Spatio-temporally precise activation of engineered receptor tyrosine kinases by light

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Abstract

Receptor tyrosine kinases (RTKs) are a large family of cell surface receptors that sense growth factors and hormones and regulate a variety of cell behaviours in health and disease. Contactless activation of RTKs with spatial and temporal precision is currently not feasible. Here, we generated RTKs that are insensitive to endogenous ligands but can be selectively activated by low-intensity blue light. We screened light-oxygen-voltage (LOV)-sensing domains for their ability to activate RTKs by light-activated dimerization. Incorporation of LOV domains found in aureochrome photoreceptors of stramenopiles resulted in robust activation of the fibroblast growth factor receptor 1 (FGFR1), epidermal growth factor receptor (EGFR) and rearranged during transfection (RET). In human cancer and endothelial cells, light induced cellular signalling with spatial and temporal precision. Furthermore, light faithfully mimicked complex mitogenic and morphogenic cell behaviour induced by growth factors. RTKs under optical control (Opto-RTKs) provide a powerful optogenetic approach to actuate cellular signals and manipulate cell behaviour.

Keywords aureochrome; cell signalling; light-oxygen-voltage (LOV)-sensing domain; optogenetics; synthetic biology; synthetic physiology

Subject Categories Methods & Resources; Signal Transduction

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Introduction

Cell surface receptors act as relays between extracellular stimuli and intracellular signalling cascades and thereby play vital and diverse roles in normal physiology and disease (Robertson et al., 2000; Ashcroft, 2006; Lagerstrom & Schioth, 2008; Casaletto & McClatchey, 2012). A wide range of techniques has been developed to manipulate receptor signalling in order to study how activation affects cell behaviour. Genetic approaches enable conditional and cell type-specific overexpression and deletion of receptor genes, and pharmacological approaches permit activation and inhibition of selected receptor families. Current evidence suggests that receptor signalling is spatially and dynamically regulated. However, it remains challenging to activate or inactivate receptors on timescales shorter than those defined by receptor biosynthesis and degradation, or by drug diffusion and pharmacokinetics. It is further difficult to manipulate receptor function locally, such as in compartments of cells or in selected cells for which no specific genetic markers are available.

Recent work in neurobiology has harnessed the power of light for the control of cellular activity with spatial and temporal precision (Szobota & Isacoff, 2010; Fenno et al., 2011). General advantages of optogenetics over pharmacology and genetics include the ease with which signals can be applied and withdrawn and the ability of rapid and local signal activation. Complementary to light-sensing ion channels and ion pumps, which are now utilized routinely, two approaches were recently developed for optogenetic activation specifically of G protein-coupled receptors (GPCRs). Chimeric proteins combine domains of light-sensing GPCRs with domains of ligand-responsive GPCRs (Airan et al., 2009), while photochromic ligands are synthetic compounds that activate GPCRs by mimicking low-molecular-weight ligands (Levitz et al., 2013). It has been argued that light-activated proteins that perform comparably to physiologically relevant signalling molecules are difficult to engineer (Fenno et al., 2011; Christie et al., 2012; Muller & Weber, 2013; Pathak et al., 2013). The challenges include avoiding background signalling in the absence of light and achieving physiological levels of pathway activation. Indeed, for receptor families other than GPCRs, activation with light has not been achieved.

Here, we utilized rational protein engineering to design receptor tyrosine kinases (RTKs) that are under optical control (Opto-RTKs). RTKs are a large and essential surface receptor family that senses growth factors and hormones. Because extracellular ligands induce and stabilize RTK dimers (Lemmon & Schlessinger, 2010; Simi & Ibanez, 2010), we reasoned that light-activated dimerization may induce receptor activation with light. We first identified protein domains that undergo homodimerization in response to blue light, and we then used these domains to induce receptor trans-phosphorylation and signal propagation to intracellular pathways. Finally, we applied Opto-RTKs to control cellular signalling and cell behaviour.
in human cancer and endothelial cells with spatial and temporal precision.

**Results**

**Selection and expression of light-sensing proteins**

We selected blue light-sensing protein domains that belong to the large light-oxygen-voltage (LOV)-sensing domain superfamily as candidates for light-activated dimerization of RTKs. Light-sensing LOV domains bind flavins as prosthetic groups and act as reversible light sensors in bacteria, fungi and plants. LOV domains control heterogeneous effector domains such as enzymes [for instance, in the flowering plant *Arabidopsis thaliana* and the green alga *Chlamydomonas reinhardtii* (Huang et al., 2002; Kinoshita et al., 2001)] or transcriptional regulators [for instance, in the fungus *Neurospora crassa* and the yellow-green alga *Vaulcheria frigida* (Heintzen et al., 2001; Takahashi et al., 2007)]. Dimerization of LOV domains was proposed to play an important role in effector domain regulation, and a remarkable diversity in dimerization interfaces and lifetimes was observed among LOV domains (Moglich et al., 2010; Zoltowski & Gardner, 2011). As it was initially unclear whether a LOV domain would be suited to activate mammalian RTKs, we compiled an unbiased panel of diverse candidate domains (one from a proteobacterium, two from a fungus, two from algae and two from a plant; Fig 1A, Supplementary Tables S1 and S2, and Materials and Methods). As the majority of these domains was never studied in mammalian cells, we first tested in two mammalian cell lines whether the domains can be expressed and whether expression causes cytotoxicity. We found that the domains were produced efficiently (as assessed by detection of a fluorescent protein fused to the domains; Fig 1B and D) and with no detectable cytotoxicity (as assessed by cellular reduction of a tetrazolium dye; Fig 1C and E).

