Optogenetic delivery of trophic signals
in a genetic model of Parkinson’s disease

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Abstract

Optogenetics has been harnessed to shed new mechanistic light on current therapies and to develop future treatment strategies. This has been to date achieved by the correction of electrical signals in neuronal cells and neural circuits that are affected by disease. In contrast, the optogenetic delivery of trophic biochemical signals, which support cell survival and thereby may modify progression of degenerative disorders, has never been demonstrated in an animal disease model. Here, we reengineered the human and Drosophila melanogaster REarranged during Transfection (hRET and dRET) receptors to be activated by light, creating one-component optogenetic tools termed Opto-hRET and Opto-dRET. Upon blue light stimulation, these receptors robustly induced the MAPK/ERK proliferative signaling pathway in cultured cells. In PINK1B9 flies that exhibit loss of PTEN-induced putative kinase 1 (PINK1), a kinase associated with familial Parkinson’s disease (PD), light activation of Opto-dRET suppressed mitochondrial defects, tissue degeneration and behavioral deficits. In human cells with PINK1 loss-of-function, mitochondrial fragmentation was rescued using Opto-dRET via the PI3K/NF-κB pathway. Our results demonstrate that a light-activated receptor can ameliorate disease hallmarks in a genetic model of PD. The optogenetic delivery of trophic signals is cell type-specific and reversible and thus has the potential to overcome limitations of current strategies towards a spatio-temporal regulation of tissue repair.
Significance Statement

The death of physiologically important cell populations underlies a wide range of degenerative disorders, including Parkinson’s disease (PD). Two major strategies to counter cell degeneration, soluble growth factor injection and growth factor gene therapy, can lead to the undesired activation of bystander cells and non-natural permanent signaling responses. Here, we employed optogenetics to deliver cell type-specific pro-survival signals in a genetic model of PD. In *Drosophila* and human cells exhibiting loss of the PINK1 kinase, akin to autosomal recessive PD, we efficiently suppressed disease phenotypes using a light-activated tyrosine kinase receptor. This work demonstrates a spatio-temporally precise strategy to interfere with degeneration and may open new avenues towards tissue repair in disease models.
Introduction

Biology occurs over a wide range of time and length scales, from milliseconds and nanometers for protein folding, to days and centimeters for organism development. In recent years, powerful research methods have been developed that permit the manipulation of biological processes on even the smallest length and shortest time scales. In optogenetics, natural or reengineered photoreceptors are expressed in genetically defined cell populations to optically activate or inhibit, e.g., neuronal action potential firing or cell signaling. The use of light provides unprecedented precision in space and time as a way to answer previously unresolvable questions in a multitude of disciplines, including microbiology, cell/developmental biology, synthetic biology, and neuroscience. In particular, spatio-temporally precise perturbation of selected cells in intact organisms can reveal cause-consequence-relationships that are a critical determination for understanding central nervous system function or animal development (1-3). Optogenetics also provides access to the reversible and rapid activation of cell signaling pathways that is required for dissection of their dynamic properties (4, 5) and for development of new drug discovery platforms (6). Inspired by these successes, optogenetics is continuously translated into new research areas, including disease mechanism and therapy.

Shortly after its inception, optogenetics was beginning to be employed in the study of neural circuits that are known to be affected by neurological and neurodegenerative disorders, including spinal cord injury, stroke and Parkinson’s disease (PD) (7, 8). In this field, optogenetics has shed new light on the mechanisms of currently utilized therapies (e.g., deep brain stimulation in PD) or therapies of the future (e.g., stem cell-based tissue regeneration) (9, 10). This work was followed by the development of light-gated prosthetic approaches in which a genetically introduced photoreceptor senses either natural light, e.g. for vision restoration (11), or light from a prosthetic source, e.g. for heart or skeletal muscle pacing (12, 13). Notably, these pioneering studies harnessed optogenetics to excite or inhibit electrical signals through regulated ion flow ions across the cell membrane. In apparent contrast, the optogenetic delivery of trophic signals, which support cell survival and are
central to treatment strategies in a variety of degenerative disorders, has never been
demonstrated in a disease model. It is unclear if this is feasible as hypo- or hyperactivity of
pro-survival pathways is linked to undesired cellular outcomes (see below).

We and others have recently engineered light-activated variants of key signaling
proteins that now provide a basis for the optogenetic delivery of trophic signals. Particular
success was reported for receptor tyrosine kinases (RTKs) (14-18). RTKs are expressed in
virtually all human cell types and respond to growth factors (GFs) with conformational
changes and/or oligomerization state changes that result in receptor trans-phosphorylation.
Trans-phosphorylation is then followed by recruitment of intracellular secondary messengers
in, e.g., the mitogen-activated protein kinase/extracellular signal-regulated kinase
(MAPK/ERK) or phosphatidylinositol-3 kinase/AKT (PI3K/AKT) signaling pathways. Because
of their ability to activate these proliferative and pro-survival pathways, RTKs are prime
targets in several neurodegenerative disorders. In the context of PD, the RET RTK (19) has
been intensively investigated in both preclinical and clinical studies. hRET is activated by
glial cell line-derived neurotrophic factor (GDNF) family ligands (GFLs; these are GDNF,
neurturin, artemin, and persephin) that bind GDNF family receptor α (GFRα) co-receptors
(GFRα1-4) to recruit dimeric RET into a ternary complex (Figure 1A). GFLs are linked to the
development and maintenance of dopaminergic (DA) midbrain neurons and have been
pursued as disease-modifying agents in PD, either by local injection or by gene delivery
using adeno-associated viruses (20, 21). Despite initial success in animal models, outcomes
in clinical trials were limited (22, 23), which was attributed to difficulties in GFL delivery,
limited responsiveness of the targeted DA neurons and advanced PD in some of the
recruited patients. In addition, there are concerns that the continuous delivery of GFLs can
lead to counter-productive compensatory effects (24-27). These observations and
considerations have highlighted a need for methods that can control the GFL-RET-axis in a
reversible and more precise manner (28, 29).

