

ORIGINAL ARTICLE

Identification of 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase as a novel autophagy regulator by high content shRNA screening

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Deregulation of autophagy has been linked to multiple degenerative diseases and cancer, thus the identification of novel autophagy regulators for potential therapeutic intervention is important. To meet this need, we developed a high content image-based short hairpin RNA screen monitoring levels of the autophagy substrate p62/SQSTM1. We identified 186 genes whose loss caused p62 accumulation indicative of autophagy blockade, and 67 genes whose loss enhanced p62 elimination indicative of autophagy stimulation. One putative autophagy stimulator, 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase 4 (PFKFB4), drives flux through pentose phosphate pathway. Knockdown of PFKFB4 in prostate cancer cells increased p62 and reactive oxygen species (ROS), but surprisingly increased autophagic flux. Addition of the ROS scavenger N-acetyl cysteine prevented p62 accumulation in PFKFB4-depleted cells, suggesting that the upregulation of p62 and autophagy was a response to oxidative stress caused by PFKFB4 elimination. Thus, PFKFB4 suppresses oxidative stress and p62 accumulation, without which autophagy is stimulated likely as a ROS detoxification response.

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INTRODUCTION

Autophagy maintains protein and organelle homeostasis and enables survival in the absence of exogenous nutrients. Although active in most tissues in low levels, autophagy is markedly induced by stress and starvation where it eliminates damaged proteins and organelles, and recycles their components into metabolic and biosynthetic pathways. Autophagy is critical to survive starvation and to prevent the toxic buildup of damaged or superfluous intracellular components.¹

Autophagy-mediated preservation of mitochondrial metabolism is exploited by tumor cells and is associated with promotion of mammary, lung and pancreatic cancers, suggesting that autophagy-inhibiting strategies may have therapeutic efficacy in these cancer subtypes.^{2–12} In other settings, autophagy may suppress tumorigenesis by mitigating chronic tissue damage,^{13–17} leading to the general consensus that the role of autophagy in cancer is context dependent.¹² Defective autophagy has been linked to neurodegenerative disorders such as Alzheimer's and Parkinson's diseases, as well as liver disease, all characterized by the accumulation of toxic protein aggregates.^{18,19} Autophagy prevents muscle wasting, and impaired autophagy has been implicated in various models of muscular dystrophy.^{20–23} Taken together, this suggests that autophagy-modulating therapies will be of great clinical interest: autophagy stimulators for neurodegenerative and degenerative musculo-skeletal disorders and perhaps also cancer prevention; and autophagy inhibitors

to target a metabolic vulnerability in aggressive cancers. Although targeted small molecule modulators of autophagy are in development, the optimal targets remain unknown.

Studies assessing the functional consequences of pharmacologic inhibition of autophagy in multiple cancers using the lysosomotropic agents and chloroquine (CQ) and hydroxychloroquine (HCQ) are underway. These inhibitors block autophagy at the terminal step of cargo degradation by interfering with lysosome function. CQ and HCQ suppress tumor growth in mouse models of cancer and in human cancer cell line xenografts, and have shown efficacy in patient-derived xenograft models alone and in combination with cancer therapeutics.^{9,24–32} These results have prompted clinical trials with HCQ to block autophagy and enhance the activity of cancer therapy in refractory malignancies.^{33,34} Although the early findings of these trials are promising^{35–40} there is an unmet need for more specific tool compounds to probe actionable targets in the autophagy pathway. The central problem lies in the fact that HCQ nonspecifically interferes with lysosome function and likely has off-target effects.⁴¹ The current challenge is to identify the genes and regulatory points in the autophagy pathway that merit anticancer drug development.

To identify new autophagy regulators, we designed and executed a novel image-based high-throughput short hairpin RNA (shRNA) screen based on functional elimination of the autophagy cargo receptor and substrate p62/SQSTM1.

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p62 localizes to the site of autophagosome formation where it recognizes ubiquitinated cargo via its ubiquitin-binding domain and delivers it to the autophagosome via its LC3 interaction region domain.^{42–44} Importantly, p62 is an autophagy substrate that is degraded along with its cargo; thus, monitoring p62 levels can be used as a surrogate of the functional state of autophagy in a given cell. Deletion of the essential autophagy genes *Atg5* or *Atg7* in mice leads to profound p62 accumulation, often in large aggregates.¹² Screening shRNA libraries for genes that alter the ability of cells to eliminate aggregated p62 following metabolic stress and recovery is a novel approach to identify autophagy regulators upstream of the terminal substrate degradation step in the pathway. Although our screen was designed to identify autophagy regulators, it is likely our candidate gene lists also contain p62 modulators, which are of interest independent of their regulation of autophagy given the importance of p62 in cancer.

p62 is a multi-functional protein that has key roles in nuclear factor-κB signaling (via TRAF6), bone remodeling (mutations in p62 are associated with Paget's disease) and cancer where its overexpression has been linked to tumorigenesis.^{15,45–49} p62^{-/-} mice are obesity prone and develop diabetes with age.⁵⁰ More recently, p62 has been linked to amino-acid sensing via TORC1, controlling the localization and activity of the kinase via its interaction with TRAF6.^{51,52} Thus, p62 is emerging as a nutrient sensor regulating redox balance and mammalian target of rapamycin (mTOR) activation, as well as autophagy.

We used the ability of cells to eliminate p62 following ischemic stress as a readout of the functional autophagy status of the cell. We identified 186 positive and 67 negative regulators of p62 accumulation from function-based libraries encompassing the protein kinome, components of the vesicle trafficking machinery and a collection of GTPases. Our results highlight key interactions with core components of the endocytic and vesicle transport system, actin cytoskeleton, nutrient sensing and calcium signaling pathways required for functional autophagy. Metabolically relevant kinases and a subset of Rab GTPases were identified among the negative regulators of pathway. Many of these genes, particularly those of the vesicle trafficking library, are poorly annotated and this work will locate them within the context of autophagic regulation.

One of the high-priority candidate genes identified by the screen, the bi-functional enzyme 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase-4, PFKFB4, was selected for further investigation. PFKFB4 belongs to a family of four rate-limiting enzymes (PFKFB 1–4), which control entry into glycolysis by regulating fructose-2,6-bisphosphate (F2,6BP) levels.^{53–55} We report here that PFKFB4 regulates autophagy by influencing redox balance in the cell. We also implicate the role of many new genes in the regulation of the autophagy pathway, which has the potential to inform the design of autophagy-modulating therapies for cancer and other autophagy-related diseases.

RESULTS

p62 modulator screen to identify autophagy regulators

We developed and executed a novel cell-based shRNA screen designed to capture both positive and negative regulators of autophagy at all steps upstream of substrate degradation. This was done in a single assay using cells stably expressing a fluorescent version of the autophagy cargo receptor and substrate p62 (EGFP-p62) as a reporter of functional autophagy status (*Beclin1*^{+/-}; EGFP-p62 iBMK cells). The screen is predicated on the fact that p62 accumulates under stress in all cells and is eliminated by functional autophagy. In the absence of autophagy, this stress-induced accumulated p62 persists, and is more marked depending on the severity of the autophagy defect (that is, cells with

homozygous deletion of *Atg5* exhibit a greater impairment in p62 elimination than cells with allelic loss of *Becn1*).^{15,48} Thus, monitoring the levels of p62 in cells following stress and recovery can be used as a surrogate of autophagy functionality.

We designed our autophagy screen differently from those conducted previously.^{56–62} Our assay used murine immortalized epithelial cells possessing an allelic loss of the essential autophagy gene, *Becn1*, chosen over 293T or HeLa cells that are common choices for cell-based lentiviral screens, as the tumorigenic potential of the mouse epithelial cells is known and high-priority candidate genes can be rapidly validated in syngeneic mouse models. Moreover, the *Becn1* heterozygous cells display a partial defect in autophagy that amplifies the p62 signal, which enables the identification of both positive and negative regulators and maximizes the discovery potential of the screen.

Our assay stimulus, metabolic stress (1% oxygen, and glucose deprivation) (Figure 1a), is highly physiologically relevant; autophagy localizes to hypoxic regions of tumors thus identifying genes capable of regulating autophagy under these conditions would be advantageous for cancer therapy.¹³ Importantly, our screen provides a functional readout of autophagy by measuring clearance of fluorescently tagged autophagy substrate EGFP-p62 containing aggregates during recovery, differentiating it from LC3-based screens that are limited to the detection of autophagosome formation at the initial step of autophagy activation. Thus, our assay enables the identification of autophagy modulators at both the beginning and throughout the late stages of the pathway in a single screen in addition to identifying modulators of p62 homeostasis.

We first evaluated the consequences of lentiviral infection and selection with puromycin on the elimination of p62 from autophagy-competent or -deficient cell lines. Although both infection and selection modestly increased baseline p62 levels in this system, EGFP-p62 accumulated in both *Becn1* wild-type and heterozygote cell types under stress, and a clear difference in the ability of the autophagy-competent and -deficient cells to eliminate p62 was retained as measured by western blot and immunofluorescence, consistent with prior studies¹⁵ (Figures 1a and c). Predictably, a markedly shorter timeframe of stress and recovery was needed to visualize accumulation and elimination of p62 when coupled to the additional stresses of infection and selection (7-h stress, 18-h recovery compared with 3-day stress, 2-day recovery).¹⁵ A schematic of the screening protocol and representative images from the screen is provided (Figure 1d). Three classes of candidate genes were expected: those whose expression loss restored clearance of p62 (putative-negative regulators of autophagy), genes whose expression loss further impaired p62 elimination leading to an accumulation of p62 (putative-positive regulators of autophagy) and genes whose expression loss had no effect on p62 elimination.