**Engineering a light-activated fibroblast growth factor receptor**

Our experiments focused on fibroblast growth factor (FGF) receptor 1, a highly conserved key regulator of cell behaviour in, for instance, embryonic development, adult neurogenesis and tumour formation (Deng et al., 1994; Zhao et al., 2007; Yang et al., 2013). We constructed chimeric receptors where each of the seven LOV domains was linked to the intracellular catalytic domain of murine FGFR1 (mFGFR1; Fig 2A). The extracellular ligand-binding modules of mFGFR1 were omitted to obtain chimeras that are not responsive to native ligands. Cells expressing these chimeras should respond to blue light with activation of signalling pathways characteristic for mFGFR1. We performed cell signalling experiments in a custom-built incubator that allows illumination of cells and tissues with light of defined intensity and colour (Materials and Methods). We first examined the mitogen-activated protein kinase (MAPK)/extracellular signal-regulated kinase (ERK) pathway, a key signalling pathway that is activated by FGFs via FGFRs and highly dynamic (Ma et al., 2009; Albeck et al., 2013). To quantify MAPK/ERK pathway activation, we utilized a reporter assay that enables simultaneous evaluation of chimeras in multi-well plates. In this assay, luciferase is produced under the control of an engineered Elk1 transcriptional activator and pathway activation is represented as raw light units (RLU). As a positive control, we employed a modified, chemically inducible mFGFR1 (imFGFR; Welm et al., 2002), which lacks the endogenous ligand-binding modules and is activated by binding of the small chemical ligand AP20187 to a single, engineered FK506 binding protein (FKBP; Clackson, 1998). We found that the chimera incorporating the LOV domain of aureochrome1 from *V. frigida* (mFGFR1-VIAU1-LOV) activated the MAPK/ERK pathway similarly to the positive control (Fig 2B). In particular, no augmented basal pathway activation in the absence of light was observed and pathway induction by light was of comparable magnitude to that by ligand. All other chimeras either exhibited no activity or constitutive activity (Fig 2B). Control experiments showed that: (i) ERK1/2 is phosphorylated upon blue light stimulation in cells transfected with mFGFR1-VIAU1-LOV (Supplementary Fig S1), (ii) blue light had no effect on cells transfected with imFGFR1 (thereby excluding reporter activation by light alone; Fig 2C, left), (iii) blue light had no effect on cells transfected with mFGFR1-VIAU1-LOV with Y271F and Y272F substitutions that result in loss of autophosphorylation and kinase activity (thereby demonstrating that kinase activity of the receptor is required; Fig 2C, middle) and (iv) green light or red light had no effect on cells transfected with mFGFR1-VIAU1-LOV (thereby demonstrating wavelength specificity; Fig 2C, right). Collectively, these results show that mFGFR1-VIAU1-LOV, a chimeric receptor consisting of the catalytic domain of a mammalian RTK and an algal LOV domain, activates the canonical MAPK/ERK pathway in response to blue light. In addition to phosphorylation of ERK, we observed phosphorylation of AKT, the key adapter protein fibroblast growth factor receptor substrate 2 (FRS2) and phospholipase C1 (PLC1) in response to blue light (Supplementary Fig S1). Using luciferase reporters, we also detected activation of additional pathways linked to mFGFR1 (Supplementary Fig S2). We termed the chimeric mFGFR1-VIAU1-LOV receptor ‘Opto-mFGFR1’.

**Dimerization is required for Opto-mFGFR1 activation**

We next investigated dimerization during activation of Opto-mFGFR1 with two independent methods. The first method utilizes a single charge inversion substitution (R195E in Opto-mFGFR1 or imFGFR1; R557E in full-length FGFR1). This substitution prevents formation of a functionally essential, asymmetric kinase domain dimer in FGFR1 (Bae et al., 2010) and inhibited MAPK/ERK pathway activation by imFGFR1 (Fig 3A). We introduced this substitution into Opto-mFGFR1 as a probe for dimer formation during signalling. In HEK293 cells transfected with Opto-mFGFR1-R195E, we detected no MAPK/ERK pathway activation in response to blue light (Fig 3A), indicating that receptor dimerization is required for receptor activation with light. We utilized protein cross-linking as a second method and directly observed Opto-mFGFR1 dimers in response to blue light (Fig 3B). In addition, and as expected, we did not detect trans-activation of the endogenous FGFR1 in response to light stimulation of Opto-mFGFR1 in HEK293 cells that express similar levels of chimeric and endogenous receptors (Supplementary Fig S3). These results, together with the observation that VIAU1-LOV also induces dimerization of a transcription factor in response to blue light (Supplementary Fig S4), point to dimerization as the mechanism underlying Opto-mFGFR1 activation.