Here, we explored optogenetics as a means for delivery of trophic signals in a genetic
model of PD. We first reengineered full-length hRET and its Drosophila orthologue dRET.
(30-32) to be activated by light in optogenetic tools termed Opto-hRET and Opto-dRET. We then showed that temporally precise dRET activation in vivo can be used to induce degeneration in a tissue sensitive to ectopic RTK signaling. Optogenetic delivery of RET signals was then successfully applied in a genetic fly model of PD. Mutations in the PINK1 gene are linked to autosomal recessive PD (33-35), and Drosophila has been shown to be a suitable model to study consequences of PINK1 loss-of-function (36-38). We suppressed Drosophila phenotypes associated with PINK1 deficiency and identified the involved downstream signaling pathways in a human cellular model. This work demonstrates the use of optogenetics as a cell-type specific and remote controlled method to exert beneficial trophic effects of in a genetic disease model.

Results

Light-activated hRET and dRET receptors

hRET assembles in dimers in the activated ternary GFL2-GFRα2-RET2 complex (39, 40) (Figure 1A) and forced dimerization by mutations or synthetic binding domains has been shown to induce signaling of hRET (41, 42) and dRET (43, 44). Based on these observations, we converted hRET and dRET into optogenetic tools by incorporating a light-activated dimerization switch. To achieve this switch, we utilized the light-oxygen-voltage-sensing (LOV) domain of the AUREOCHROME1 photoreceptor from the yellow-green algae Vaucheria frigida (AU1-LOV) (45) (Figure 1B). AU1-LOV is a member of the large LOV domain superfamily and responds to blue light with formation of a symmetric homodimer (46) (Figure S1). AU1-LOV is smaller than other photoreceptors commonly used in optogenetics (145 aa in length; this corresponds to ~a third of the length of cryptochromes or phytochromes) (47, 48) and relaxes slower than many other LOV domains from the light-activated ‘lit’ state (that is predominantly dimeric) to the dark-adapted state (that is predominantly monomeric; relaxation time constant ~600 s) (49). We and others have shown that small size and slow cycling make AU1-LOV well suited for assembly and activation of membrane receptors (14, 16, 17, 50). We placed AU1-LOV at the far C-terminus of the RET
receptors because fluorescent proteins (FPs) were previously incorporated at this site without negative impact on receptor signaling or trafficking (51, 52). To functionally test the generated Opto-hRET and Opto-dRET, we took advantage of the fact that Drosophila RTKs can couple to the mammalian MAPK/ERK pathway via Ras (44). We and others utilize human embryonic kidney 293 (HEK293) cells for testing new optogenetic methods because these cells do not exhibit native light-induced signaling events. Using transcriptional reporters (14, 47), we found robust induction of MAPK/ERK signaling upon blue light stimulation of HEK293 cells transfected with Opto-hRET and Opto-dRET (intensity (I) = 250 μW/cm², wavelength (λ) = 470 nm) (Figure 1C). Whilst Opto-hRET activated transcriptional responses more strongly than Opto-dRET, Opto-dRET activation levels were comparable to those reached by the Opto-dRET\textsuperscript{MEN2B} variant (Figure 1C). Opto-dRET\textsuperscript{MEN2B} contains a Met to Thr gain-of-function substitution in the kinase domain that was discovered in multiple endocrine neoplasia (MEN) Type 2B as causative for potent receptor hyperactivation in the absence of GFLs (43, 53). These results show that signaling activity can be induced by blue light through Opto-h/dRET receptors.