Loss of *mTOR*, the master negative regulator of autophagy, restored functional autophagy and clearance of p62 (Figure 1d). Loss of the gene that encodes IKK-β, a known autophagy promoter,⁶³ *Ikkb*, resulted in significant accumulation of p62 (Figure 1d). Those genes whose loss had no effect on p62 homeostasis, illustrated by cells infected with a hairpin targeting luciferase, constitute a third category (shRNA Control, Figure 1d). Single-cell image analysis of p62 abundance was achieved via a customized Cellomics algorithm (see Materials and methods section for complete algorithm settings) (Figure 1e). Representative images of cells infected with control shRNA (pLKO-emptyT) or shRNA targeting p62 (pLKO-p62 #3) are shown with and without overlay of the algorithm.

Screen performance

We screened > 11 000 hairpins targeting 1361 unique genes from function-based libraries encompassing the protein kinome,

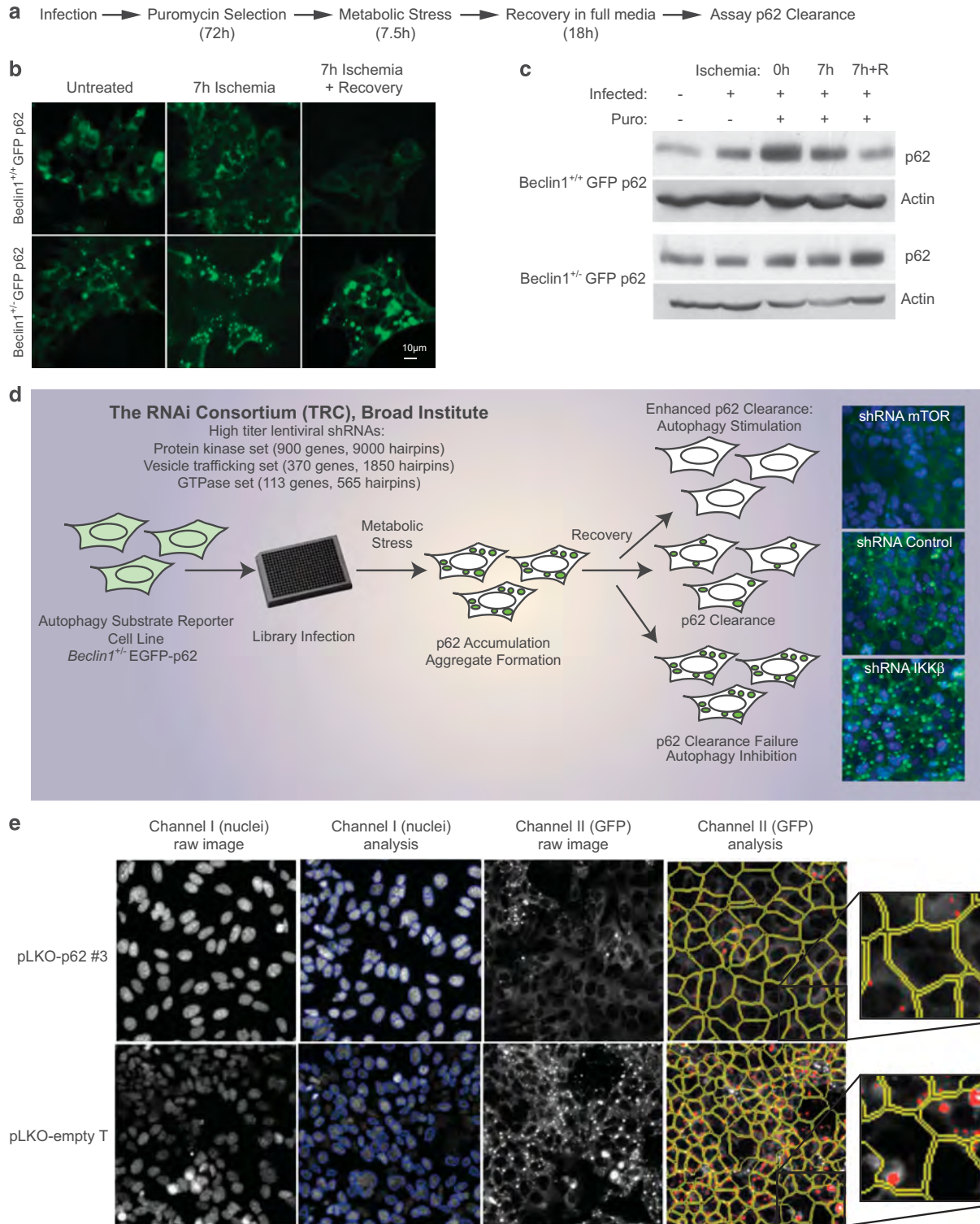


Figure 1. Identification of autophagy substrate modulators by high content shRNA screening. **(a)** Experimental timeline detailing sequence of infection, selection and assay for p62 elimination. **(b)** *Beclin1^{+/+}* or *Beclin1^{-/-}* iBMK cells stably expressing EGFP-p62 were infected with shRNA targeting Luc1510, selected with puromycin and subjected to a time course of 7 h ischemia and recovery. Representative images of GFP-p62 accumulation during stress and recovery demonstrating impaired elimination of p62 from the *Beclin1^{-/-}* iBMK cell line during recovery in full media. Scale bar is 10 μm. **(c)** Western blot analysis of GFP-p62 accumulation and elimination in *Beclin1^{+/+}* or *Beclin1^{-/-}* iBMK cells stably expressing EGFP-p62 infected with shRNA targeting Luc1510, selected with puromycin and subjected to a time course of 7-h ischemia and recovery as indicated. **(d)** Schematic representation of the p62 modulator screen. Representative images from the screen are provided for each class of regulator. **(e)** A customized Cellomics algorithm was used to score p62 abundance at the single-cell level. Images from pilot studies using p62 or non-targeting (emptyT) hairpins are provided with and without the overlaid analysis. Cells were counterstained with Hoechst 33342 to visualize nuclei (blue, Channel 1). Channel 2 maps the cell borders, and detects GFP-p62 levels within a designated radius from the nucleus (ring spot total intensity). Nine images per field were collected to ensure representative coverage. A p62 aggregate score equal to mean ring spot total intensity/mean nuclei was calculated for each well.

components of the vesicle trafficking machinery and a large collection of GTPases (Supplementary Table S1). These gene sets were enriched in genome-wide screens for autophagy regulators^{56,64} and are predicted to contain signaling, as well as machinery regulators of the autophagy pathway. Genes were targeted by 5–10 unique hairpins (depending on the library) and infected in quadruplicate.

Primary screen data were filtered to remove hairpins with poor infection efficiency or nuclei counts < 1500 that likely represent autofocus errors or toxicity. The distribution of the remaining hairpins plotted against robust-z-score is shown (Figure 2a). Negligible differences in batch-to-batch distribution of hairpins were observed, enabling analysis to be conducted on the entire screen data set at once (Figure 2b). A non-Gaussian distribution of hits, weighted heavily toward identification of positive regulators of the autophagy pathway was observed. This distribution reflects the biology of the libraries screened (that is, likely many positive regulators of autophagy among vesicle trafficking components) (Figure 2c).

Data were analyzed by the Broad Institute's proprietary Gene-E software.⁶⁵ Taking the union of the second best and weighted sum metrics, we identified 186 high-priority positive and 67 negative regulators of autophagy (Tables 1 and 2, and Supplementary Table S1). Candidate gene lists were compared with a March 2014 build of the murine genome hosted by NCBI. Genes with a 'withdrawn' status were removed from our lists, whereas genes whose entry's had been updated were retained. Criteria for inclusion in the positive regulator list was a gene ranking within the top 200 'live' genes in the screen possessing two or more hairpins scoring within the top 1000 hairpins in the overall screen. This cut-off represents a 14% hit rate. We extended our criteria to capture known regulators of the pathway and maximize discovery. Negative regulators were defined as genes with two or more hairpins in the top 500 hairpins in the screen, which corresponds to a 5% hit rate.

Many known positive and negative regulators of the pathway were identified increasing our confidence in the candidate gene lists. PANTHER gene ontology analysis of the p62 modulators revealed an equivalent distribution of candidate genes within the biological process categories between the positive and negative regulators of the pathway (Figures 2d and e and Supplementary Table S2).⁶⁶

Candidate-positive regulators of autophagy

Genes whose loss was associated with further impairment of p62 elimination are putative stimulators of autophagy (Table 1). This high-priority candidate gene list was enriched with genes involved in metabolism and nutrient sensing (*Ulk1*, *Becn1*, *Prkaa1*, *Prkaa2*, *Stradb*, *Prkag1*, *Mark4* and *Pfkfb4*), core components of the endocytosis and protein trafficking machinery (*Cav3*, *Rab7*, *Snap29*, *Dmn2*, *Ap3m2*, *Sec23a* and *Vti1b*), interactions with the actin cytoskeleton (*Pik3cg*, *Arcp1a*, *Fgfr4*, *Rac3*, *Arcp2*, *Pak4*, *Rras2*, *Rac1*, *Araf* and *Hras1*) and calcium/calmodulin-dependent signaling (*Camk1g*, *Camk4*, *Phkg1*, *Camk2g*, *Phkg2*, *Phka1*, *Itpka*, *Camkk1* and *Doc2b*). Intracellular calcium levels may be required for vesicle fusion events, which occur at various steps within the autophagy pathway, or may lead to 5' adenosine monophosphate-activated protein kinase (AMPK)-mediated activation of the autophagy machinery via Ca²⁺/calmodulin-dependent kinase kinase-β (CamKKβ).^{67,68} Genes involved in Toll-like (*Pik3cg*, *Irak1*, *Irak3*, *Map2k3*, *Rac1*, *Mapk9* and *Ikbkb*) and Ephrin receptor signaling (*Epha5*, *Fgfr4*, *Epa7*, *Ephb6*, *Epha6*, *Ryk*, *Mst1r* and *Tie1*) were also highly enriched in our positive regulator list. Moreover, multiple mitogen-activated protein kinase pathway components were associated with functional autophagy in this system (*Map2k3*, *Araf*, *Mknk2*, *Mapk9*, *Map3k14* and *Hras1*), consistent with work from our group demonstrating that Ras transformation upregulates the autophagic machinery.⁸

