorescent reporters for imaging kinase-signaling dynamics in animals. The SPARK reporters will find important biological applications in small animals, including *Drosophila* and zebrafish (Figure S10).

The SPARK Technology Can Be Employed to Develop Reporters of Other Kinases

The SPARK technology is based on multivalent PPI-induced phase separation and formation of fluorescent droplets. While the multivalency is introduced by the paired HOTags, the PPI is based on two modular motifs: the kinase-specific substrate domain and the phosphopeptide-binding domain (PBD). The PPI motifs in the SPARK technology are compatible with most FRET-based kinase reporters since they also use these two motifs to sense kinase activity. Therefore, the kinase activity-sensing motifs from the FRET reporters can likely be utilized to develop phase separation-based reporters of many kinases (Oldach and Zhang, 2014). In general, the kinase activity-sensing motif can be adapted from specific kinase substrates to develop the SPARK technology-based kinase reporters, and the PBD may be obtained from the proteins that bind phosphorylated substrates.

In summary, we have developed phase separation-based kinase reporters, SPARKs, which achieve large fluorescence increase, high brightness, and fast kinetics. Because of intensive

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fluorescence, simple signal pattern, and rapid response, SPARKs enable easy and robust visualization of dynamic kinase signaling in living animals in a qualitative way, based on raw images with no requirement for quantitative data analysis. Because of the modular design, other fluorescent proteins may be used to develop multicolor SPARK for other contexts. Furthermore, development of other homo-oligomeric coiled coils that are orthogonal to HOTag3/6 will enable simultaneous imaging of multiple kinase activities. Such multicolor SPARKs could be valuable tools for visualizing signaling networks in living animals.

Limitations

The main limitation of the SPARK technology is spatial resolution, which is poorer than the FRET-based kinase reporters. On the other hand, compared to the KTR-based kinase reporters, its spatial resolution is better since, in principle, it can distinguish kinase activities in the cytoplasm versus nucleus.

STAR ***** METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Chemicals, Peptides, and Recombinant Proteins		
Calcium Phosphate Transfection Kit	ThermoFisher Scientific	CAT# K278001
Lipofectamine 2000 Transfection Reagent	ThermoFisher Scientific	SKU#11668-019
Isoproterenol	Sigma-Aldrich	1351005; CAS: 51-30-9
Ser-Phe-Leu-Leu-Arg-Asn-amide trifluoroacetate salt [PAR-1 (Protease-Activated Receptor) Selective Activating Peptide]	Sigma-Aldrich	S1820; CAS: 141923-40-2 (free base)
Carvedilol	Sigma-Aldrich	C3993; CAS: 72956-09-3
Adenosine	Sigma-Aldrich	A9251; CAS: 58-61-7
Caffeine	Sigma-Aldrich	C0750; CAS:58-08-2
JNK Inhibitor VIII	Sigma-Aldrich	420135; CAS 894804-07-0
P38 inhibitor, SB 203580	Sigma-Aldrich	S8307; CAS: 152121-47-6
ERK inhibitor, PD 0325901	Sigma-Aldrich	PZ0162 ; CAS: 391210-10-9
H-89 dihydrochloride hydrate	Sigma-Aldrich	B1427; CAS:130964-39-5 (anhydrous)
Experimental Models: Cell Lines		
293T/17	ATCC	CRL-11268
Experimental Models: Organisms/Strains		
D. melanogaster: Wild type: w1118	Bloomington Drosophila Stock Center	BDSC: 3605
D. melanogaster: btl-GAL4	Sato and Kornberg, 2002	N/A
D. melanogaster: hs-FGF	Sato and Kornberg, 2002	N/A
D. melanogaster: UAS-FGFR ^{DN}	Reichman-Fried and Shilo, 1995	N/A