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Coupling of Opto-mFGFR1 to cellular signalling pathways

We further investigated how Opto-mFGFR1 couples to cellular signalling pathways. Inhibition of MAPK/ERK kinase 1/2 (MEK1/2) confirmed that signals of the Elk1 reporter originate from activation of the MAPK/ERK pathway (Supplementary Fig S5). We introduced single-site substitutions into Opto-mFGFR1 to test if receptor phosphorylation is required for MAPK/ERK pathway activation. In

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agreement with previous work (Mohammadi et al., 1996), we found that Y271 and Y272 are required for activation of the MAPK/ERK pathway (Fig 2C, middle), while Y81, Y201, Y203, Y348 and Y384 are dispensable (these residues correspond to Y653, Y654, Y463, Y583, Y585, Y730 and Y766 of mFGFR1; Fig 3C). We also tested if a functionally relevant phosphorylation site has been introduced through generation of the chimera. Y447 is the sole surface-accessible Tyr in VfAU1-LOV as judged by inspecting a recent crystal structure of VfAU1-LOV (Mitra et al., 2012), and substitution at this site does not affect Opto-mFGFR1 function (Fig 3C).

**Figure 2. Design and function of mFGFR1-LOV domain chimeric receptors.**

A. Receptor tyrosine kinases such as mFGFR1 consist of the extracellular ligand-binding domain (LBD), single-span transmembrane domain (TMD) and intracellular domain (ICD) [kinase domain (KD) and a C-terminal tail domain (CTD)]. In mFGFR1-LOV domain chimeras, only the ICD is retained to render the protein insensitive to endogenous ligand. The ICD is attached to the membrane using a myristoylation domain (MYR) and LOV domains are incorporated at the ICD C-terminus.

B. MAPK/ERK pathway activation in response to blue light for HEK293 cells that were transfected with chimeric proteins of mFGFR1-ICD and LOV domains. Activation is expressed as induction of a luciferase reporter gene. imFGFR1 is activated by the small molecule dimerizer AP20187.

C. MAPK/ERK pathway activation in response to blue, green and red light for HEK293 cells that were transfected with imFGFR1, Opto-mFGFR1 (mFGFR1-VfAU1-LOV) or kinase dead Opto-mFGFR1 (Y271F, Y272F).

Data information: For (B) and (C): 24 h after transfection, cells were stimulated with light for 8 h followed by detection of luciferase. Light intensity was 1.7–25 μW/mm². Mean values ± SEM for four to 16 independent experiments each performed in triplicates are shown.

LOV domains respond to blue light with formation of a covalent photoadduct between a cysteine residue and the flavin prosthetic group. The photoadduct undergoes thermal reversion, and photoadduct lifetime is an important parameter in the function of light-sensing proteins (Zoltowski et al., 2009). We found that VIAU1-LOV exhibits a photoadduct lifetime that is intermediate among known
LOV domains ($\tau = 625$ s at 20°C, Supplementary Fig S7 and Supplementary Table S1; Moglich et al., 2010; Zoltowski & Gardner, 2011). We then investigated how photoadduct lifetime contributes to Opto-mFGFR1 signalling by reducing the lifetime through a point substitution. It was previously shown that substitution of a cysteinyl-methionine of Opto-mFGFR1 harbouring Tyr substitutions at positions 81 (468), 201 (583), 203 (585), 348 (730), 384 (766) and 447 (3) or combined substitutions to Ala (‘AA’) at positions K37, P40, L41 and R43 (K419, P422, L423 and R425; numbering is relative to start-methionine of Opto-mFGFR1, corresponding positions in mFGFR1 or VIAU1-LOV are given in parentheses). These residues were previously shown to be phosphorylated (Tyr) or required for association of FRS2 (LysProLeuArg).

Data information: For (A) and (C), 24 h after transfection, cells were stimulated with blue light for 8 h followed by detection of luciferase. Mean values ± SEM for two to 16 independent experiments each performed in triplicates are shown. For (A–C): Light intensity was ~25 μW/mm².

Transfer of design principle to other RTKs and LOV domains

A highly desirable feature of engineered molecular systems is generalizability in which novel regulatory components are compatible with several different catalytic components. To test whether VIAU1-LOV is capable of activating other RTKs, we combined it with the catalytic domain of the human epidermal growth factor receptor (hEGFR) and human rearranged during transfection (hRET). We followed the design established in Opto-mFGFR1 and utilized membrane anchors previously shown to allow functional anchoring lifetime of wild-type VIAU1-LOV but longer than that of VIAU1-LOV after substitution.
of these catalytic domains (Muthuswamy et al., 1999; Freche et al., 2005; Allocca et al., 2007). In line with their known signalling capability, robust activation of the MAPK/ERK pathway with blue light was observed in cells transfected with hEGFR and hRET chimeras (Fig 4B). These chimeras were termed ‘Opto-hEGFR’ and ‘Opto-hRET’. We next tested whether LOV domains that resemble VIAU1-LOV can activate mFGFR1. Using database searches, we identified VIAU1-like proteins in the eustigmatophyte *Nannochloropsis gaditana* [N. gaditana putative aureochromel (NgPA1)] and in the golden algae *Ochromonas danica* [O. danica putative aureochromel (OdPA1)]. For mFGFR1 chimeras incorporating the LOV domains of NgPA1 and OdPA1 (NgPA1-LOV and OdPA1-LOV), we also observed activation of the MAPK/ERK pathway with blue light and amplitudes similar to that of the original Opto-mFGFR1 (Fig 4C). We note that the LOV domain is located C-terminally to the effector domain in a similar to that of the original Opto-mFGFR1 (Fig 4C). We note that the LOV domain is located C-terminally to the effector domain in