The endosomal sorting complexes required for transport (ESCRT) machinery sort ubiquitinated cargo into multivesicular bodies and mediate membrane fusion and abscission events within the cell. Defects in the machinery have been associated with protein aggregate accumulation and neuronal toxicity. Although the precise details of the defects remain unclear, it is suspected that these phenotypes arise from problems with vesicle docking and fusion with lysosomes. Indeed, loss of multiple ESCRT components in mouse and human cell lines is associated with an accumulation of autophagosomes and amphisomes.^{69,70} Our present work confirms the involvement of the ESCRT-0 component *Hrs* in the positive regulation of autophagy noted previously,⁷¹ and additionally identifies the ESCRT-II component *Vps36*, ESCRT-III components *Chmp4B* (the homolog of *Snf7b*) and *Chmp5* (*Vps60* homolog), and *Vps4b* and *Vti1b*, components of the abscission complex, as positive regulators of autophagy.

Nucleoside diphosphate kinase B (Nme)-2 and -7 were identified as positive regulators of autophagy in our screen. Intriguingly, *Nme2* was recently noted to be required for mitophagy in a genome-wide screen.⁵⁸ Other intriguing regulators identified in this work are *Mark4* and *Agap3*. *Mark4* is an AMPK-related protein kinase recently identified in *Drosophila* as a negative regulator of mTORC1 activation via interaction with the Rag complex, preventing amino-acid induction of mTOR.⁷² *Agap3* is an ARF family GTPase previously described to be critical for degradation of poly-glutamine containing proteins that responds to reactive oxygen species (ROS)-induced by poly-glutamine accumulation and leads to degradation of these aggregated proteins via ubiquitin-proteasome system.⁷³ Its role in autophagy has yet to be described.

Candidate-negative regulators of autophagy

Genes whose loss enhanced p62 elimination are putative-negative regulators of autophagy (Table 2). This list included known inhibitors of the autophagy pathway (*mTOR*, *Akt1*, *Dgka*, *Egfr* and *Mapk14*), and genes involved in intracellular protein transport, particularly vesicle docking (*Scfd1*, *Pldn*, *Stxbp3a* and *Vps33*) and Rab GTPases (*Rab39b*, *Rab34*, *Rab71* and *Rab9*). A significant enrichment of membrane-associated guanylate kinase homologs was observed (*Mpp3*, *Dlg4*, *Cask* and *Lrguk*). These proteins share conserved SH3, PDZ and guanylate kinase domains, and are thought to function primarily as scaffolds to cluster receptors for signaling events. Of note, *CASK* is known to interact with the E3 ligase Parkin, a tumor-suppressor and key mediator of mitophagy via its PDZ domain.⁷⁴

Validation of known autophagy regulators identified in the screen
A subset of candidate genes was validated to confirm on-target knockdown of expression of the gene and the scoring of the p62 phenotype. This was achieved by evaluating p62 clearance or accumulation by fluorescence (EGFP-p62) and target protein expression by western blot following infection with two control hairpins targeting *RFP* and all of the available hairpins for the gene in question (minimum of five unique hairpins). Loss of the essential autophagy gene *Atg7*, not contained in the screen set itself but added as a proof of concept control, or the known positive regulators of the autophagy pathway *Ulk1*, *Becn1* or *Prkaa1* (which encodes the alpha1 subunit of AMPK) resulted in a significant accumulation of p62 (Figures 3a and b). Note that the antibody used to probe AMPK recognizes both the α1 and α2 subunits, while the hairpins in this validation experiment solely target *Prkaa1*. Similarly, silencing the ESCRT-0 component *Hrs* or the late endosomal small GTPase *Rab7*, known to be required for proper fusion of the autophagosome with the lysosome,^{75–77} resulted in a failure to eliminate p62 during recovery, confirming the need for these genes for functional autophagy. Elongation of the immature autophagosome depends on two ubiquitin-like

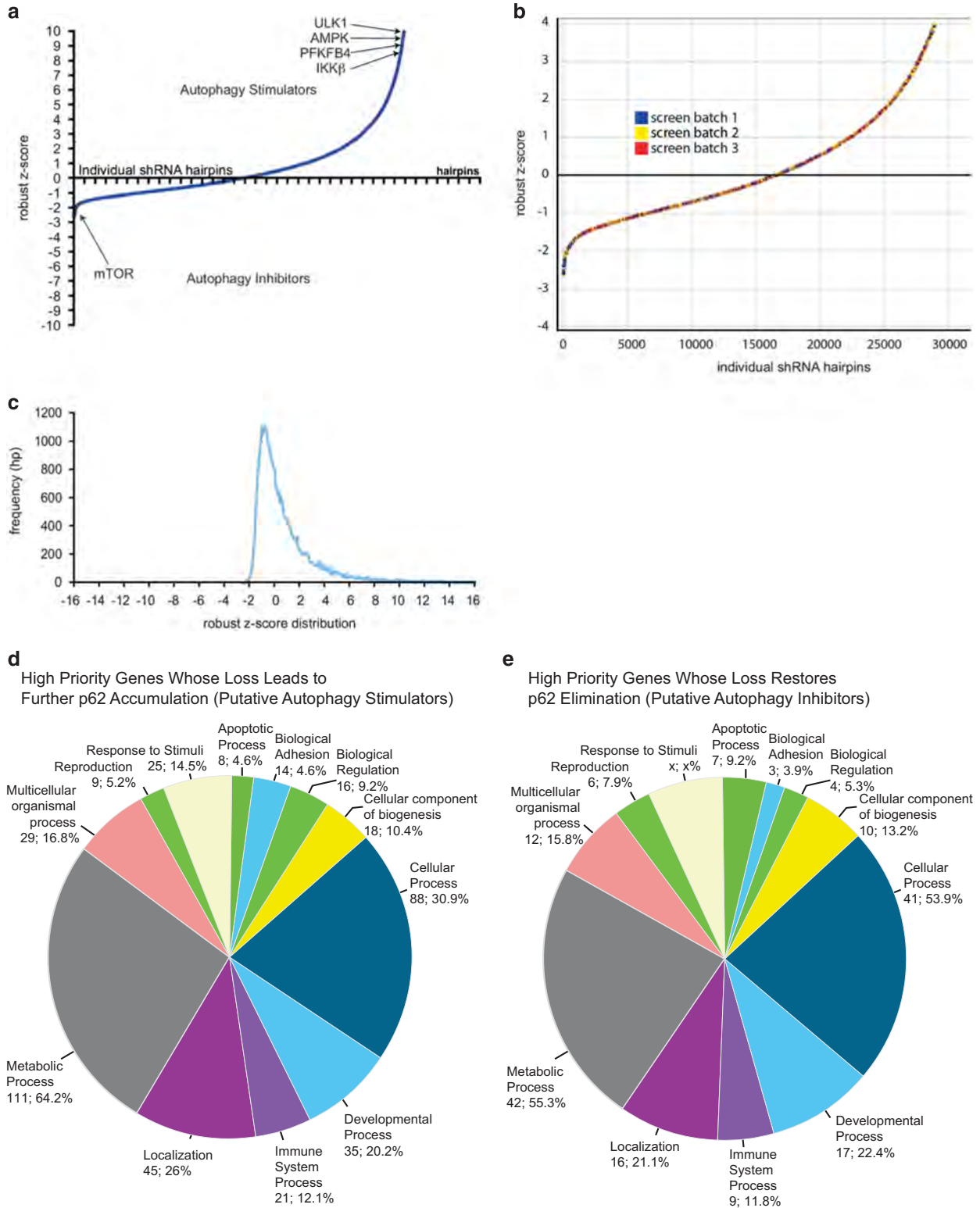


Figure 2. Screen performance and analytics. **(a)** Distribution of all hairpins in the screen after filtering to remove hairpins with poor infection efficacy and wells containing < 1500 nuclei, likely representing autofocus errors, plotted against robust-z-score. The location known positive (ULK1, AMPK and IKK β) and negative regulators of the autophagy pathway (mTOR) is shown. **(b)** Overlay of hairpin distribution for all three of the screen batches demonstrating little to no batch-to-batch variation in the data. **(c)** Histogram of hairpin frequency plotted against robust z-score **(d,e)**. Classification of autophagy screen hits arranged by Biological Process by the PANTHER system. # genes in the screen data set found within each category; % of genes in the data set contained within each category.