REAGENT or RESOURCE	SOURCE	IDENTIFIER
D. melanogaster: tub-Gal4	Bloomington <i>Drosophila</i> Stock Center	BDSC:5138
D. melanogaster: PKA-SPARK	This paper	N/A
D. melanogaster: ERK-SPARK	This paper	N/A
D. melanogaster: ERK-SPARK (T mutated to A)	This paper	N/A
Recombinant DNA		
Plasmid: H2B-mIFP	Yu et al., 2015	Addgene: 56223
Plasmid: mCherry	Shaner et al., 2004	
Plasmid: pcDNA3-FKBP-EGFP-HOTag1	This paper	N/A
Plasmid: pcDNA3-FKBP-EGFP-HOTag2	This paper	N/A
Plasmid: pcDNA3-FKBP-EGFP-HOTag3	This paper	N/A
Plasmid: pcDNA3-FKBP-EGFP-HOTag4	This paper	N/A
Plasmid: pcDNA3-FKBP-EGFP-HOTag5	This paper	N/A
Plasmid: pcDNA3-FKBP-EGFP-HOTag6	This paper	N/A
Plasmid: pcDNA3-FKBP-EGFP-HOTag7	This paper	N/A
Plasmid: pcDNA3-Frb-EGFP-HOTag1	This paper	N/A
Plasmid: pcDNA3-Frb-EGFP-HOTag2	This paper	N/A
Plasmid: pcDNA3-Frb-EGFP-HOTag3	This paper	N/A
Plasmid: pcDNA3-Frb-EGFP-HOTag4	This paper	N/A
Plasmid: pcDNA3-Frb-EGFP-HOTag5	This paper	N/A
Plasmid: pcDNA3-Frb-EGFP-HOTag6	This paper	N/A
Plasmid: pcDNA3-Frb-EGFP-HOTag7	This paper	N/A
Plasmid: pcDNA3-LRRATLVD-EGFP- HOTag3 2A FHA1-HOTag6 (i.e., PKA- SPARK)	This paper	N/A
Plasmid: pcDNA3-LRRAALVD-EGFP- HOTag3 2A FHA1-HOTag6	This paper	N/A
Plasmid: pcDNA3- PDVPRTPVDKAKLSFQFP-EGFP-HOTag3 2A WW-EGFP-HOTag6 (i.e., ERK-SPARK)	This paper	N/A
Plasmid: pcDNA3- PDVPRAPVDKAKLSFQFP-EGFP-HOTag3 2A WW-EGFP-HOTag6	This paper	N/A
Plasmid: pcDNA3-PDVPRTPVDKAKLS- EGFP-HOTag3 2A WW-EGFP-HOTag6	This paper	N/A
Plasmid: pIHEU-LRRATLVD-EGFP-HOTag3 2A FHA1-HOTag6 (PKA-SPARK for expression in <i>Drosophila</i>)	This paper	N/A
Plasmid: pIHEU-PDVPRTPVDKAKLSFQFP- EGFP-HOTag3 2A WW-EGFP-HOTag6 (ERK- SPARK for expression in <i>Drosophila</i>)	This paper	N/A
Plasmid: pIHEU-PDVPRAPVDKAKLSFQFP- EGFP-HOTag3 2A WW-EGFP-HOTag6	This paper	N/A
Plasmid: pminiTol2-LRRATLVD-EGFP- HOTag3 2A FHA1-HOTag6 (PKA-SPARK for expression in zebrafish)	This paper	N/A

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Software and Algorithms		
NIS-Element	Nikon	N/A
Fiji ImageJ	Open	https://fiji.sc/

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for reagents may be directed to, and will be fulfilled by, the Lead Contact, Xiaokun Shu (Xiaokun. shu@ucsf.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animals—*D. melanogaster w1118* was used as the wild-type strain. Fly crosses were maintained at 25°C. The *Drosophila* transgenic lines were generated in this study by using attP insertion on 3rd chromosome.

Cell culture—The HEK293T/17 cells were passaged in Dulbecco's Modified Eagle medium (DMEM) supplemented with 10% Fetal Bovine Serum (FBS), non-essential amino acids, penicillin (100 units/mL) and streptomycin (100 µg/mL). All culture supplies were obtained from the UCSF Cell Culture Facility.