**Control of signalling in FGF2-responsive cells with spatial and temporal precision**

We next tested the ability of Opto-mFGFR1 to control signalling with spatial and temporal precision in cells that are naturally responsive to FGFs. We first screened cells from different tumour entities for effects of FGF2, a prominent FGF ligand, and we identified two cell lines derived from malignant pleural mesothelioma to be FGF2-responsive (M38K and SPC212; Schmitten et al., 1992; Kahlos et al., 1998). We also investigated human blood endothelial (hBE) cells as an additional cellular model (Schoppmann et al., 2004). We delivered Opto-mFGFR1 into these cells using virus and propagated cells with stable marker gene expression. Although expression levels of Opto-mFGFR1 in these stable M38K, SPC212 and hBE cells were higher than those of endogenous FGRs (Supplementary Fig S8), cell resting states and endogenous receptor function were not compromised (see below). Immunofluorescence microscopy revealed that Opto-mFGFR1 localizes predominantly to the plasma membrane in these cells and also in HEK293 cells (Supplementary Fig S9). In M38K*Opto-mFGFR1* and SPC212*Opto-mFGFR1* cells, we found that 1 min of blue light was sufficient to induce phosphorylation of the receptor and ERK1/2 (Fig 5A). We also observed phosphorylation of AKT and PLCγ1 in SPC212*Opto-mFGFR1* cells (Fig 5B). After cessation of light, phosphorylation of the receptor returned to pre-stimulation levels within 5 min, followed by phosphorylation of ERK1/2, AKT and PLCγ1 within 15–30 min (Fig 5A and B). In hBE*Opto-mFGFR1* cells, 5 min of blue light stimulation was sufficient to produce phosphorylation of the receptor and ERK1/2 (Fig SC). This result demonstrates control over cellular signalling in several cell types with temporal precision on the minute timescale. A remarkable diversity in the temporal signature of cell responses to this brief illumination was observed that likely reflect different activity of adaptor proteins or feedback mechanisms. To demonstrate spatially precise activation of signalling, we created monolayers of SPC212*Opto-mFGFR1* and hBE*Opto-mFGFR1* cells and limited illumination to small circular areas in the layers (areas contain ~500 SPC212*Opto-mFGFR1* cells in Fig 6A or ~2,500 hBE*Opto-mFGFR1* cells in Fig 6B; ~50,000–250,000 surrounding cells were not illuminated). Immunodetection of ERK1/2 phosphorylation revealed activation of the MAPK/ERK pathway in illuminated but not neighbouring cells. This result and illumination of HEK293 cells in arbitrary patterns (Fig 6C) demonstrates local signal activation in genetically homogenous cell layers.

**Controlling cell behaviour with light**

We next analysed whether Opto-mFGFR1 could functionally substitute for ligand activation in M38K and hBE cells. In M38K*Opto-mFGFR1* cells, one-h light stimulation caused an increase in cell proliferation and induced EMT-like morphological and gene expression changes (Fig 7A–C, Supplementary Figs S10, S11 and S12), features characteristic of increased malignancy of cancer cells. In these experiments, changes in cell behaviour induced by light were within error of those induced by FGF2 (Fig 7A and B). Blue light had no effect on control cells or cells that were pre-treated with PD166866, a selective FGFR1 kinase inhibitor (Fig 7B and C). In addition to their prominent role in cancer, growth factors are important inducers of angiogenesis in physiological and pathophysiological conditions (Ferrara et al., 2003; Presta et al., 2005). The ability of several
growth factors to induce angiogenesis is reflected by their ability to activate endothelial cell sprouting in vitro (Ferreras et al., 2012). Blue light illumination induced sprouting in hBE\textsuperscript{Opto-mFGFR1} spheroids to a similar extent as the major proangiogenic peptide VEGFA (Fig 7D and E). These results indicate that activation of Opto-mFGFR1 can functionally substitute signalling by ligands and control the behaviour of cancer and endothelial cells.

Discussion

We demonstrate that RTKs can be light-activated through substitution of ligand-induced dimerization by light-induced dimerization. In the absence of precedence for light-activated mammalian receptor dimerization, and because of the structural and functional diversity of LOV domains (Moglich et al., 2010; Zoltowski & Gardner, 2011), we initially evaluated seven LOV domains from five different species. This evaluation resulted in the identification of VIAU1-LOV, an algal LOV domain that was never expressed in mammalian cells or incorporated into a chimeric protein. In contrast to phytochromes and LOV domains that were engineered to bind small proteins (Muller & Weber, 2013; Pathak et al., 2013), VIAU1-LOV responds to light with homodimerization (Toyooka et al., 2011). For this reason, a single transgene is sufficient to reconstitute optical activation of cellular signalling, and having to deliver only one transgene greatly facilitates application of the engineered receptors. VIAU1-LOV is capable of activating multiple, distantly related RTKs, as demonstrated here for mFGFR1, hEGFR and hRET. This observation suggests that design principles can be developed that are valid for multiple families of a large receptor superfamiliy and potentially beyond (see below). With the exception of a recently published gene transcription system (Wang et al., 2012), LOV domain homodimerization has not been exploited in optogenetics yet. Our work thus extends the portfolio of useful light-sensing domains with homodimerizing aureochromes and supports the notion that nature offers a large repertoire of light-sensing molecular functionalities that can be harvested by optogenetics (Chow et al., 2010).