Opto-dRET function in vivo

We next tested if Opto-dRET can be applied in vivo to conduct a temporally-targeted gain-of-function experiment (Figure 2A). We choose the Drosophila retina for this experiment because RTKs and their downstream pathways are tightly regulated during its morphogenesis, and because RTK hyperactivation during retina development results in marked phenotypes. For instance, two RTKs, the Drosophila epidermal growth factor receptor (DER) and Sevenless, orchestrate retinal cell growth, differentiation and regulated death (54, 55). These RTKs act in late larval and early pupal stages to form the fourteen cells that compose each ommatidium unit eye and the ommatidial lattice (56). Hyperactivation of RTK signaling during these stages has been shown to result in irregular ommatidia size and spacing leading to a disrupted tissue pattern termed ‘roughening’ (57). We generated transgenic flies in which the Opto-dRET gene is placed downstream of five UAS elements.
(Figure S2). We also generate analogous flies expressing the constitutively-active Opto-
dRET\textsuperscript{MEN2B}. We then targeted Opto-dRET or Opto-dRET\textsuperscript{MEN2B} to the retina using the GMR-
GAL4 driver, which induces expression in cells located posterior of a morphogenetic furrow
that sweeps in anterior direction to initiate mitosis and cell differentiation (55). In Opto-
dRET\textsuperscript{MEN2B} flies, scanning electron microscopy (SEM) revealed a marked rough retina
phenotype (compare Figure 2B and C). Roughening was previously observed in flies
expressing dRETMEN2B (43), and based on the severe outcome observed for Opto-dRET\textsuperscript{MEN2B}
we concluded that AU1-LOV attachment does not negatively impact receptor function. We
next illuminated control flies and Opto-dRET flies in a time window that captures ommatidia
and lattice formation (from third instar larva through to the second day after pupariation; I =
385 \mu W/cm\textsuperscript{2}, \lambda = 470 nm; Figure 2A). In control flies without Opto-dRET, we did not observe
light-induced roughening indicating that light alone did not have an effect on the retina
(compare Figure 2D and E; in agreement with previous studies, we observed mild
phenotypes upon GAL4 expression with the GMR driver (58)). In apparent contrast, we found
that light stimulation resulted in a marked increase in roughening in Opto-dRET flies
(compare Figure 2F and G). To quantify this effect, we manually metered in each retina
image the ‘fused area’ (the area without identifiable ommatidia) and also applied
computational methods to count individual ommatidia (~600 ommatidia can be assigned in
our frontal view images) as two measures of tissue integrity. We found that upon illumination
the fused area increased and the number of identified structures decreased specifically in the
illuminated Opto-dRET flies (Figure 2H and I). For these and control flies, we also
determined lattice regularity, which is defined as the ratio of the mean and the standard
deviation (SD) of the ommatidia nearest-neighbor distance (NND) distribution (59). Regularity
decreased from 3.98 $\pm$ 0.39 in WT flies to 2.15 $\pm$ 0.55 in illuminated Opto-dRET flies, and
these values are indicative of near-perfect regularity and near-random assembly,
respectively (60). The potent effects induced by Opto-dRET upon light stimulation and the
lack of light responses in the absence of Opto-dRET establish the suitability of this
optogenetic approach to modify tissue behavior in vivo.
Suppression of defects in a genetic model of PD

With Opto-dRET in hand, we went on to explore if defects in a genetic disease model can be ameliorated using optogenetics (Figure 3A). PINK1 is a Ser/Thr kinase that localizes to mitochondria and supports their integrity and function. Loss-of-function mutations and dominant negative mutations in the PINK1 gene are associated with autosomal recessive PD (33-35). In Drosophila, loss of X-linked PINK1 leads to a striking phenotype, including tissue degeneration, locomotor defects and disruption of mitochondrial structure and function (61-64). To test if optogenetics can suppress phenotypes associated with PINK1 deficiency, we expressed Opto-dRET in indirect flight muscles (IFMs) of PINK1 B9 flies using the MEF2-GAL4 driver (65). IFMs are frequently studied in Drosophila models of PD and PINK1 loss-of-function leads to a marked ‘crushed’ thorax phenotype and reduced locomotion. We first compared PINK1 B9 flies to Opto-dRET PINK1 B9 flies that were not illuminated. Similar penetrance of thoracic defects (58 and 61% of flies exhibited a crushed thorax, respectively) shows that the engineered optogenetic receptor did not affect the phenotype in the absence of light (Figure 3B). When proceeding to light stimulation, we took into consideration that the opaque case and cuticle of pupa and adults may reduce blue light penetration to IFMs. To address this, we first confirmed that AU1-LOV can be activated by light of 1-5 \( \mu \text{W/cm}^2 \) intensity, which corresponds to the product of minimal blue light transmission through the case or cuticle (\( \sim 0.5\% \) (66, 67)) and the light intensity applied in our light chambers (\( I = 320 \mu \text{W/cm}^2 \); Figure S3A). We observed that light of this intensity is indeed sufficient to activate AU1-LOV (Figure S3B). We then went on to light stimulate Opto-dRET PINK1 B9 flies during late pupal stages and adult states (Figure 3A) (these stages coincide with the onset of degeneration (64, 68)). Strikingly, we observed phenotype suppression in Opto-dRET PINK1 B9 flies resulting in only 16% of flies with defects (Figure 3B). This result indicates marked improvement in tissue integrity and was comparable to the improvement observed previously upon PINK1 overexpression in the PINK1 B9 model (Park et al, 2006). We also tested if illumination restored the climbing deficits that accompany PINK1 loss-of-function.
This was indeed the case with illuminated Opto-dRET flies reaching climbing pass rates similar to those of WT flies (Figure 3C).