METHOD DETAILS

Plasmid construction-All plasmid constructs were created by standard molecular biology techniques and confirmed by exhaustively sequencing the cloned fragments. To create FKBP-EGFP-HOTag fusions, FKBP was first cloned into pcDNA3 containing EGFP. Various HOTag (HOTag1 to HOTag7) was then each cloned into the pcDNA3 FKBP-EGFP construct, resulting in pcDNA3 FKBP-EGFP-HOTagX (X = 1, 2, 3, ..., 7). Similar procedures were carried out to produce pcDNA3 Frb-EGFP-HOTagY (Y = 1, 2, 3, ..., 7). To create PKA-SPARK, DNA fragments encoding LRRATLVD was inserted to replace FKBP in the pcDNA3 FKBP-EGFP-HOTag3 construct. DNA sequence encoding the FHA1 domain from the FRET-based PKA reporter AKAR was fused to HOTag6. A 2A sequence was added before the FHA1. The DNA fragment encoding 2A-FHA1-HOTag6 was then cloned into the pcDNA3 LRRATLVD-EGFP-HOTag3 to produce pcDNA3 LRRATLVD-EGFP-HOTag3 2A FHA1-HOTag6 (i.e., PKA-SPARK). To create ERK-SPARK, a DNA sequence encoding PDVPRTPVDKAKLSFQFP was introduced to replace FKBP in the pcDNA3 FKBP-EGFP-HOTag3 construct. The DNA sequence encoding WW domain was introduced to replace Frb in the pcDNA3 Frb-EGFP-HOTag6. The 2A sequence was then added before WW domain. The DNA fragment encoding 2A-WW-EGFP-HOTag6 was then cloned into the pcDNA3 PDVPRTPVDKAKLSFQFP-EGFP-HOTag3 to produce the final construct pcDNA3 PDVPRTPVDKAKLSFQFP-EGFP-HOTag3 2A WW-EGFP-HOTag6 (i.e., ERK-SPARK). The phosphothreonine was changed to alanine in the PKA-SPARK and ERK-SPARK using site-directed mutagenesis (Agilent). To create plasmids for making transgenic Drosophila, the DNA fragement encoding PKA-SPARK or ERK-SPARK was cloned into pIHEU construct. To create plasmid for expression in zebrafish, PKA-SPARK was cloned into pminiTol2 construct.

Characterization of HOTags in mammalian cells—To robustly induce protein phase separation or droplet formation via protein interaction, the HOTags should meet several requirements. First, HOTags themselves should not form protein droplets even at high concentration in cells. We therefore chose coiled coils that have been reported to be highly soluble. For example, the pentameric HOTag1 was previously shown to be soluble in aqueous solution (Huang et al., 2014). However, most of these *de novo* designed coiled coils were characterized using synthesized peptides and had not been expressed in mammalian cells. It was thus unclear whether they fold properly in the cellular context and whether they are soluble in mammalian cells. We transfected the fusion constructs (FKBP-EGFP-HOTag) into human embryonic kidney 293 (HEK293) cells, and fluorescence imaging indicated that HOTag3 and HOTag6 (Figure 1B), along with HOTag2, 5, and 7 (Figure S1B), were soluble even at high expression level (~20 μ M [see "Estimation of absolute concentrations of EGFP in cells," Figure S2]). HOTag1 was not soluble even at low expression level (~0.3 μ M, Figure S1B). Thus, we decided to abandon HOTag1 and focus on the other six HOTags.

In order to achieve PPI-inducible protein phase separation, the second task was to identify pairs of HOTags that do not cross-interact to form protein droplets in the absence of PPI-induction. To find soluble HOTag pairs, we co-expressed two fusion constructs (FKBP-EGFP-HOTagX and Frb-EGFP-HOTagY) in HEK293 cells (Figure 1C; Figures S1C and S1D). Fluorescence imaging indicated that none of the fifteen pairs formed protein droplet at ~10 μ M expression level in cells (Figure S1C). We then examined whether the fusion proteins could be induced to form droplets. Upon addition of rapamycin, time-lapse fluorescence imaging revealed that the pairs formed protein droplets at ~10 μ M (Figures S1C and S1D). However, only two pairs, HOTag2/7 and HOTag3/6, formed droplets at ~1 μ M (Figures S1C and S1D). At ~0.3 μ M expression level, only the HOTag3/6 pair formed droplets upon rapamycin-induced PPI (Figures S1C and S1D). Therefore, our data indicate that the hexameric HOTag3 and the tetrameric HOTag6 (Figure S1E) can robustly drive protein phase separation upon protein interaction. We thus chose HOTag3/6 as the pair for use in kinase reporters.

Western blot of dually phosphorylated ERK—Cells were lysed in 1X SDS sample buffer and resolved in NuPAGE 4%–12% Bis-Tris Protein Gels. After transferring, Phosphop44/42 MAPK (Erk1/2) (Thr202/Tyr204) (D13.14.4E) Rabbit mAb (CST, 4370S) were used at 1:1000. HRP-Donkey-anti-Rabbit (Invitrogen A16029) antibodies were used at 1:10,000. Pierce ECL standard substrate buffer was used for visualization against film. The membrane was then stripped with Pierce Western Blot Stripping Buffer. HRP-anti-actin (Santa Cruz Biotechnology, sc-47778 HRP) was used at 1:10,000 for detecting beta-actin as a loading control.