For receptor activation, VIAU1-LOV utilizes flavin mononucleotide (FMN) as the light-sensing element. As FMN is present in animal cells, the function of VIAU1-LOV does not require addition of exogenous molecules. Furthermore, low-intensity blue light (~2.5 μW/mm\(^2\)) is sufficient for activation of all Opto-RTKs. This light intensity did not produce phototoxicity, is several orders of magnitude lower than those achieved in standard fluorescence microscopes, and is readily achieved in animal models (Janovjak et al., 2010; Ye et al., 2011). Opto-mFGFR1 enabled the precise control of receptor activation on physiologically relevant time scales and activation of cellular signalling with high spatial precision in genetically homogenous cell layers. Furthermore, activation with light triggered complex cellular programs like DNA synthesis, EMT and angiogenic sprouting in cancer cells and blood endothelial cells, which represent new cellular model systems in which behaviour is under optical control.

Beyond improved spatial and temporal precision, optical activation of RTKs will enable experiments that are difficult to realize using diffusible ligands. Light enables contactless application and complete withdrawal of stimuli without perturbing cellular environments. Furthermore, stimulation by light is limited to the receptor type that was genetically engineered, and thus light offers specificity beyond that of many ligands with promiscuous activation of receptors (Zhang et al., 2006). Work in the nervous system has demonstrated that combining optical activation with genetic targeting provides a powerful approach to manipulate cellular signalling in complex tissues. Recently, light-induced membrane localization of son of sevenless was applied to probe the dynamics of the Ras/ERK pathway (Toettcher et al., 2013). In contrast, light-activation of Opto-RTKs encoded each in a single gene enables control of multiple signalling pathways such as MAPK/ERK, AKT and PLC\gamma1 that all contribute to RTK-mediated biological responses. In the light of the critical involvement of RTKs in developmental biology, stem cell research and regenerative medicine, we expect that Opto-RTKs will open new avenues in these research areas. We also envision their use for cytoprotection and restoration of cell behaviours affected by disorders, particularly in those tissues for which growth factor gene therapy is already considered. Similar design principles may be developed for other cellular signals as receptor dimerization is also implicated in the function of receptor serine/threonine kinases, T-cell receptors and GPCRs (Massague, 1998; Cochran et al., 2001; Lohse, 2010).
Materials and Methods

mFGFR1 receptor constructs

pSH1/M-FGFR1-Fv-Fvls-E [D.M. Spencer, Baylor College of Medicine (Welm et al, 2002)] was obtained from Addgene (Cambridge, MA, USA). The ICD of mFGFR1 flanked by MYR, two FKBP domains and an hemagglutinin (HA)-epitope was transferred from pSH1 to pcDNA3.1(−) (Invitrogen-Life Technologies, Vienna, Austria) using NotI and EcoRI restriction enzymes. Using inverse PCR, a single or both FKBP domains were deleted to yield constructs imFGFR1 and imFGFR1-DFKBP. In this reaction, amplification using
Genes coding for the LOV domains of \textit{A. thaliana} phototropin 1 (AtPH1-LOV2, residue 449–586 of Uniprot entry O48963), \textit{A. thaliana} phototropin 2 (AtPH2-LOV2, residue 363–500 of Uniprot entry P93025), \textit{C. reinhardtii} phototropin (CrPH-LOV1, residue 16–133 of Uniprot entry A8IXU7), \textit{N. crassa} vivid (NcVV-LOV, residue 37–186 with 5Y50 substitution of Uniprot entry Q932Y6), \textit{N. crassa} white collar 1 (NcWC1-LOV, residue 357–586 of Uniprot entry Q01371), \textit{Rhodobacter sphaeroides} PCC 17025 phototropin 2 (AtPH2-LOV2, residue 449–586 of Uniprot entry CSNWS6) were synthesized with mammalian codon optimization according to the supplier’s recommendation (Epoch Life Science, Inc., Missouri City, TX, USA; Supplementary Table S4). NgPA1-LOV and OdPA1-LOV were synthesized with restriction sites. Final receptor constructs were re-inserted into the plasmid GAVPO (Y. Yang, East China University of Science and Technology) containing a Ga4 DNA-binding domain, NcVV-LOV and a trans-activation domain (Wang et al., 2012; Supplementary Fig S4). Activation of GAVPO was detected with the luciferase reporter plasmid of the MAPK/ERK pathway assay (the plasmid contains multiple UAS sequences). VfAU1-LOV was amplified by PCR and oligonucleotides 44 and 45 (Supplementary Table S3) and inserted in GAVPO using BglII and EcoRI restriction enzymes. Luciferase detection was performed as described below except that mFGFR1 plasmids were replaced with 50 ng GAVPO per well.