Table 1. Positive regulators of p62 elimination (putative autophagy stimulators)

<i>Symbol</i>	<i>Gene ID</i>	<i>Symbol</i>	<i>Gene ID</i>	<i>Symbol</i>	<i>Gene ID</i>	<i>Symbol</i>	<i>Gene ID</i>
2810408M09Rik	66566	Epha6	13840	Nkiras2	71966	Rragc	54170
Acvr2a	11480	Epha7	13841	Nme2	18103	Rras2	66922
Acvr2b	11481	Ephb6	13848	Nme7	171567	Ryk	20187
Adrbk2	320129	Etnk1	75320	Ntpcr	66566	Sbk3	381835
Agap3	213990	Etnk2	214253	Nuak1	77976	Sec23a	20334
Ak2	11637	Fastkd5	380601	Pak4	70584	Septin11	52398
Aldh18a1	56454	Fgfr4	14186	Pank3	211347	Sh3gl2	20404
Ankk1	244859	Gbp3	55932	Pdgfrl	68797	Shpk	74637
Anxa7	11750	Gck	103988	Pdpk1	18607	Sik1	17691
Ap1g1	11765	Gm318	545204	Pfkfb3	170768	Smg1	233789
Ap1s1	11769	Gm4776	212225	Pfkfb4	270198	Smok2a	27263
Ap3m2	64933	Gm4862	229879	Pftk1	18647	Smok4a	272667
Araf	11836	Gm5285	383956	Phka1	18679	Snap29	67474
Arf5	11844	Gm5374	385049	Phkg1	18682	Snx13	217463
Arfrp1	76688	Gm5425	432482	Phkg2	68961	Snx16	74718
Arl4d	80981	Gm9824	432447	Pi4ka	224020	Snx2	67804
Arpc1a	56443	Golga5	27277	Pik3c2a	18704	Snx3	54198
Arpc2	76709	Gpn1	74254	Pik3cg	30955	Snx4	69150
B2m	12010	Grk6	26385	Pkmyt1	268930	Stk22d	117545
Becn1	56208	Gsg2	14841	Plaur	18793	Stk35	67333
Camk1g	215303	Gtpbp2	56055	Prkaa1	105787	Stradb	227154
Camk2g	12325	Gtpbp4	69237	Prkaa2	108079	Stx16	228960
Camk4	12326	Gucy2e	14919	Prkar1	19082	Synj2	20975
Camkk1	55984	H2bfm	69389	Prkar2b	19088	Syt15	319508
Cav3	12391	Hgs	15239	Prkcd	18753	Syt8	55925
Cdadcl	71891	Hipk2	15258	Prkcdbp	109042	Tesk2	230661
Cdc2l1	12537	Hras1	15461	Prkc22	404705	Tie1	21846
Cdc42bpg	240505	Hspb8	80888	Prps11i	75456	Tjp2	21873
Cdk3	69681	lkbkb	16150	Psmc1	19179	Trp53rk	76367
Cdkl5	382253	Irak1	16179	Rab14	68365	Trpm6	225997
Cerk	223753	Irak3	73914	Rab15	104886	Tspan6	56496
Chek1	12649	Itpka	228550	Rab19	19331	Tspan7	21912
Chmp4b	75608	Keap1	50868	Rab20	19332	Tssk3	58864
Chmp5	76959	Khk	16548	Rab3d	19340	Txk	22165
Ciita	12265	Kpnb1	16211	Rab6b	270192	Uck1	22245
Ckmt2	76722	Map2k3	26397	Rab7	19349	Uck2	80914
Csnk1g1	214897	Map3k14	53859	Rab8a	17274	Ulk1	22241
Cyth3	13132	Mapk9	26420	Rac1	19353	Vps13a	271564
Dab2	217480	Mapkapk3	102626	Rac3	170758	Vps36	70160
Dgkb	217480	Mark4	232944	Rap1a	109905	Vps39	269338
Dlg4	13385	Mast3	546071	Rasl10b	276952	Vps4b	20479
Dnm2	13430	Mknk2	17347	Rerg	232441	Vps8	209018
Doc2b	13447	MLK1	74568	Rhoc	11853	Vti1b	53612
Drg2	13495	Mst1r	19882	Rhog	56212	Wnk4	69847
Eef1a2	13628	Mylk4	238564	Rhot1	59040	Yes1	22612
Eif2s3y	26908	N4bp2	333789	Rhov	228543		
Epha5	13839	Nkiras1	69721	Rps6kl1	238323		

Table 2. Negative regulators of p62 elimination (putative autophagy inhibitors)

<i>Symbol</i>	<i>Gene ID</i>	<i>Symbol</i>	<i>Gene ID</i>	<i>Symbol</i>	<i>Gene ID</i>	<i>Symbol</i>	<i>Gene ID</i>
1810024B03Rik	329509	Dlg4	13385	Mtor	56717	Ripk3	56532
4932415M13Rik	211496	Egrf	13649	Musk	18198	Sar1a	20224
Adk	11534	Eif2ak1	15467	N4bp2	333789	Scfd1	76983
Akt1	11651	Ephb1	270190	Nek5	330721	Snx10	71982
Alpk3	116904	Ern1	78943	Pfkfb4	270198	Stxbp3a	20912
Arf3	11842	Gm14147	381390	Pik3c3	225326	Sybu	319613
Arhgap24	231532	Gm4922	237300	Pldn	18457	Tlk2	24086
Atp6v0a1	11975	Gm5374	385049	Plk2	20620	Tpr	108989
Bmpr1a	12166	Grk4	14772	Prkaca	18747	Trib3	228775
Brdt	114642	Hip1	215114	Prkc2	18762	Trrap	100683
Cask	12361	Hip1r	29816	Rab34	19376	Tssk3	58864
Cdk6	12571	Kpna2	16647	Rab39b	67790	Txndc3	73412
Cdk8	264064	Lrguk	74354	Rab71l	226422	Vamp8	22320
Chmp1a	234852	Map3k11	26403	Rab9	56382	Vapa	30960
Cpne3	70568	Map4k3	225028	Rabl2a	68708	Vps33b	233405
Ddr2	18214	Mapk14	26416	Rhoa	11848	Vta1	66201
Dgka	13139	Mpp3	13384	Rhoh	74734		

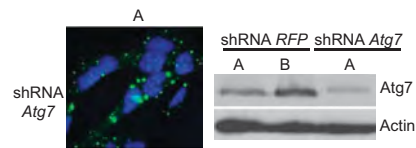
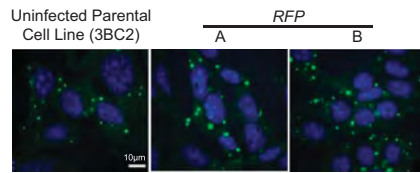
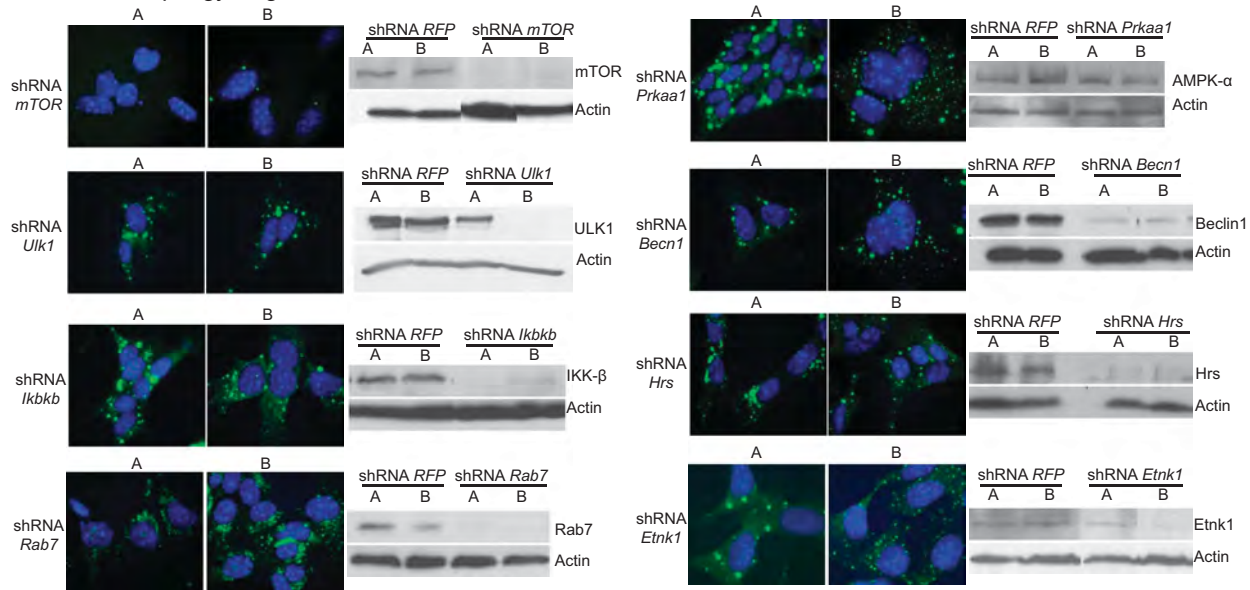
a Control Hairpins**b** Known Autophagy Regulators

Figure 3. Validation of selected screen hits. **(a)** Representative low-throughput validation images from experiments with uninfected parental *Beclin1*^{+/-} iBMK cells (3BC2- clone #D3) or *Beclin1*^{+/-} iBMK cells infected with two distinct hairpins targeting *RFP* or a hairpin targeting *Atg7*, assaying GFP-p62 clearance following metabolic stress (7.5 h) and recovery (18 h) in full media. Nuclei were counterstained with Hoechst 33342 to facilitate visualization. Scale bar is 10 μm. Non-validating hairpins were spliced out of the western blots presented for aesthetic reasons. All lanes presented for a given gene were contained within a single gel. Complete gels are shown in Supplementary Figure S1. **(b)** Validation of a subset of known autophagy regulators identified in our screens. For a candidate gene to pass validation at least two individual hairpins targeting distinct regions of the message must silence expression of the target gene by western blot and correlate well with GFP-p62 phenotype observed microscopically. *Beclin1*^{+/-} iBMK cells were infected with hairpins targeting *RFP* or *mTOR*, *Ulk1*, *Prkaa1*, *Becn1*, *Ikbkb*, *Hrs*, *Rab7* or *Etnk1*, subjected to selection, and assayed for GFP-p62 clearance following metabolic stress (7.5 h) and recovery (18 h) in full media by immunofluorescence. Nuclei were counterstained with Hoechst 33342 to facilitate visualization. Scale bar is 10 μm. Confirmation of on-target gene silencing was obtained by western blot analysis. TRC hairpin identifiers and antibody details are contained in the Supplemental methods section. Non-validating hairpins were spliced out of the western blots presented for aesthetic reasons. All lanes presented for a given gene were contained within a single gel. Complete gels are shown in Supplementary Figure S1.

conjugation systems that cleave and conjugate LC3 with phosphatidylethanolamine that is required for autophagosome maturation closure. We identified genes needed for phosphatidylethanolamine biosynthesis, ethanolanamine kinase 1 and 2 (*Etnk1*, *Etnk2*) (Table 2 and Figure 3b) as essential components for functional autophagy.