Drosophila husbandry—Flies were reared on standard cornmeal and agar medium at 25°C. The L3 wandering larvae were staged in cold PBS. Wing discs and associated trachea were immersed in a drop of PBS underneath a coverslip. A previously established fly line Btl-RFP-moe is used to specifically label the tracheal progenitors (Chen and Krasnow 2014). Btl-RFP-moe does not encompass the entire enhancer of breathless gene as btl-Gal4, but it

turns out to be the best reagent to specifically label tracheal progenitor cells (Chen and Krasnow 2014).

Live Drosophila imaging—White pupae of *Drosophila* (0hr APF) were brief washed and cleaned. Pupae were mounted within a drop of halocarbon oil 700 (Sigma) between two stripes of vacuum grease. Next, pupae were mixed well with oil, positioned with forceps and rolled so that a single trachea is up for optimal imaging of Tr4 and Tr5 metamere. Then, pupae were immobilized by a 22X30mm No.1.5 cover glass.

Fluorescence imaging was conducted on a Nikon Eclipse Ti inverted microscope equipped with a Yokogawa CSU-W1 confocal scanner unit (Andor), a digital CMOS camera ORCA-Flash4.0 (Hamamatsu), a ASI MS-2000 XYZ automated stage (Applied Scientific Instrumentation), a Nikon CFI Plan Apochromatic 20X dry (N.A. 0.75) objective, CFI apochromatic confocal LWD 40X WI λ S water objective (N.A. 1.15), CFI apochromatic TIRF 60X oil objective (N.A. 1.49). Laser inputs were provided by an Integrated Laser Engine (Spectral Applied Research) equipped with laser lines (Coherent) 488 nm for GFP imaging, 561 nm for mCherry imaging, and 642 nm for mIFP imaging. The confocal scanning unit was equipped with the following emission filters: 525/50-nm for GFP imaging, 610/60-nm for mCherry imaging, 732/68-nm for mIFP imaging.

Fluorescence imaging of *Drosophila* larvae or pupae was carried out in an incubation chamber maintained at 25°C. The animals were imaged using a 20X dry objective for ~10 hours with image acquisition every 10 minutes.

Live cell imaging—HEK293T/17 cells were transiently transfected with the plasmid using calcium phosphate transfection reagent or lipofectamine. Cells were grown in 35 mm glass bottom microwell (14 mm) dishes (MatTek Corporation). Transfection was performed when cells were cultured to ~50% confluence. For each transfection, 4.3 μ g of plasmid DNA was mixed with 71 μ L of 1X Hanks' Balanced Salts buffer (HBS) and 4.3 μ L of 2.5M CaCl₂. Cells were imaged 24 hours after transient transfection.

Time-lapse imaging was performed with the aid of an environmental control unit incubation chamber (InVivo Scientific), which was maintained at 37°C and 5% CO₂. Fluorescence images were acquired with an exposure time of 50 ms for EGFP, 100 ms for mCherry, 200 ms for mIFP. Chemical reagents were carefully added to the cells in the incubation chamber when the time-lapse imaging was started. In particular, to induce interaction between FKBP and Frb, 100 nM rapamycin was added. To activate β ARs, 10 μ M isoprenaline was added. To demonstrate specificity of PKA-SPARK, cells were pre-incubated with 10 μ M H89 before addition of 10 μ M isoprenaline. For inhibition of bARs, cells were incubated with 10 μ M carvedilol for 5 minutes before addition of 2 μ M isoprenaline. To demonstrate specificity of ERK-SPARK, cells were pre-incubated with 10 μ M ERK inhibitor PD 0325901, or 10 μ M JNK inhibitor VIII, or 10 μ M p38 inhibitor SB 203580, before addition of 1 μ M EGF. To activate PAR1, cells were stimulated with 10 μ M PAR1 agonist SFLLRN. Image acquisition was controlled by the NIS-Elements and ImageJ (NIH).

For FRET imaging, cells were transfected with AKAR2 or EKAR. Isoprenaline was added to HEK293 cells expressing AKAR2. EGF was added to HEK293 cells transfected with EKAR. Images at CFP and YFP channel were taken using 405nm laser. The fluorescence images were processed in ImageJ.

Live zebrafish imaging—To demonstrate whether PKA-SPARK can be used in zebrafish, we injected DNA encoding the reporter into zebrafish embryo at one-cell stage. Muscle cells at 24 hr APF were imaged upon addition of isoprenaline, which showed EGFP droplet formation. Our data thus suggests that PKA-SPARK visualizes activation of PKA in zebrafish. Therefore, the SPARK reporters can be used in imaging kinase activity in zebrafish.