Mitochondrial dysfunction is a major pathological alteration observed in sporadic and familial PD and also the primary cellular consequence of loss of PINK1. We therefore tested the effect of illumination on mitochondrial function and integrity in Opto-dRET PINK1B9 flies. PINK1B9 flies exhibited reduced muscle ATP levels and these levels could be restored by Opto-dRET and light stimulation (Figure 4A). To examine mitochondrial integrity, we conducted ultrastructure analysis using transmission electron microscopy (TEM). PINK1B9 muscles exhibited broadening of the myofibril Z-line and enlarged mitochondria with fragmented cristae (compare Figure 4B and C). Illumination of Opto-dRET PINK1B9 flies was clearly beneficial with a reduced fraction of impaired mitochondria and an increased fraction of mitochondria with WT-like cristae structure (compare Figure 4D and E) that approached levels of control flies (Figure 4F). Overall, these results on the cell- and tissue-level demonstrate optogenetic suppression of consequences of PINK1 loss-of-function in a Drosophila model. In these experiments, we took advantage of temporally precise light stimulation to prevent undesired side effects of continuous growth signal delivery, specifically lethality associated with dRET overactivation in muscle at early developmental stages (69).

Amelioration of mitochondrial defects in PINK1-deficient human cells

Finally, we tested if light activation of RET signaling can revert defects induced by loss of PINK1 in human cells. We performed these experiments in dopaminergic neuroblastoma SH-SY5Y cells that have been previously applied to study how mutations observed in PD, including those in the PINK1 gene (70), impact mitochondrial integrity. We transfected the cells with either control siRNA or PINK1 siRNA as well as expression vectors for Opto-dRET, Opto-dRETMEN2B or an inactive ‘kinase-dead’ (KD) Opto-dRET (Opto-dRETKD). Western blot (WB) analysis revealed efficient downregulation of PINK1 levels (Figure 5A) and that expression levels of the Opto-dRET variants were comparable (Figure 5B). Silencing of the PINK1 gene resulted in severe mitochondrial defects with ~65% of cells exhibiting
fragmented mitochondria (Figure 5C, rows 1 and 2, Figure 5D, bars 1 and 2). As shown previously, mitochondrial integrity was restored in this model through endogenous RET stimulated with GDNF/GFRα1 for 4 h (69, 71). In this paradigm, the fraction of cells with fragmented mitochondria was reduced to 20%, which is comparable to cells treated with control siRNA (Figure 5C, rows 1 and 3, Figure 5D, bars 1 and 3). We then analyzed Opto-dRET\textsuperscript{MEN2B} and Opto-dRET cells and found that either expression of Opto-dRET\textsuperscript{MEN2B} or light stimulation of Opto-dRET cells (I = 232 μW/cm², λ = 470 nm, 4 h) rescued mitochondria with similar efficiency (~25% of cells displaying fragmentation; Figure 5C, rows 4 to 6, Figure 5D, bars 4 to 6). Similarly to the Drosophila experiments, no rescue was observed upon Opto-dRET expression in dark conditions, indicating limited basal receptor activity in the absence of the light stimulus (Figure 5C, rows 2 and 5, Figure 5D, bars 2 and 5). We also verified that the kinase activity of dRET is required for rescue (Figure 5C, rows 5 to 8, Figure 5D, bars 5 to 8) and that light alone did not influence mitochondrial morphology (Figure S4). We also tested which signaling pathways downstream of RET are involved in mediating mitochondrial integrity. Of the main pathways activated by RET, we found that reversion of mitochondrial fragmentation depended on both the PI3K and nuclear factor 'kappa-light-chain-enhancer' of activated B-cells (NF-κB) pathway, but not on the MAPK/ERK pathway (Figure 5E). This result is in agreement with previous studies showing that the protein network regulated by GFL/RET overlaps with that involved in PINK1 function (69, 71). Collectively, these findings show that beneficial trophic signals can be delivered to a human cellular model of PD using optogenetics.

Discussion

Choreographed signaling of GFs and their cognate RTKs underlies tissue morphogenesis and homeostasis, whereas their aberrant activity is linked to human disorders. For instance, in the case of RET, gain-of-function is implicated in several forms of cancer and loss-of-function is linked to developmental disorders and neurodegeneration (72, 73). Motivated by the importance of RET, we engineered human and Drosophila RET receptors that can be
activated by light. Recent work has demonstrated light activation of RTKs generally following seminal designs that built on either dimerizing (14) or oligomerizing (15) photoreceptor domains. We developed Opto-hRET and Opto-dRET using the homodimerizing AU1-LOV domain, and whilst LOV domains have enabled dimerization of isolated kinase domains in the past, we here show that this approach is suited for activation of full-length RET receptors. This suggests that enforced association at the RET C-terminus can overcome autoinhibition by elements of the extracellular domain that can counteract ligand-independent dimerization. These light-activated human and Drosophila RTKs add to an optogenetic arsenal that already consists of light-activated enzymes and optically-recruited signaling proteins, some of which have already permitted the optogenetic control of cell behavior in Drosophila (1, 2).

In the first experiment, we applied Opto-dRET in the Drosophila retina to interfere with tissue morphogenesis. Retina development depends on concerted cell proliferation and differentiation events, and RTKs play a key role in ommatidia formation and ommatidial lattice generation. We observed retina malformations specifically in flies that were illuminated during tissue formation stages, in agreement with earlier observations that downstream pathways are not operating at maximal levels during retina development because of multiple and reiterative uses of RTKs (54, 55, 74). This experiment complements recent optogenetic studies in Drosophila tissues other than the retina that have incorporated spatio-temporal regulation to identify tissues and stages with high sensitivity to ectopic signals (75-78).