The bi-functional enzyme PFKFB4 was among the highest scoring genes that inhibited p62 elimination in our screen; found in the top 20 of all screened genes with two or more scoring hairpins in both weighted sum and second best metrics, a ranking shared by the well-known autophagy regulators *Ulk1* and *Prkaa2*. The strength of this hit, coupled with the growing list of PFKFBs as druggable cancer targets and emerging evidence of increased expression of PFKFB4 in multiple solid tumor types including breast, colon and prostate led us to select PFKFB4 for further validation⁷⁸⁻⁸¹ (Table 1).

PFKFB4 deficiency could have two consequences in terms of autophagy (Figure 4a). Depletion of the enzyme could lead to increased glycolytic flux, ATP generation, activation of mTOR and autophagy inhibition. Alternatively, PFKFB4 deficiency

could lead to decreased flux into the PPP, compromising nicotinic adenine diphosphonucleotide, reduced (NADPH) generation, leading to enhanced ROS levels sufficient to induce autophagy.

Validation of PFKFB4 as an autophagy regulator

As PFKFB4 has been reported to be essential for survival of multiple cancer types,^{78,82,83} we speculated that lentiviral-mediated stable knockdown would be problematic, and moved into a transient silencing system. To confirm the original screening result in this second system, we transiently silenced expression of PFKFB4 in the *Beclin1*^{+/-}; EGFP-p62 iBMK cells using Dharmacon Smartpool small interfering RNAs. An experimental timeline is provided (Figure 4b). PFKFB4 knockdown was associated with EGFP-p62 accumulation, impaired cell growth and significantly decreased viability (approximately 50% reduction) (Figure 4c) in agreement with published reports that PFKFB4 is essential for cancer cell survival.^{78,82,83} Moreover, PFKFB4 knockdown induced robust EGFP-p62 accumulation as measured by fluorescence microscopy and western blotting (Figures 4c and d). Surprisingly, despite inducing p62 accumulation,

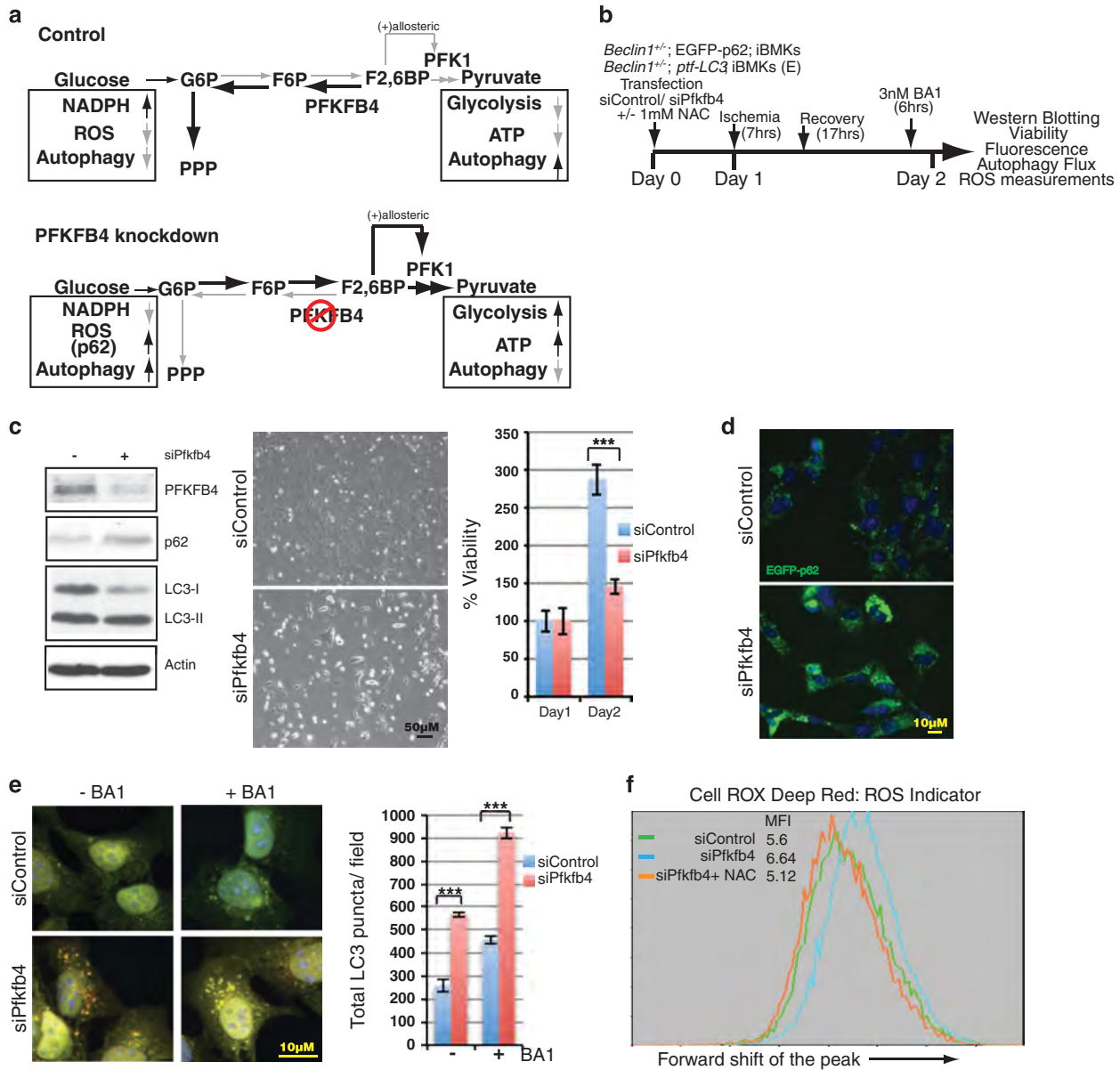


Figure 4. Validation of PFKFB4 as an autophagy regulator in *Beclin1^{+/+}*; EGFP-p62 iBMKs. **(a)** Proposed mechanism of action of autophagy modulation by PFKFB4. **(b)** Experimental timeline for assessment of PFKFB4-mediated autophagic regulation in the iBMK system. **(c)** Western blot analysis of PFKFB4, LC3 and p62 expression in *Beclin1^{+/+}*; EGFP-p62 iBMKs, 2 days post-transfection with siControl or siPfkfb4. Actin is used as loading control. Representative phase contrast images and viability graphs of *Beclin1^{+/+}*; EGFP-p62 iBMKs 2 days post-transfection with siControl or siPfkfb4. Error bars represent \pm s.d. ($***P < 0.0001$). Scale bar is 50 μ m. **(d)** Representative fluorescence images showing EGFP-p62 accumulation in *Beclin1^{+/+}*; EGFP-p62 iBMKs, 2 days post-transfection with siPfkfb4. Scale bar is 10 μ m. **(e)** Autophagy flux analysis of *Beclin1^{+/+}*; iBMKs stably expressing ptf-LC3 in absence and presence of 3 nM BA1 2 days post-transfection with siControl or siPfkfb4. Representative fluorescence images and corresponding quantitation are provided. Error bars represent \pm s.d. ($***P < 0.0001$). Scale bar is 10 μ m. **(f)** Cell ROX analysis of ROS levels, 2 days post-transfection in the presence or absence of 1 mM NAC. Representative histograms are provided.

PFKFB4 silencing led to increased processing of microtubule-associated protein I (LC3-I) to LC3-II indicative of increased autophagic flux (Figure 4c).

Conventionally, p62 accumulation is associated with autophagy inhibition, whereas conversion of LC3-I to LC3-II is associated with autophagy induction. However, LC3-II can also accumulate because of uncleared autophagosomes secondary to inhibition of or defects in lysosomal function. To more precisely define the role of PFKFB4 in autophagy, we performed flux analysis of *Beclin1^{+/+}*; iBMKs stably expressing tandem-tagged ptf-LC3 (RFP-GFP-LC3). This assay enables the differential identification of autophagosomes (visualized as yellow puncta, the merge of both

red fluorescent protein (RFP) and green fluorescent protein (GFP) fluorescence) and autolysosomes, (visualized by red puncta because of quenching of GFP signal in the acidic environment of the lysosomes) in a single assay.^{84,85} The total number of LC3 puncta (yellow and red) in PFKFB4-depleted cells per field was almost double that of the control cells, indicative of increased autophagy flux (Figure 4e). We confirmed this result with a pharmacologic inhibitor of lysosomal acidification, Bafilomycin A1 (BA1).⁸⁶ In agreement with the tandem-tagged LC3 results, BA1 further increased LC3 puncta upon Pfkfb4 knockdown indicating that the increased number of autophagosomes was due to autophagy induction (Figure 4e).