**Custom incubator for light stimulation of cells**

For light stimulation of cells, an incubator (PT2499; ExoTerra/HAGEN, Holm, Germany) was equipped with 300 light-emitting diodes (JS-FS5050RGB-W30 with JS-CON-004 controller; Komerci, Ebern, Germany; \( \lambda_{\text{max}} \approx 630 \text{ nm (red light)} \) and \( \lambda_{\text{max}} \approx 530 \text{ nm (green light)} \), \( \lambda_{\text{max}} \approx 470 \text{ nm (blue light)} \). Light intensity was controlled with an analogue dimmer and measured with a digital power meter (PM120VA; Thorlabs, Munich, Germany). Intensities at maximal output were 2.3 (red light), 2.6 (green light) and 3.3 (blue light) \text{ W/m}^2.\) For stimulation over extended time periods (> 8 h), an aluminium box was equipped with the same light-emitting diodes and placed in a tissue culture incubator (also see below).

**Cell culture and transfection (HEK293 and CHO K1 cells)**

Cells were maintained in DMEM supplemented with 10% FBS, 100 U/ml penicillin and 0.1 mg/ml streptomycin in a humidified incubator with 5% CO\textsubscript{2} atmosphere. After trypsination, \( 5 \times 10^4 \) cells were seeded in each well of 96-well plates that were coated with poly-L-ornithine (Sigma, Vienna, Austria). Either transparent plates or black clear bottom plates were used. Cells were transfected using Lipofectamine 2000 (Invitrogen/Life Technologies) or polyethyleneimine (Polysciences, Eppelheim, Germany).

**LOV domain expression and cell viability (HEK293 and CHO K1 cells)**

An expression plasmid based on pcDNA3.1(−) was prepared in which a BspEI restriction site followed the fluorescent protein mVenus (Nagai et al., 2002) and a glycine- and serine-rich linker. LOV domains were inserted into this plasmid using PCR (see above). Cells were transfected with 100 ng expression plasmid in each well of 96-well plates (three to four wells were prepared for each construct throughout the study). Expression was assessed by...
measuring mVenus fluorescence in a plate reader (BioTek Synergy H1, Bad Friedrichshall, Germany) 16–18 h after transfection. Transfection of pcDNA3.1(−) or a FKB-B-mVenus fusion protein served as controls. Viability measurements used a tetrazolium dye following the supplier’s protocol (EZ4U Cell Proliferation and Cytotoxicity Assay; Biomedica, Vienna, Austria). Absorbance measurements at 450 nm with 620 nm reference were taken after fluorescence measurements.

**Stimulation and detection of MAPK/ERK pathway (HEK293 cells)**

Activation of the MAPK/ERK pathway was assayed with the PathDetect Elk1 trans-Reporting System (Agilent, Vienna, Austria) consisting of an Elk1 phosphorylation-dependent trans-activator and a luciferase-based trans-reporter. Cells were transfected with 213.3 ng total DNA per well (receptor, trans-activator and trans-reporter at a ratio of 1:3:60 or 1:3:600). Six hours after transfection, medium was replaced with CO₂-independent reduced serum starve medium (Gibco/Life Technologies; supplemented with 0.5% FBS, 2 mM L-glutamine, 100 U/ml penicillin and 0.1 mg/ml streptomycin) and cells were incubated for 18 h at 37°C. Cells were transferred to the stimulation incubator and for 8 h either subject to constant illumination or, for control cells throughout the study, kept in the dark. For stimulation of imFGFR1, 10 nM AP20187 (ARIAD Pharmaceuticals, Cambridge, MA, USA; Clackson, 1998) was added before transfer to the stimulation incubator. After incubation, plates were washed with PBS and processed with Luciferase 1000 WesternL (Promega, Glostrup, Denmark) in PBS for 10 min at 37°C. Luminescence was detected with a microplate reader (Tecan Infinite 200 Pro, Männedorf, Switzerland). Some samples were processed with ONE-Glo (Promega) and detected with the BioTek microplate reader. Differences in absolute intensities due to assay and plate reader were corrected using Opto-mFGFR1 as a standard if intensities were plotted in the same figure. UO126 was obtained from LC Laboratories (Woburn, MA, USA).

**Stimulation and immunoblot (HEK293 cells)**

2 × 10⁶ cells were seeded in 25-cm² dishes and transfected with 2 μg Opto-mFGFR1. After 5 h, medium was replaced with reduced serum medium. After additional 20 h, cells were illuminated for 5 min. Cells were harvested in 70 μl lysis buffer on ice and lysates were sonicated and centrifuged (12,000 g, 5 min, 4°C). Thirty μg protein per lane were separated by SDS-PAGE and electro-blotted onto PVDF membranes. Blots were incubated with primary antibodies (ERK1/2, #9102; pERK1/2, #9101; PLCγ1 #2822; pPLCγ1 #2821; AKT #9272; pAKT #4058S; pFRS2α(Tyr196), #3864; pFRS2α(Tyr436) #3861; Cell Signaling Technology, Danvers, MA, USA dilution 1:1,000; FRS2 (A-5) Santa Cruz Biotechnology, Santa Cruz, CA, USA, dilution 1:500; β-actin, Sigma, dilution 1:8,000) in blocking solution (3% BSA or 5% skim milk in TBST) overnight at 4°C. Secondary antibodies (HRP-coupled α-rabbit or α-mouse IgG; Dako, Glostrup, Denmark) were applied at a dilution of 1:10,000 for 2 h at RT. Chemiluminescence was developed with WesternC reagent (Biorad, Hercules, CA, USA) and signals recorded on X-ray film (GE Healthcare, Chalfont St. Giles, UK). For FRS2α-RFP transfection, an expression plasmid based on pcDNA3.1(−) was prepared in which a Nhel restriction site precedes the red fluorescent protein and human FRS2α was inserted into this plasmid using PCR (oligonucleotides 46–47, Supplementary Table S3). For detection of transactivation, stable transfectants were selected with 500 μg/ml G418.