We then explored if optogenetics can suppress phenotypes in a genetic model of PD. Previous optogenetic studies in the context of neurological and neurodegenerative disorders focused on understanding or correcting aberrant electrical activity in excitable cells (7, 8), whilst our goal was the delivery of trophic effects through the optical control of biochemical pro-survival pathways. Our model was Drosophila with loss of PINK1, a Ser/Thr kinase that causes autosomal recessive PD (33-35). Although evidently not able to recapitulate all features of human PD, we chose Drosophila as the model because PINK1 loss-of-function manifests in robust phenotypes that have previously helped to delineate pathways implicated in mitochondrial physiology and in PD pathogenesis (61, 64, 68, 79-86). Cell degeneration in
this model occurs most strongly in cells outside of the nervous system, such as in IFMs, likely because of their high energy demand. Activation of Opto-dRET resulted in efficient suppression of mitochondrial alterations, tissue degeneration and attendant locomotion fitness. We also demonstrated rescue of mitochondrial morphology in PINK1-deficient human cells, and this second model allowed us to identify signaling pathways downstream of dRET that are essential for reversion of the defects. PI3K and NF-κB activity were required to reestablish the healthy mitochondrial network. These pathways are known to act as an important node of crosstalk downstream of tyrosine kinases (87-89), and their involvement is in line with previous observations that the protein networks regulated by GFL/RET overlap with those altered in PD (69, 71). We noted that PINK1 deficiency phenotypes in flies and human cells were only modified by Opto-dRET upon stimulation with light but not in the dark, indicating little background activity of the receptor in the absence of activation of the introduced optical switch. It will be interesting to determine in future studies whether Opto-dRET is efficacious in ameliorating phenotypes in mammalian in vivo models of PD.

The new ability to remotely and spatio-temporally control cellular events relevant to human disease has previously inspired the pursuit of optogenetics-based treatment strategies (see above); but what makes optogenetics an attractive method for pro-survival signal delivery, in general or in the context of PD models? The protection or regeneration of cells is a key target in the treatment of a variety of disorders, but the practical application of GFs is challenging. Many GF receptors have widespread tissue distribution and thus systemic growth factor administration may result in off-target effects, such as toxicity or undesired proliferation, in cells other than those targeted (90, 91). Furthermore, many GFs exhibit limited half-life in the circulation or do not reach target tissues (92, 93). Additionally, GF gene therapy results in permanent hyperactivation of signaling pathways that can be linked to side effects and potential safety issues (94). In PD specifically it is not clear if the cellular signaling machinery of degenerating DA neurons can respond to GFLs, e.g. because of impaired RTK retrograde trafficking or expression (23, 95). Optogenetics has properties that may address some of these challenges. For instance, optogenetic control can be
reversible and limited to specific cell populations. In addition, optogenetic receptors do not rely on ligand binding in neuronal projections. It has been recently demonstrated that RET downregulation in a mouse model of PD can be compensated by a virally delivered of RET (96). This finding provides an encouraging basis for further exploration of Opto-RETs in mammalian models of PD. Translation of optogenetics into the brain may be facilitated by innovations that are currently pursued by many groups, such as wirelessly-powered microscale electronics that are implantable and biocompatible, or transcranial energy delivery. In this study, we demonstrated in a genetic model of PD that ligand-independent optical delivery of trophic signals is in principle possible, paving the way for future studies in animal models of PD and potentially also other disorders linked to the GF-RTK axis.
Author contributions (CRediT taxonomy)

Conceptualization, A.I.P., D.S., P.S., K.W. and H.J.; Funding Acquisition, D.S., P.S., K.W. and H.J.; Methodology, A.I.P., P.S., K.W. and H.J.; Project Administration, H.J.; Investigation, A.I.P., N.F., N.H., M.P., E.R. and V.Z.; Data curation, A.I.P., N.F., S.C., N.H., M.P.; Resources, A.G. and J.B.; Supervision, D.S., P.S., K.W. and H.J.; Writing - Original Draft, A.I.P. and H.J.; Writing - Review and Editing, D.S., P.S., K.W. and H.J.

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Materials and Methods

Engineering light-activated RET receptors. The gene encoding full-length dRET with the MEN2B M955T substitution \((dRETMEN2B)\), a kind gift of Ross Cagan, Icahn School of Medicine at Mount Sinai, NY) was amplified from an expression vector by PCR and inserted into pUAST. To obtain \(Opto-dRETMEN2B\) in pUAST, \(AU1-LOV\) (14) was inserted at the far C-terminus of the receptor. To obtain \(Opto-dRET\) in pUAST, the M955T substitution of \(dRETMEN2B\) was reverted using site-directed mutagenesis. To express \(Opto-dRET\) in mammalian cells, the gene was amplified by PCR and sub-cloned into pcDNA3.1(-) including a hemagglutinin (HA)-epitope. To obtain \(Opto-dRETMEN2B\) in pcDNA3.1(-), the M955T substitution was introduced using site-directed mutagenesis. To obtain the KD variant, the K805M substitution was introduced using site-directed mutagenesis. To express \(Opto-hRET\) in mammalian cells, the full-length receptor was inserted into a pcDNA3.1(-) vector containing \(AU1-LOV\) (14). All constructs were verified by DNA sequencing. Sequences of the receptors are given in Tables S1 and S2.