Having established that PFKFB4 depletion induces autophagy in *Beclin1*^{+/-} iBMKs, we turned our attention to identification of the underlying mechanism. We hypothesized that PFKFB4 depletion could lead to autophagy induction by generating ROS. Therefore, we investigated whether PFKFB4 knockdown altered ROS levels by flow cytometry. Peaks were well separated and the peak for PFKFB4-depleted cells showed a forward shift when compared with control peak indicative of increased ROS levels (Figure 4f). Supplementation of PFKFB4 knockdown cells with 1 mM N-acetyl cysteine (NAC) during transfection, shifted the peak backward, demonstrating scavenging of ROS. This observation is consistent with reports that PFKFB4 knockdown leads to ROS generation in prostate cancer cells.⁷⁸ Taken together, loss of PFKFB4 increases cellular ROS levels, autophagic flux and p62 accumulation.

PFKFB4 inhibits autophagy

To determine if PFKFB4 acts upstream or downstream of the essential autophagy gene *Atg7* in the pathway, knockdown experiments were carried out in an isogenic pair of autophagy-competent and -deficient tumor-derived cell lines (TDCLs) (*K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{+/+} and *K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{-/-}).³ ATG7 converts LC3-I to LC3-II, enabling autophagosome formation. Twenty-four hours post-PFKFB4 knockdown, cells were exposed to ischemic conditions in order to evaluate PFKFB4 function under autophagy-inducing conditions. BA1 was used to assess autophagic flux as depicted in the experimental timeline (Figure 5a). PFKFB4 expression was markedly reduced in TDCLs at both 1 and 2 days post-transfection, while p62 accumulation and increased LC3-II levels were observed 2 days post-transfection (Figures 5b and c). Of note, ischemia led to accumulation of p62 that was further enhanced by PFKFB4 depletion (Figures 5b and c). Moreover, treatment with BA1 led to increased accumulation of p62 in PFKFB4-depleted cells suggestive of enhanced flux through the pathway. Similarly, LC3-I and LC3-II levels were reduced in PFKFB4-depleted cells by day 2, but were rescued by blocking flux with BA1, suggesting that the low levels of LC3 in PFKFB4-depleted cells were due to increased turnover resulting from increased autophagy flux. Under ischemia, PFKFB4 silencing led to increased processing of LC3-I to LC3-II, evidence of increased flux relative to controls. PFKFB4 knockdown in autophagy-deficient TDCLs that lack conversion of LC3-I to LC3-II also increased p62 accumulation in ischemic conditions and had similar accumulation of p62 and LC3-I independent of BA1 treatment (Figure 5c). This suggests that PFKFB4 depletion acts upstream of ATG7 consistent with increased oxidative stress that induces autophagy and p62 upregulation.

Phase contrast images of TDCLs 2 days post-transfection with siPfkfb4 revealed that PFKFB4 depletion impaired cell growth that was further impaired by ischemia, irrespective of *Atg7* status (Figures 5d and e). Importantly, growth was more impaired in *Atg7*^{-/-} than *Atg7*^{+/+} TDCLs, which may be due to greater impairment of redox balance in the absence of autophagy although viability was not significantly altered (Figures 5d and e). To further study the impact of PFKFB4

depletion on autophagy flux in TDCLs, we generated autophagy-competent TDCLs (*K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{+/+}) stably expressing the ptf-LC3 reporter and performed an autophagy flux assay in the presence or absence of BA1. PFKFB4 knockdown significantly increased autophagy flux, confirming the autophagy-suppressive function of PFKFB4 in a second experimental system (Figure 5f).

To directly test our hypothesis that PFKFB4 depletion leads to autophagy induction by generating ROS, we measured ROS levels by flow cytometry. We observed increased basal ROS in autophagy-deficient TDCLs likely due to autophagy deficiency compromising ROS detoxification.⁸⁷ In both autophagy-competent and -deficient TDCLs, PFKFB4 depletion generated ROS that was efficiently scavenged by NAC (Figures 5g and h).

PFKFB4 suppresses autophagy and p62 accumulation by mitigating ROS

PFKFB4 has been reported to be essential for the survival of prostate cancer cells.⁷⁸ Given the physiological relevance of PFKFB4 in prostate cancer, we investigated its role in the negative regulation of autophagy in the human prostate cancer cell line PC3. An experimental timeline for studies in this system is provided (Figure 6a). We first examined the consequences of PFKFB4 depletion on p62 accumulation and LC3-I to LC3-II conversion. Analysis of cell lysates revealed that PFKFB4 knockdown induced p62 and accumulation of both LC3-I and -II (Figure 6b).

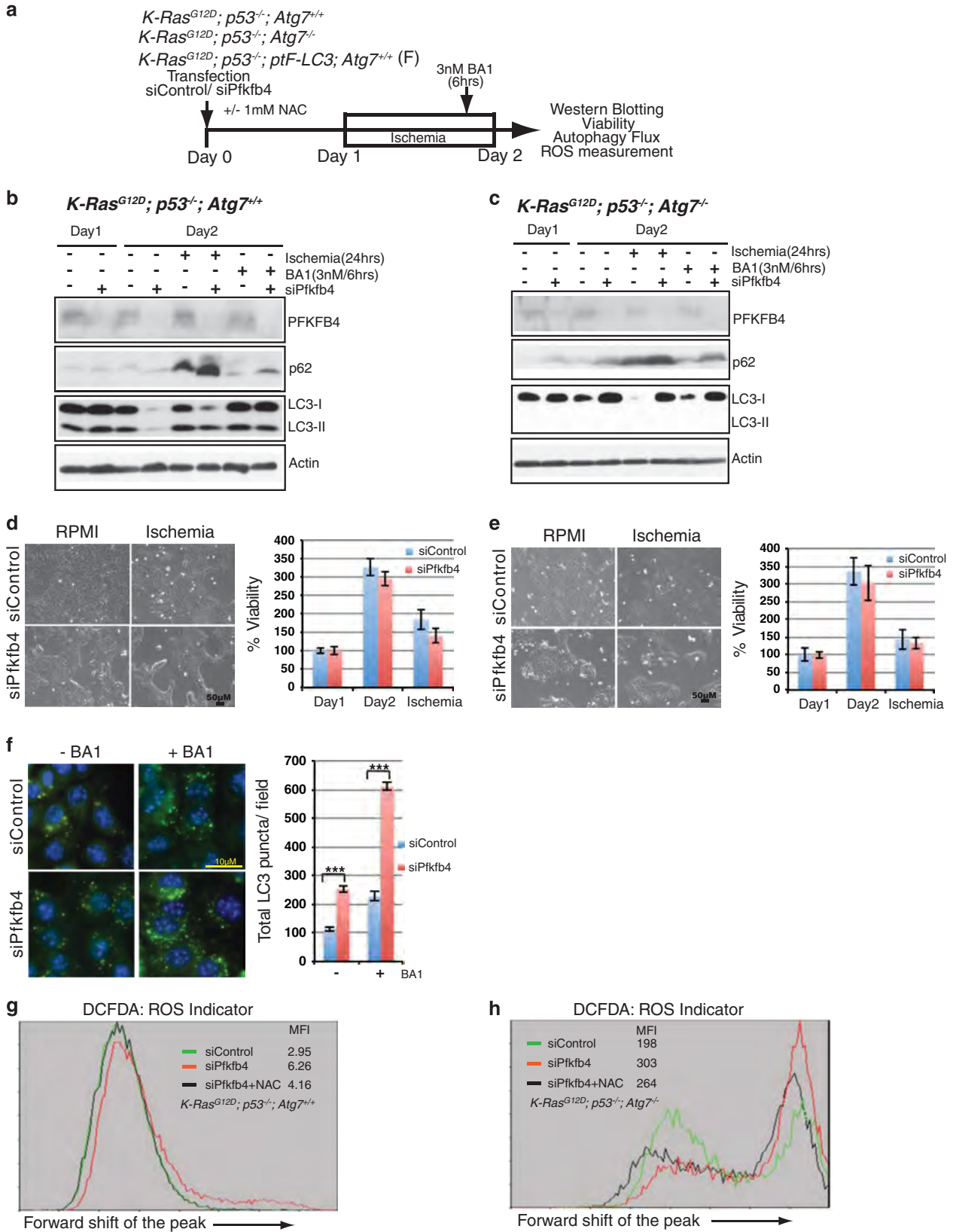
Phase contrast images and viability assays at 3 days post-transfection demonstrated that PFKFB4 silencing significantly impaired cell growth and viability in PC3 cells (Figure 6b), consistent with data from the iBMK cells (Figure 4c) and that of Ros et al.⁷⁸ We next measured autophagy flux in PC3 cells stably expressing the tandem-tagged ptf-LC3 in the absence and presence of BA1 following PFKFB4 knockdown. Consistent with PFKFB4 deficiency promoting autophagy, increased accumulation of p62 and LC3-II were observed by western blot in cells treated with BA1 (Figure 6c). Fluorescence images and quantitation of total LC3 puncta revealed significantly enhanced autophagy flux associated with PFKFB4 silencing (Figure 6c). Importantly, PFKFB4 depletion led to increased ROS levels that could be quenched by NAC supplementation (Figure 6d). Thus, PFKFB4 depletion led to accumulation of p62, stimulation of autophagic flux, ROS generation in PC3 cells.