Estimation of absolute concentrations of EGFP in cells—To estimate concentration of expressed EGFP in HEK293 cells, we followed a method described in a previous study (Shin et al., 2017). We first purified EGFP and determined its concentration based on its absorbance and extinction coefficent. Then we prepared EGFP at various concentrations between 0.3 μ M and 28 μ M. Next we imaged the EGFP samples under the fluorescence microscope and recorded fluorescence intensity (counts/pixel) for each sample. The net intensity was determined after subtracting the dark background (without sample). The relationship between EGFP concentration and the net fluorescence intensity (counts/ pixel) was plotted (Figure S2). To estimate EGFP concentration in HEK293 cells, we imaged the cells using the same imaging parameters including the laser power, exposure time, and magnification as those used for purified EGFP samples. The pixel intensity in each cell was averaged, which was subtracted by the intensity of a nearby dark area. The plotted curve was used to estimate EGFP concentration in HEK293 cells based on the net intensity.

QUANTIFICATION AND STATISTICAL ANALYSIS

Image analysis—For quantitative analysis of the SPARK signal, images were processed in imageJ. The sum of droplets pixel fluorescence intensity and the cells pixel intensity were scored using Analyze Particle function in imageJ.

Drosophila viability quantification—In viability assay, the numbers of ERK-SPARKexpressing adults and their siblings with balancers were counted. The survival percentage was calculated as observed progeny carrying the ERK-SPARK to the expected number, that is, observed number of non-balancer progeny to the offspring with a balancer. The adult progeny were obtained from at least three independent crosses. The survival ratios were plotted as bar graph in Prism.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- Rapamycin-inducible protein phase transition is developed
- The kinase reporter SPARK achieves intensive brightness and simple signal
 pattern
- SPARK reveals dynamic ERK signaling during tissue homeostasis in vivo
- SPARK technology enables modular design of other kinases' reporters



Figure 1. Rational Design of Inducible Protein Phase Separation by Protein-Protein Interaction (A) *De novo*-designed coiled coils are used as homo-oligomeric tag (HOTag), which introduces multivalency.

(B) HOTags fused to EGFP and FKBP were expressed in HEK293 cells.

(C) Time-lapse images of cells expressing HOTag fusions upon the addition of rapamycin. FKBP-EGFP-HOTag3 and Frb-EGFP-HOTag6 were co-transfected. Histograms are shown corresponding to the dashed line in the fluorescence images.

(D) Schematic diagram of induced protein phase separation with HOTags. Left: FKBP-EGFP (not shown)-HOTag3 hexamer and Frb-EGFP (not shown)-HOTag6 tetramer are shown. Right: protein condensate or droplet is shown. Scale bar, 10 µm.



Figure 2. Design and Characterization of Phase Separation-Based Activity Reporter of Protein Kinase A

(A) Schematic diagram of the reporter. P represents the phosphate group.

(B) Fluorescence images of the HEK293 cells expressing the PKA reporter PKA-SPARK before and after the addition of beta adrenergic agonist isoprenaline. Cells were not preincubated with other molecules unless otherwise noted. +H89, the PKA inhibitor H89 was added 5 min before the addition of isoprenaline; $T \rightarrow A$, the phosphothreonine in PKA-SPARK was mutated to alanine; HOTag3-, HOTag3 was removed; HOTag6-, HOTag6 was removed; HOTag3-/6-, HOTag3 and HOTag6 were removed.

(C) Time-lapse fluorescence images of cells expressing PKA-SPARK. Cells were preincubated with isoprenaline for 5 min. Isoprenaline was then removed. Upper images were taken after the addition of H89. Lower images were taken after the addition of medium solution without H89. Middle: definition of the SPARK signal is given; see details in the

STAR Methods. Right: normalized SPARK signal (from the cell shown in the left images) was plotted against time.

(D) Fluorescence images of cells stimulated with isoprenaline. Cells expressed PKA-SPARK. Right: normalized PKA-SPARK signal (averaged from the cells shown on images in the left) was plotted against time.