Cell culture, transfection and MAPK/ERK pathway activation (HEK293). The MAPK/ERK pathway was assayed in HEK293 cells with the Elk1 trans-reporting system (PathDetect, Agilent). HEK293 cells were maintained in DMEM supplemented with 10% FBS, 100 U/ml penicillin and 0.1 mg/ml streptomycin in a humidified incubator (37°C, 5% CO₂). 50'000 cells were seeded in each well of white clear bottom 96-well plates (triplicates for each construct) coated with poly-L-ornithine (Sigma). Cells were reverse transfected with 3 to 25 ng receptor vector and ~200 ng combined reporter vectors per well using polyethylenimine (Polysciences). Six h 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). Cells were then either illuminated with blue light in a custom incubator (PT2499, ExoTerra) for 8 h or protected from light with foil as described previously (97). After incubation, plates were processed with a
luciferase assay (One-Glo, Promega) and luminescence was detected in a microplate reader (Synergy H1, BioTek). Low-light stimulation (Figure S3B) was performed as previously described (47) using a light blocking sample with an optical density of 1 and an external light intensity of 15 \( \mu \)W/cm\(^2\) (resulting in a final intensity of 1.5 \( \mu \)W/cm\(^2\)).

**Cell culture, RNA interference, transfection and treatments (SH-SY5Y cells).** SH-SY5Y cells (DSMZ ACC 209) were maintained in DEMEM/F-12 (1:1) supplemented with 15% FBS (Sigma), 1% non-essential amino acid solution, 100 U/ml penicillin and 100 \( \mu \)g/ml streptomycin (Life Technologies) in a humidified incubator (37°C, 5% CO\(_2\)). 1.5 \( \times \) 10\(^5\) cells were seeded in each well of a 6-well plate containing two 15 mm coverslips per well. Transient co-transfection of siRNA oligos and DNA plasmids were performed using Lipofectamine 2000 (Thermo Fisher). The following three \( PINK1 \) siRNAs were used at a final concentration of 60 pmol/ml each: siRNA \( PINK1 \) HSS127945/127946/185707 (Life Technologies). To identify transfected cells by fluorescence microscopy, a plasmid encoding GFP was co-transfected (0.2 \( \mu \)g/well; in total, 1.2 \( \mu \)g/well were transfected). Mitochondrial morphology was analyzed 2 days after transfection as described below. For illumination of the cells, the 6-well plate was placed in a LED illumination unit inside the incubator. Cells were illuminated for 4 h at a wavelength of 470 nm and an intensity of 232 \( \mu \)W/cm\(^2\). To activate endogenous RET, cells were treated with recombinant human GDNF (Shenandoah Biotechnology) and human GFR\( \alpha_1 \) (R&D Systems) for 4 h at a final concentration of 100 ng/ml. Signaling pathway inhibitors were added to cells 1 h prior to illumination at the following concentrations: 50 \( \mu \)M LY294002 (PI3K inhibitor, Cell Signaling) or 30 \( \mu \)M PD98059 (MEK1 inhibitor, Cell Signaling). The HA-I\( \kappa B-2S32A/S36A \) plasmid (I\( \kappa B-2S/A \) (98)) was generated by overlap extension PCR using the following primers: mut-I\( \kappa B-2S-fwd \) CCACGACGCGCCCTGGAGCGGCTGAGAAG, mut-I\( \kappa B-2S-rev \) CGTCTTTCATGGCGTCCAGGCCGGCGTCG, BamHI-I\( \kappa B-2S-fwd \) ATATGGATCCTTCAGGGCGAGCGAGCGCTCCAACACACAGTC. The amplified
fragments were digested with BamHI and NotI and cloned into the pcDNA3.1-N-HA vector.

pEGFP-N3 was purchased from Clontech.

**Analysis of mitochondrial morphology (SH-SY5Y cells).** Mitochondria in SH-SY5Y cells growing on 15 mm coverslips were stained for 15 min with 25 nM MitoTracker red CMXRos (Life Technologies) diluted in cell culture media and then washed twice with medium. Mitochondrial morphology of living cells was immediately analyzed with a fluorescence microscope (Nikon Eclipse E400). Cells displaying an intact network of tubular mitochondria were classified as tubular. When this network was disrupted and mitochondria appeared either globular or rod-like, they were classified as fragmented (70). For quantification of mitochondrial morphology, five independent experiments were performed. At least 150 transfected cells were analyzed per condition for each experiment.

**WB analysis (SH-SY5Y cells).** SH-SY5Y cells were analyzed two days after transient transfection for expression of Opto-dRET constructs and PINK1 silencing efficiency. For stabilization of endogenous full-length PINK1, cells were treated with 10 μM FCCP (Agilent) for 2 h before cell lysis. Proteins were detected by WB using a monoclonal rabbit anti-PINK1 antibody (1:1000; Cell Signaling, D8G3) or an anti-HA antibody (1:1000; Covance, 16B12) for the Opto-dRET constructs. Data were normalized to monoclonal mouse anti-β-actin staining (1:2000; Sigma, AC-74).