ROS activation can lead to autophagy induction,⁸⁸⁻⁹⁰ as well as transcriptional upregulation of p62 as an antioxidant defense mechanism.⁹¹ If elevated ROS levels were driving p62 accumulation and autophagic flux in the PFKFB4-depleted cells, we postulated that the enhanced p62 accumulation and autophagy flux following PFKFB4 depletion should be rescued by treatment with NAC. To directly test this hypothesis, autophagy flux was analyzed in tandem-tagged LC3-expressing PC3 cells following PFKFB4 knockdown in the presence or absence of NAC. Consistent with our hypothesis, NAC supplementation at the time of transfection significantly decreased both p62 levels and

Figure 5. PFKFB4 functions as an autophagy inhibitor in K-Ras-driven TDCLs. **(a)** Experimental timeline for assessment of PFKFB4-mediated autophagic regulation in K-Ras-driven TDCLs. **(b,c)** Western blot analysis of PFKFB4, p62, and LC3 in *K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{+/+} **(b)** and *K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{-/-} **(c)** cell lines grown in normal media conditions or subjected to 24-h ischemia, 1 and 2 days post-transfection with siControl or siPfkfb4. Cells were incubated in the presence or absence of 3 nM BA1 as indicated. **(d,e)** Representative phase contrast images and associated viability graphs of autophagy-competent *K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{+/+} **(d)** and autophagy-deficient *K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{-/-} **(e)** cells in normal media or ischemic conditions, 2 days post-transfection with siControl or siPfkfb4. Scale bar is 50 μm. **(f)** Autophagy flux analysis of *K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{+/+} stably expressing ptf-LC3 in absence and presence of 3 nM BA1, 2 days post-transfection with siControl or siPfkfb4. Representative fluorescence images and corresponding quantitation are provided. Error bars represent ± standard deviation (****P* < 0.0001). Scale bar is 10 μm. **(g,h)** DCFDA analysis of ROS levels in *K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{+/+} **(g)** and *K-Ras*^{G12D/+}; *p53*^{-/-}; *Atg7*^{-/-} **(h)** 2 days post-transfection with siPfkfb4 in the presence or absence of 1 mM NAC. Representative histograms are provided.

autophagy flux (Figure 6e). Western blot analysis of PC3 cells revealed a profound decrease in p62 and LC3 levels upon NAC supplementation in PFKFB4-depleted cells. Importantly, NAC

supplementation led to a significant decrease in autophagy flux as evident by the quantitation of LC3 puncta in PFKFB4-deficient cells (Figure 6e). Taken together, our data demonstrate that



PFKFB4 inhibits autophagy by mitigating ROS levels, which prevents induction of p62 because of oxidative stress (Figure 6f).

DISCUSSION

Using a novel cell-based assay that measures the ability of cells to eliminate the autophagy substrate p62 and loss-of-function screening, we have identified candidate genes whose expression regulates autophagy either positively or negatively. A unique feature of this screen is the use of accumulation of an autophagy substrate as a readout of the culmination of autophagy rather than the number of LC3 puncta to monitor autophagy initiation.^{56–62} This approach enabled us to identify autophagy regulators throughout the autophagy pathway, as well as modulators of p62 itself.

Our screen used exogenously expressed p62 as a reporter. We demonstrated that autophagy-competent *beclin1*^{+/+} cells efficiently eliminate accumulated EGFP-p62 during recovery in full media but this clearance is blocked in autophagy impaired *beclin1*^{+/-} derived cell lines (Figures 1b and c). It is this differential elimination of p62 that is the basis of the screen. Importantly, we have previously published similar differential clearance of endogenous p62 for this cell line (as well as in *atg5*^{-/-} derived iBMKs) indicating that our overexpression approach faithfully recapitulates the failed elimination of p62 in the autophagy-deficient setting.¹⁵ In agreement with this work, the Mizushima lab has recently shown that endogenous and exogenously driven p62 levels decrease similarly during productive autophagy.⁹²

Another unique aspect of this work is the focus on cellular GTPases and vesicle trafficking components as potential autophagy regulators. Targeting the vesicle trafficking machinery may be a means to limit kinase activity or complex formation essential for autophagy initiation. It is well known that spatio-temporal dynamics are involved at multiple levels within the autophagy cargo loading, and vesicle fusion and docking processes. This work extends our understanding of the machinery involved in autophagosome maturation and fusion, as well as regulation of the pathway by the protein kinome.

Our candidate gene lists identify potential drug targets for autophagy-modulating therapies. As both activation and suppression of the pathway are therapeutically desirable outcomes, this work is an important resource. A small subset of autophagy regulators identified here scored in recent genome-wide screens using GFP-LC3 reporters. Importantly, these screens differed in their method of silencing and library source (Dharmacon siGENOME) from this assay, as well as autophagy stimulus: one screen measured regulation of basal autophagy in normal growth conditions in the H4 neuronal cell line,⁵⁶ another assayed autophagy induced by Sinbis virus infection in HeLa cells,⁵⁸ whereas a third group identified regulators of starvation-induced autophagy in human embryonic kidney 293T cells.⁶⁴ Although there are many reasons candidate genes may not appear in various screens, attention should be paid to those regulators that

emerge irrespective of cell type, mechanism of silencing, reporter system or the cellular stress used to activate the pathway. These genes include *Epha6*, *Grk6*, *Gtbbp4*, *Irak3*, *Pldn*, *Prkaa2*, *Prckz*, *Tpr*, *Shpk*, *Tlk2*, *Rab7*, *Nme2* and *Eif2ak1*.

In this work, we identify PFKFB4 as a novel autophagy regulator. PFKFB4 depletion impairs the growth and survival of cancer cells, enhances ROS levels and profoundly induces autophagy. We demonstrate that the induction of autophagy by PFKFB4 depletion is the consequence of ROS generation, as scavenging ROS in PFKFB4-depleted cells prevented autophagy induction. PFKFB4 depletion in multiple human prostate cancer cell lines has been reported to increase ROS levels by decreasing flux through the pentose phosphate pathway, which compromises NADPH production essential for scavenging of intracellular ROS.⁷⁸ Interestingly, PFKFB4 silencing led to p62 accumulation despite inducing autophagic flux. These seemingly paradoxical findings can be explained by induction of p62 from oxidative stress via Nrf2 binding to the antioxidant response element containing cis-element of p62;⁹¹ supported by our observation that the addition of NAC to PC3 cells prevented accumulation of endogenous p62 in PFKFB4-depleted cells (Figure 6e). p62 contributes to the activation of NRF2 target genes in response to oxidative stress via binding and inactivating the negative regulator of NRF2, KEAP1.⁹¹

Of note, *Pfkfb4* also scored on our negative regulator list. However, a July 2014 query of the TRC library revealed that the two hairpins, which scored as negative regulators of autophagy and are predicted to have a 100% match to *Pfkfb4*, are also predicted to have an 80% match to the well-known autophagy inhibitor *Rheb*. Thus, it is possible that the ranking of PFKFB4 as a negative regulator may be attributed to off-target effects of these hairpins on *Rheb*, a critical component of the mTOR signaling cascade. However, as RHEB levels were not interrogated and these hairpins were predicted to have significant efficacy against *Pfkfb4* itself, we did not remove this gene from Table 2. Importantly, the hairpins targeting *Pfkfb4* that resulted in p62 accumulation in our screen are not predicted to target *Rheb*. Critically, a second system, Dharmacon siGENOME Smartpools (made of four distinct small interfering RNAs targeting *Pfkfb4*) supports our finding that silencing *Pfkfb4* results in robust accumulation of p62. Further, we note that the increased accumulation of p62 observed post-silencing was associated with the well-documented decrease in viability reported with loss of PFKFB4.