(E) Time-dependent fluorescence ratio change of AKAR2 upon the addition of isoprenaline in HEK293 cells. The normalized ratio of yellow fluorescence of YFP (yellow fluorescent protein) over cyan fluorescence of CFP (cyan fluorescent protein) was plotted against time. (F) Time-lapse images of the zoomed-in area shown in (D). Lower row: fluorescence histograms correspond to the dashed lines in the upper images. Note: the dashed line of the 120-s image is in a relatively bright area (the upper-left area is relatively dark), and the sum of the entire cell's fluorescence compared to that at time 0 is within 10%.

(G) Fluorescence images of cells stimulated with isoprenaline. Cells expressed a recently developed PKA reporter PKA-KTR. PKA-KTR is based on GFP translocation between cytoplasm and nucleus upon PKA activation. Fluorescence histograms corresponding to the dashed line in the images are shown next to the fluorescence images. Normalized fluorescence of cytoplasm over nucleus (C/N) was plotted against time. Scale bars, 10 µm.



Figure 3. SPARK Visualizes Dynamic PKA Activity upon Adenosine-Induced GPCR Activation (A) Schematic diagram of signaling pathways from the GPCR ligand adenosine to PKA activation. AC, adenylyl cyclase.

(B) Time-lapse fluorescence images upon the addition of adenosine. Images were taken for \sim 3 hr. Scale bar, 10 μ m.

(C) Differential PKA activity in single cells upon the addition of adenosine.



Figure 4. Design and Characterization of Phase Separation-Based Activity Reporter of ERK (A) Schematic diagram of the phase separation-based ERK reporter ERK-SPARK. 2A is a self-cleaving peptide. The phosphothreonine is shown in red. P represents the phosphate group.

(B) Fluorescence images of HEK293 cells expressing ERK-SPARK and its mutant forms before and after the addition of the growth factor EGF. $T \rightarrow A$, the phosphothreonine was mutated to alanine; no FQFP, the docking site FQFP was deleted; no HOTag3, HOTag3 was deleted; no HOTag6, HOTag6 was deleted; no HOTag3/no HOTag6, HOTag6 were both deleted.

(C) Time-lapse fluorescence images of cells expressing ERK-SPARK upon the addition of EGF. Right: normalized SPARK signal (averaged from cells in the left images) was plotted against time; normalized western blot of ppERK was also plotted against time.

(D) Fluorescence images of HEK293 cells expressing ERK-SPARK upon the addition of EGF. The cells were incubated with various inhibitors (left, ERK inhibitor PD 0325901; middle, JNK inhibitor VIII; and right, p38 inhibitor SB 203580).

(E) Time-lapse images of cells expressing ERK-SPARK upon the addition of PAR1 agonist.

In (C) and (E), cells expressing ERK-SPARK were also labeled with mCherry and a nucleus-located monomeric infrared fluorescent protein (mIFP) fused to histone 2B (H2B). Scale bar, $10 \mu m$ (B–E).



does not perturb animal development

Figure 5. SPARK Reports ERK Activity in *Drosophila* Tissues without Perturbing Animal Development

(A) Left: schematic diagrams of the air sac primordium (ASP) and the FGFR/ERK-signaling pathway. Right: fluorescence images show cells in the ASP. Arrow points to EGFP droplet. $T \rightarrow A$, the phosphothreonine of ERK-SPARK is changed to alanine; FGFR^{DN}, a dominant-negative mutant of FGFR; HS-FGF, FGF level was elevated through a brief induction by heat shock (HS).

(B) Left: schematic diagrams of the wing imaginal disc and the EGFR/ERK-signaling pathway. Right: fluorescence images show cells in the wing pouch of the wing imaginal disc. $T \rightarrow A$, the phosphothreonine of ERK-SPARK is changed to alanine (the image is $5 \times$ brightened relative to the left panel).

(C) Top: schematic of an adult wing with veins labeled. Bottom: images show adult wings developed from wing discs expressing ERK-SPARK versus a control. Scale bars, 25 (A)and 50 μm (B).



Figure 6. ERK-SPARK Can Be Used in Live Imaging to Visualize Dynamics of ERK Signaling *In Vivo*

(A) Schematic diagram of the *Drosophila* trachea. Left: tracheal metamere (Tr) 6 is shown. DT, dorsal trunk; TC, transverse connective branch.

(B) Time-lapse fluorescence images of the trachea in intact *Drosophila* pupa. ERK-SPARK was expressed by crossing the UAS-ERK-SPARK transgenic fly with the btl-GAL4 line. Arrowhead points to EGFP droplets.

(C) Normalized ERK-SPARK signal over time. Stars correspond to images in (B). Scale bar, 50 μ m (B).