**Fly strains, maintenance and scoring.** Flies were raised on standard agar, cornmeal and molasses substrate supplemented with 1.5% nipagin. GMR-GAL4 flies were a kind gift of Ross Cagan. PINK1$^{FM6}$/MEF2-GAL4 flies were a gift of Alex Whitworth (University of Cambridge, UK). Transgenic flies expressing Opto-dRET and Opto-dRET$^{MEN2B}$ were generated by injection of pUAST receptor constructs (BestGene). For selection, balanced fly lines (~12 transformants/line) were crossed with GMR-GAL4 flies. Approximately 10 days after crossing, descendants were visually inspected for the presence of a rough retina.
phenotype. Rough retina and hollow thorax phenotypes were scored on a dissecting microscope equipped with a digital camera (M205FA and DFC3000G, Leica Microsystems). Genotypes of fly lines utilized in this study are summarized in Table S3.

**Light stimulation of flies.** Flies were illuminated inside their vials in the custom LED incubator (Figure S3A) set to the temperature and light intensities indicated in the text and figures. Vials containing control flies were wrapped with foil and placed in the same incubator. Light incubators were placed in a controlled environment to maintain humidity at 65%. Experiments with GAL4 drivers were conducted at 29°C to increase receptor expression.

**Scanning electron microscopy.** Adult flies were anaesthetized with CO₂, placed in 25% ethanol for 24 h at room temperature (RT) and dehydrated in a graded ethanol series for 3 days. Samples were dried from 100% ethanol with a critical point dryer (EM-CPD300, Leica Microsystems), gold-coated using a sputter coater (EM-ACE600, Leica Microsystems) and imaged at a magnification of 152X (FE-SEM Merlin compact VP, Carl Zeiss; operated at 3 kV).

**Quantification of rough retina phenotype.** Three analysis methods were applied to retinas. Fused retinal area was defined as the ratio of the total retina area divided by the total disrupted area. The disrupted area was defined as a region containing two or more fused ommatidia. The number of distinct structures was determined using a distortion algorithm (99). The output of the algorithm is mapping of boundaries surrounding single or fused ommatidia that are the structures of interest. Structure count and structure centers were then identified in Fiji. Regularity was determined based on structure centers and their nearest neighbor distance distributions using macros written in Igor Pro (Wavemetrics). Regularity was defined as the ratio of the mean nearest neighbor distance and its SD for each image (59).
Transmission electron microscopy. Thoraces were fixed in 2.5% glutaraldehyde and 2% paraformaldehyde in 0.1 M phosphate buffer (pH 7.4) for 2 h at RT. Samples were post-fixed and contrast enhanced with 1% osmium tetroxide in phosphate buffer for 1.5 h and 1% uranyl acetate in 50% ethanol/water for 45 min. Samples were then dehydrated in a graded ethanol series and embedded in Durcupan (Sigma-Aldrich). Ultrathin sections (70 nm) were sliced using a microtome (EM UC7 Ultramicrotome, Leica Microsystems) and mounted on formvar coated copper slot grids. Images were acquired at a magnification of 9000X (Tecnai 10, FEI/Thermo Fisher Scientific; operated at 80 kV, equipped with OSIS Megaview III camera). The electron dense fraction of the cytoplasm (mitochondria) was determined by manual selection, application of threshold and the area fraction command in Fiji.

ATP determination. Thoraces from two-day old flies were homogenized in 50 µl of extraction buffer (100 mM Tris-HCl, 4 mM EDTA, pH 7.8) supplemented with 6 M Guanidine-HCl using a pellet homogenizer (47747-370, VWR international). The lysate was boiled for 3 min and cleared by centrifugation at 20’000 g for 5 min. The samples were diluted 1:100 in extraction buffer before quantification using a luciferase-based ATP kit (A22066, Thermo Fisher Scientific). Values were expressed relative to total protein concentration measured by using a BCA assay (Pierce). Luminescence and absorbance at 562 nm were measured using the microplate reader. ATP levels were normalized to those of Opto-dRETF female flies.

Negative geotaxis (climbing) assay. Male flies of the indicated genotype have been exposed to blue light (I = 320 µW/cm², λ = 470 nm) or kept in the dark during the indicated developmental time points. For each experiment, males hatching on the same day were pooled. Adults were aged for 2-3 days on standard fly food. On day 3, flies were anaesthetized briefly with CO₂ and 10 flies each were placed in an acrylic glass tube of 30 cm length closed with a flyplug (Carl Roth PK13.1). Flies were allowed to recover and adapt for 1 h. Negative geotaxis climbing performance was then assayed as previously described.
Flies were tapped down and the number of flies reaching the 15 cm mark within 15 s was recorded. 10 technical repeats (1 min break between repeats) were performed for each genotype to obtain an average climbing score (defined as fly count above the 15 cm line / total fly count). For each genotype and condition, at least 5 independent experiments were performed.

Statistical analysis. HEK293 cell assays were performed in triplicate wells and in at least three independent experiments. Statistical analysis was performed using unpaired, two-tailed t-tests for comparison of dark and light conditions.