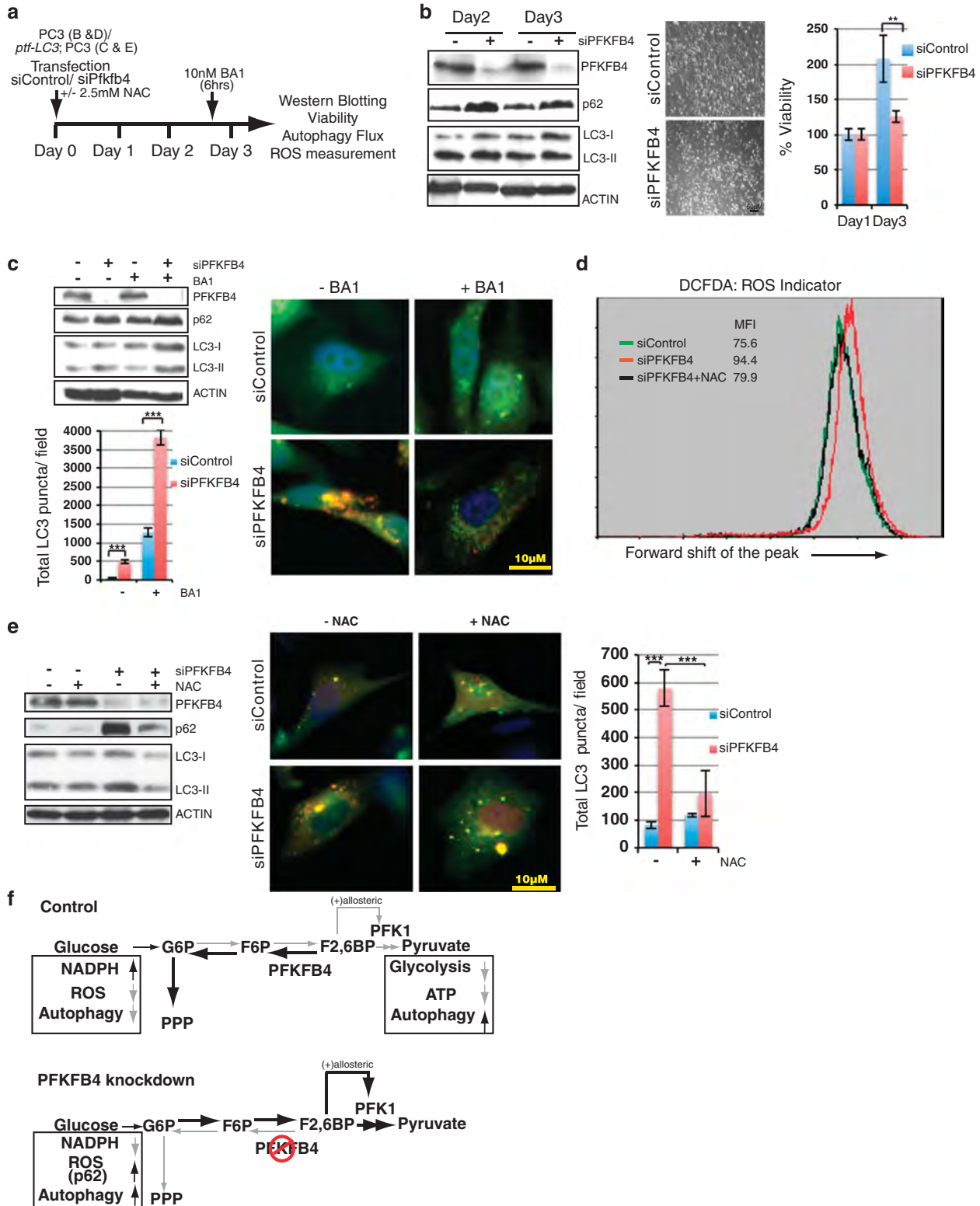
Although this study is the first to describe PFKFB4 as an autophagy regulator, a related family member, PFKFB3 has been linked to the pathway. In rheumatoid arthritis T lymphocytes, PFKFB3 acts as an autophagy inducer,^{93,94} yet in colon adenocarcinoma cells it promotes glucose uptake that suppresses autophagy by activating mTOR.⁹⁵ TIGAR, a p53-inducible protein with fructose-2,6-bisphosphatase enzymatic activity identical to PFKFB4, has a similar role in suppression of cellular ROS levels and also suppresses autophagy.^{96,97} Thus, it is clear that these key metabolic enzymes tightly regulate autophagy via multiple

Figure 6. PFKFB4 mitigates ROS to suppress autophagy and p62 accumulation in PC3 cells. **(a)** Experimental timeline for assessment of PFKFB4-mediated autophagy regulation in PC3 cells. **(b)** Western blot analysis of PFKFB4, p62 and LC3 in PC3 cells, 2 and 3 days post-transfection with siControl or siPFKFB4. Representative phase contrast images and viability graphs of PC3 cells 3 days post-transfection with siControl or siPFKFB4 are provided. Error bars represent \pm s.d. (** $P < 0.005$). Scale bar is 50 μ m. **(c)** Autophagy flux analysis of PC3 cells stably expressing ptf-LC3, 3 days post-transfection with siControl or siPFKFB4 in the presence or absence of BA1. Representative western blots of PFKFB4, p62, LC3, fluorescence images and corresponding quantitation are provided. Scale bar is 10 μ m. **(d)** DCFDA analysis of ROS levels in PC3 cells 3 days post-transfection with siControl or siPFKFB4 in the presence or absence of 2.5 mM NAC as indicated. Representative histograms are provided. **(e)** Autophagy flux analysis of PC3 cells stably expressing ptf-LC3 3 days post-transfection with siControl or siPFKFB4, in the presence or absence 2.5 mM NAC. Representative western blots of PFKFB4, p62, LC3, fluorescence images and quantitation graph are provided. Error bars represent \pm s.d. (*** $P < 0.0001$). Scale bar is 10 μ m. **(f)** Model of PFKFB4-mediated regulation of autophagy by controlling redox balance of the cell.

mechanisms in a context-dependent manner. Additional knowledge gained about PFKFB4 will be of value to the scientific community as PFKFB3 and PFKFB4 are emerging therapeutic targets for cancer.^{79,81,95}

In this work, we have shown that PFKFB4 suppresses autophagy and p62 by mitigating oxidative stress. Although it is known

that PFKFB4 promotes pentose phosphate pathway flux and NADPH production critical for antioxidant defense, our work establishes for the first time that this activity also suppresses p62 induction and autophagy. Importantly, loss of PFKFB4 promotes oxidative stress, triggering autophagy as a stress-adaptive survival mechanism.



MATERIALS AND METHODS

Cell lines and reagents

Isogenic *Beclin1^{+/+}* and *Beclin1^{+/-}* iBMK cell lines expressing EGFP-p62 were established¹³ and maintained in Dulbecco's modified Eagle's medium (Invitrogen, Grand Island, NY, USA) containing 10% fetal bovine serum and 1% penicillin–streptomycin (Invitrogen). *K-Ras^{G12D}*; *p53^{-/-}*; *Atg7^{+/+}* and *Atg7^{-/-}* TDCLs generated from 8-week-old lung tumors³ were cultured in RPMI-1640 medium containing 10% fetal bovine serum, 1% penicillin–streptomycin (Invitrogen) and 1 mM sodium bicarbonate (Invitrogen). The human prostate cancer cell line PC3 (ATCC, Manassas, VA, USA; CRL-1435) was maintained in RPMI-1640 medium containing 10% fetal bovine serum, 1% penicillin–streptomycin (Invitrogen). Glucose-free RPMI-1640 or Dulbecco's modified Eagle's medium for ischemia were obtained from Invitrogen.

The following reagents were used according to the manufacturer's instructions: Bafilomycin A1 (Sigma-Aldrich, St Louis, MO, USA; B1793) CellRox Deep Red Reagent (Invitrogen, C10422), NAC (Sigma-Aldrich, A9165), Prolong-Anti-Fade Gold mounting media (Invitrogen, P10144), Hoechst 33342 (Invitrogen, H1399), Puromycin (Sigma-Aldrich, P8833) and chloromethyl derivative of 2',7'-dichlorodihydrofluorescein diacetate (CM-H₂DCFDA, Invitrogen, C6827).

Antibodies

The following antibodies were used: LC3 (Novus Biologicals, Littleton, CO, USA; NB600-1384), p62⁽⁸⁾, actin (Sigma, St Louis, MO, USA; A1978), Atg7 (Sigma, A2856), AMPKα (Cell Signaling, Danvers, MA, USA; 2532), Etnk1 (Santa Cruz Biotechnology, Dallas, TX, USA; sc-13-754), Hrs (Santa Cruz Biotechnology, sc-23791), IKKβ (Cell Signaling, 2370), mTOR (Cell Signaling, 2972), Rab7 (Cell Signaling, 2094), ULK1 (Sigma, A7481). A rabbit polyclonal antibody against PFKFB4 was developed in conjunction with Cocalico Biologicals (Reamstown, PA, USA) (details in Supplemental Methods).

Autophagic flux assay

Cells stably expressing ptf-LC3 were analyzed for autophagic flux under normal and PFKFB4-depleted conditions. Three fields (containing a minimum of 50 cells) were scored for total number of puncta (red and yellow) per condition. Nuclei were counterstained with 4,6-diamidino-2-phenylindole. Wherein indicated, BA1 was included as a positive control for flux blockade (3 nM for iBMKs and TDCLs and 10 nM for PC3).

ROS measurements by flow cytometry

In all, 5 μM CellRox or 10 μM CM-H₂DCFDA were added to cells (in Hank's balanced salt solution for DCFDA) for 30 min at 37 °C followed by measurement of mean fluorescence by flow cytometry (FC500; Beckman Coulter, Indianapolis, IN, USA). Three independent experiments were performed analyzing 10 000 (iBMK and PC3 cells) and 5000 (*K-Ras*-TDCLs) cells in each experiment. NAC rescue was performed at 1 mM (iBMKs and TDCLs) or 2.5 mM (PC3 cell line) final concentration.

CONFLICT OF INTEREST

EW is a member of the scientific advisory board of Forma Therapeutics. EW, DMS, RP and AMS are co-inventors on a related patent application. RTA is an employee and shareholder of Pfizer, Inc.

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ABBREVIATIONS

DCFDA, 2',7'-dichlorodihydrofluorescein diacetate; AMPK, 5' adenosine monophosphate-activated protein kinase; BA1, Bafilomycin A1; ESCRT, endosomal sorting complexes required for transport; F6P, fructose-6-phosphate; F1,6BP, fructose-1,6-bisphosphate; F2,6BP, fructose-2,6-bisphosphate; LC3, microtubule-associated protein 1 light chain 3; LIR, LC3 interaction region; mTOR, mammalian/mechanistic target of rapamycin; NAC, N-acetyl-cysteine; NSCLC, non-small cell lung cancer; PFK-1, phosphofructokinase-1; PFKFB4, 6-phosphofructo-2-kinase/fructose-2,6-bisphosphatase 4; PPP, pentose phosphate pathway; ROS, reactive oxygen species; TDCLs, tumor-derived cell lines; UBA, ubiquitin-binding domain

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