**Detection of additional signalling pathways (HEK293 cells)**

Activation of additional mFGFR1-related signalling pathways was assayed with Cignal Reporter assays (Qiagen, Hilden, Germany) consisting of mixtures of inducible pathway-luciferase reporters. Cells were transfected with 100.3 ng total DNA per well (receptor and reporter at a ratio of 1:300) and treated as described above. Cells were processed with the Dual-Glo Luciferase Assay System (Promega) and Tecan microplate reader.

**Live-cell fluorescence microscopy (HEK293 cells)**

An expression plasmid based on pcDNA3.1(−) was prepared in which a BspE1 restriction site preceded the fluorescent protein mVenus and an in-frame glycine- and serine-rich linker. Human PLCγ1 was inserted into this plasmid using PCR and the Agel restriction enzyme (oligonucleotides 48 and 49, Supplementary Table S3). 4 × 10⁶ HEK293 cells were seeded in 5-cm dishes that were coated with poly-L-lysine and transfected with 200 ng expression plasmid per well (receptor and PLCγ1-mVenus at a ratio of 3:4). Medium was replaced with reduced serum medium 24 h before imaging. Stimulation with LEDs (see above) and live-cell microscopy was performed a Leica SP5 confocal microscope.

**Live-cell luminescence in spatially confined illumination experiments (HEK293 cells)**

3 × 10⁶ HEK293 cells were simultaneously seeded and transfected with 24 μg expression plasmid (receptor, trans-activator and trans-reporter at a ratio of 1:3:60) in a 10-cm dish. After 16 h, cells were treated and illuminated as described above. Live cells were incubated with 0.15 mg/ml D-luciferin (PEQLab, Erlangen, Germany) in PBS for 10 min at 37°C. Luminescence was detected with a PEQLab Fusion SL imaging system (PEQLab, Erlangen, Germany).

**Generation of stable Opto-mFGFR1 transgenic cell lines and virus construction (M38K, SPC212 and hBE cells)**

M38K and SPC212, two malignant pleural mesothelioma cell lines, were maintained in RPMI1640 supplemented with 10% FBS. Telomerase-immortalized microvascular hBE cells were maintained in Clonetics EGM2 MV endothelial growth medium (Lonza, Wakersville, MD) supplemented with 5% FBS. For retrovirus generation, Opto-mFGFR1 or mCherry (Shaner et al, 2004) were subcloned into pQCXIP (Clontech, Mountain View, CA, USA) using NotI and EcoRI restriction enzymes. Viral particles were generated in HEK293 cells by co-transfection with the helper plasmids pSVSV-G (Clontech) and p-gag-pol-gpt (Markowitz et al, 1988). Supernatants were used to transduce M38K, SPC212 or hBE cells grown to 50% confluency in 6-well plates. Cells were selected with 0.8 μg/ml puromycin for 10 days, and transgene expression was verified by immunoblotting.
Stimulation and immunoblot (M38K, SPC212 and hBE cells)

For immunoblotting, 5 × 10^5 cells were seeded in each well of 6-well plates. After 24 h, medium was replaced with reduced serum medium (M38K and SPC212 cells). After additional 20 h, cells were transferred to the stimulation incubator and illuminated for 1, 5 or 15 min. Cells were then either immediately or after additional 5, 15 or 30 min in the dark washed and lysed in 50 μl lysis buffer per well on ice. Lysates were sonicated and centrifuged as described above. Fifteen μg protein per lane were separated by SDS-PAGE and electroblotted onto PVDF membranes. Blots were incubated with primary antibodies (FGFR1, #9740 Cell Signaling Technology, dilution 1:1,000; FGFRpY653/654, Thermo Scientific, Vienna, Austria, dilution 1:1,000) and secondary antibodies for processing as described above.

Immunofluorescence and confocal microscopy (HEK293, M38K, SPC212 and hBE cells)

For localization of Opto-mFGFR1, Opto-hEGFR and Opto-hRET in HEK293 cells, 3 × 10^5 cells per well were seeded into each well of 6-well plates and transfected with 2 μg expression plasmid. After 24 h, 1 × 10^5 cells were seeded into fibronectin-coated chamber slides (BD Falcon, Heidelberg, Germany). After additional 24 h, cells were fixed with 3.8% formaldehyde, blocked with 1% BSA in PBS and incubated with monoclonal rat α-HA (3F10, Roche, Mannheim, Germany, 1:100) overnight at 4°C. This was followed by Alexa Fluor 546-linked goat α-rat antibody for 1 h at RT. Slides were mounted in Vectashield mounting medium containing DAPI and images recorded on a Zeiss LSM700 confocal microscope. For localization of Opto-mFGFR1 in stable cell lines, 5 × 10^5 M38KOpto-mFGFR1, SPC212Opto-mFGFR1 and hBEOpto-mFGFR1 cells were seeded into chamber slides and stained and imaged as above.