Fly assays were performed in at least three independent experiments with the number of flies indicated in the figures. Statistical analysis of numerical outcomes was performed using one-way analysis of variance (ANOVA) followed by Bonferroni corrected multiple t-test comparisons. For categorical outcomes (thorax defect and climbing pass), SEMs shown in the figure derived from binomial distributions. Statistical significance was tested using Fisher’s exact tests. Climbing experiments were performed in 10 repeats for each animal group consisting of 10 animals. Statistical significance is indicated using the ‘compact letter display’.

SH-SY5Y cell fragmentation assays were performed in five independent experiments with at least 150 cells per condition in each experiment. Statistical analysis was performed using one-way analysis of variance (ANOVA) followed by Bonferroni corrected multiple t-test comparisons. Statistical significance is indicated using ‘compact letter display’.
Figure 1. Engineering of light-activated RET receptors. (A) hRET and dRET consist of an extracellular ligand-binding domain (LBD), single-span transmembrane domain (TMD) and intracellular domain (KD: kinase domain, CTD: C-terminal tail domain). Activation by GFL and GFRα was shown to result in the formation of a human ternary complex (binding model of Schlee et al. (39)). (B) In light-activated Opto-h/dRET, the LOV domain of the AUREOCHROME1 photoreceptor of V. frigida is incorporated at the receptor C-terminus. (C) MAPK/ERK pathway activation in response to blue light (I = 250 µW/cm², λ = 470 nm) for HEK293 cells transfected with Opto-hRET, Opto-dRET or Opto-dRETMEN2B. Light units (LU; mean ± SD) for dark treated cells and illuminated cells are given (n = 9 to 18, three independent experiments, t-test, *: p<.0001).
**Figure 2.** Induction of retina roughening and phenotype quantification. (A) Developmental time window targeted by light in retina experiments. (B-G) Representative retina SEM images. Scale bar: 0.1 mm. (H and I) Quantification of rough retina phenotypes of one-day old flies as fused area and the number of structures identified. “M2B” denotes Opto-dRET^{MEN2B}. For H and I, mean ± SD for the indicated number of flies is given (at least three independent experiments). Means sharing the same label are not significantly different (ANOVA/Bonferroni corrected t-tests, p>.04). Light intensity was 385 µW/cm².
Figure 3. Suppression of thorax defects and locomotion deficits. (A) Time window targeted by light in experiments with PINK1^{B9} flies. Illumination of pupal and adult stages prevented lethality observed upon Opto-dRET signaling in earlier stages (e.g., Opto-dRETMEN2B flies were grown at 18°C to prevent lethality during development; see Main Text). (B) Percentage of two-day old flies with a degenerate thorax phenotype. Representative bright field thorax images shown on the right. Hollow thorax is highlighted by the red arrow. (C) Climbing ability of flies. “M2B” denotes Opto-dRETMEN2B. PINK1 “+” denotes the WT gene. For B and C, counts ± SE for the indicated number of flies (n) is given (see Materials and Methods for repetitions in climbing assays). Means sharing the same label are not significantly different (Fisher’s exact test, p>.04). Light intensity was 320 μW/cm².
Figure 4. Improved mitochondrial structure and function. (A) ATP content in fly thoraces from PINK1<sup>B9</sup> flies at the indicated conditions. (B-E) Representative TEM images of thoracic indirect flight muscles. Arrow heads indicate mitochondria that are either electron dense (B: controls, E: illuminated PINK1<sup>B9</sup> Opto-dRET flies) or malformed with disintegrated cristae (C: PINK1<sup>B9</sup> flies, D: PINK1<sup>B9</sup> Opto-dRET flies in the absence of light). Scale bar: 2 µm. (F) Analysis of mitochondrial density in TEM images. “M2B” denotes Opto-dRET<sup>MEN2B</sup>. PINK1 “+” denotes the WT gene. For A, mean ± SD for the indicated number of flies is given (at least three independent experiments). For F, mean ± SD for the indicated number of micrographs is given (at least three independent experiments). Means sharing the same label are not significantly different (ANOVA/Bonferroni corrected t-tests, p>.04). Light intensity was 320 µW/cm<sup>2</sup>.
Figure 5. Rescue of mitochondrial fragmentation in human cells. (A and B) WB analysis of PINK1 knock-down by siRNA and Opto-dRET expression. (C) Representative images for fragmentation of mitochondria induced by PINK1 silencing. Red: MitoTracker. Green: GFP marker. Scale bar: 200 (column 1, 2) or 20 µm (columns 3, 4). (D) Quantification of mitochondrial fragmentation upon stimulation of RET, Opto-dRET, Opto-dRET\textsuperscript{MEN2B} or Opto-dRET\textsuperscript{KD}. (E) Quantification analysis of mitochondrial fragmentation upon light activation of Opto-dRET and inhibition of NF-kB signaling (by IκB-2S/A), PI3K (by LY294002) or MEK1 (by PD98059). For D and E, mean ± SD for five independent experiments is given (at least 150 cells per condition in each experiment). Means sharing the same label are not significantly different (ANOVA/Bonferroni corrected t-tests, p>.04).
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