Opto-mFGFR1 cross-linking (M38K cells)

For chemical cross-linking, 10^5 cells were collected and washed three times with PBS. Cells were stimulated for 15 min with blue light. Disuccinimidyl suberate was added to a final concentration of 5 mM and incubated at RT for 30 min in blue light or dark. The reaction was quenched with 10 mM Tris, pH 7.5 for 15 min with blue light. Cell lysis and immunoblotting were performed as described above.

ERK1/2 phosphorylation in spatially confined illumination experiments (SPC212 and hBE cells)

For detection of localized ERK1/2 phosphorylation in cell monolayers, SPC212 or hBE cells were grown to confluency in 6-cm petri dishes (SPC212) or 12-well plates (hBE cells). SPC212 were starved in medium without serum for 24 h before illumination. Templates with pinholes of 2 (SPC212) or 5 (hBE cells) mm diameter were used for localized illumination for 5 min. Afterwards, cells were washed with cold PBS and fixed with Histofix (Lactan, Graz, Austria) for 10 min. After washing with PBS and permeabilization with Triton X-100 (0.25% in PBST) and blocking in 1% BSA in PBST, dishes were incubated with pERK1/2 (#9101; Cell Signaling Technology, 1:500 for SPC212, 1:100 for hBE cells) for 1 h. Signal was developed using the UltraVision LP detection system (Thermo Scientific) and 3,3′-diaminobenzidine as chromogen. Haematoxylin was used for counterstaining of cell nuclei.

Cell proliferation (M38K cells)

2 × 10^4 cells were seeded in each well of 96-well plates. After 24 h, cells were stimulated for 1 h. FGFR2 (Sigma, St. Louis, MO) was added to a final concentration of 10 ng/ml. After 24 h, cells were incubated with 10 μM EdU for 2 h. Subsequently, newly synthesized DNA was stained with Click-it EdU (Life Technologies) following the manufacturer’s protocol and counterstained with 5 μg/ml Hoechst dye. Cells were photographed on a Nikon Ti300 inverted microscope. To determine the percentage of cells with newly synthesized DNA, Hoechst positive nuclei and EdU positive nuclei were counted.

Cell morphology (M38K cells)

10^5 cells were seeded in each well of 6-well plates. After 24 h, cells were stimulated for 1 h. FGFR2 (Sigma) or PD168666 (Pfizer Global Research and Development, New London, CT) were added to a final concentration of 10 ng/ml or 10 μM, respectively. After additional 24 h, cells were photographed on the Nikon Ti300 microscope. For quantification of cell morphology, all cell perimeters in randomly selected sections of phase contrast images were traced and aspect ratios (defined as length of major axis divided by length of minor axis of a fitted ellipse) calculated with ImageJ software (National Institute of Health). Sixty individual values contributed to each average. Automated analysis yielded comparable results.

Cell cycle (M38K cells)

Cell cycle distribution was analysed by flow cytometry. 5 × 10^5 cells were seeded in 25-cm² tissue culture flasks. After 24 h, cells were stimulated for 1 h. After additional 24 h, cells were fixed in ethanol (70%), treated with 50 μg/ml RNase A and 50 μg/ml propidium iodide (PI). Flow cytometry was performed on a FACSCalibur (BD Biosciences, Schwechat, Austria) and cell cycle distribution calculated with ModFit LT software (Verity Software House, Topsham, ME).

Actin staining (M38K cells)

5 × 10^5 cells were seeded onto coverslips in 6-well plates. After 24 h, cells were illuminated with a cycle of 5 min blue light/15 min dark for 48 h. Cells were fixed (3.8% formaldehyde), permeabilized (0.5% Triton X-100 in PBS), stained with TRITC-phalloidin (70%), treated with 50 μg/ml RNase A and 50 μg/ml propidium iodide (PI). Flow cytometry was performed on a FACSCalibur (BD Biosciences, Schwechat, Austria) and cell cycle distribution calculated with ModFit LT software (Verity Software House, Topsham, ME).

Gene expression analysis (M38K, SPC212 and hBE cells)

Total RNA was extracted with TRIzol (Life Technologies) from sub-confluent flasks of M38KOpto-mFGFR1, SPC212Opto-mFGFR1 and hBEOpto-mFGFR1 cells and the respective controls. RNA was reverse transcribed with MMLV reverse transcriptase (Thermo Scientific). cDNAs corresponding to 50 ng RNA per sample were used for SYBR
Supplementary information for this article is available online: http://emboj.embopress.org

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Conflict of interest
The authors declare that they have no conflict of interest.

References

heparan sulfate 6-O-sulfation levels regulate angiogenic responses of endothelial cells to fibroblast growth factor 2 and vascular endothelial growth factor. J Biol Chem 287: 36132 – 36146