Apoptotic Cells Activate AMP-activated Protein Kinase (AMPK) and Inhibit Epithelial Cell Growth without Change in Intracellular Energy Stores*

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Background: Nonprofessional phagocytes, like epithelial cells, recognize apoptotic cells.

Results: Apoptotic cells mimic the effects of intracellular energy depletion and inhibit the growth (cell size) of epithelial cells with which they interact.

Conclusion: Apoptotic cells activate AMP-activated protein kinase (AMPK) and inhibit cell growth.

Significance: By acting as sentinels of environmental stress, apoptotic targets enable nearby cells to monitor and adapt to local change.

Apoptosis plays an indispensable role in the maintenance and development of tissues. We have shown that receptor-mediated recognition of apoptotic target cells by viable kidney proximal tubular epithelial cells (PTECs) inhibits the proliferation and survival of PTECs. Here, we examined the effect of apoptotic targets on PTEC cell growth (cell size during G1 phase of the cell cycle). Using a cell culture model, we show that apoptotic cells potently activate AMP-activated protein kinase (AMPK), a highly sensitive sensor of intracellular energy stores. AMPK activation leads to decreased activity of its downstream target, ribosomal protein p70 S6 kinase (p70S6K), and concomitant inhibition of cell growth. Importantly, these events occur without detectable change in intracellular levels of AMP, ADP, or ATP. Inhibition of AMPK, either pharmacologically by compound C or molecularly by shRNA, diminishes the effects of apoptotic targets and largely restores p70S6K activity and cell size to normal levels. Apoptotic targets also inhibit Akt, a second signaling pathway regulating cell growth. Expression of a constitutively active Akt construct partially relieved cell growth inhibition but was less effective than inhibition of AMPK. Inhibition of cell growth by apoptotic targets is dependent on physical interaction between apoptotic targets and PTECs but independent of phagocytosis. We conclude that receptor-mediated recognition of apoptotic targets mimics the effects of intracellular energy depletion, activating AMPK and inhibiting cell growth. By acting as sentinels of environmental change, apoptotic death may enable nearby viable cells, especially nonmigratory epithelial cells, to monitor and adapt to local stresses.

Apoptosis plays an indispensable role in the maintenance and development of tissues. At the most straightforward level, apoptosis provides a means for the rapid and efficient removal of aged, damaged, or excess cells without harm to surrounding tissues (1, 2). In addition to this essentially passive activity, apoptosis also contributes in a more dynamic manner to tissue homeostasis. Cells dying by apoptosis acquire multiple new activities, both secreted and cell-associated, that allow them to modulate the function of nearby live cells (3–9). Although earlier studies focused on the ability of apoptotic cells to suppress inflammation (10–15), apoptotic cells also affect a broad range of cellular functions, including such vital activities as survival (3, 8, 9), proliferation (3, 8, 9), differentiation (16), and migration (17). Moreover, these effects are not limited to professional phagocytes, like macrophages (md), but extend to virtually all...
cell types and lineages, including traditionally nonphagocytic cells, such as epithelial and endothelial cells (6, 8, 9, 17, 18).

Importantly, the specific effects elicited in nearby viable cells following exposure to apoptotic cells depend on multiple factors relating both to the viable responding cells and to the apoptotic cell itself. For example, although the responses by murine mφ and kidney proximal tubular epithelial cells (PTECs) are similar with respect to proliferation, with inhibition of proliferation occurring in both cases, the responses of these two cells differ with respect to survival (3, 5, 8, 9). Apoptotic cells promote mφ survival, whereas they induce the apoptotic death of PTECs (3, 5, 8, 9). Even for responding cells of the same lineage, responses can differ depending on their organ of origin (e.g. PTECs versus mammary epithelial cells) (9) or state of activation (e.g. neutrophils) (20). Conversely, apoptotic cells may evoke different responses in the same cell depending on the nature of the apoptotic stimulus (9, 21) or the time elapsed from administration of the apoptotic stimulus to interaction between apoptotic and responding cells (9, 13).

In light of this complexity, it is interesting to speculate about the in vivo consequences of a local increase in apoptotic death. As we have previously hypothesized (9, 22), such an increase may serve as a signal of environmental change or stress. A tissue’s overall response, comprising the integrated responses of its component cells, may then represent an attempt at adaptation. A physiologically relevant example would be the vasoconstriction or partial occlusion of an artery supplying a segment of an organ such as the kidney. This will lead to a reduced delivery of oxygen and nutrients. Within the affected zone, the response of individual viable cells to nearby dead or dying target cells will depend on the responding cell’s lineage and anatomic location, among other factors, both intrinsic and extrinsic. Some cells, such as infiltrating mφ, will demonstrate increased survival, reflecting their importance in clearance of debris and repair (3, 5). Other cells, in contrast, such as kidney PTECs, will evince decreased proliferation and survival, reflecting the need to decrease metabolic demand in the face of reduced supply (8, 9).

AMP-activated protein kinase (AMPK) is a highly sensitive sensor of intracellular energy stores (23, 24). Activation of AMPK occurs primarily as a result of an increase in the ratio of either AMP or ADP to ATP (23, 25). Upon activation, AMPK acts as a metabolic switch with profound effects on intermediary cell metabolism. The net outcome is the augmentation or conservation of intracellular energy stores, through promotion of ATP production, inhibition of ATP consumption, and facilitated cellular uptake of nutrients (23, 24). A major downstream target of AMPK is the mammalian target of rapamycin complex 1 (mTORC1), a kinase critical for cell growth (increase of cell mass) and proliferation (increase of cell number) (23, 24, 26–29). Inhibition of mTORC1 by AMPK leads to inhibition of cell growth, and thereby cell size, by preventing mTORC1-mediated phosphorylation and activation of the ribosomal proteins p70 S6 kinases 1 and 2 (p70S6K) (27–29).

Here, using a cell culture model, we test the hypothesis that exposure of murine kidney PTECs to apoptotic target cells acts as an extracellular stress, mimicking the effects of intracellular depletion of energy stores. We show that apoptotic targets potently activate AMPK, leading to decreased activity of p70S6K and concomitant inhibition of cell growth. Importantly, these events occur without detectable change in intracellular energy stores. Inhibition of AMPK, either pharmacologically by compound C or molecularly by shRNA, diminishes the effects of apoptotic targets and largely restores p70S6K activity and cell size to normal levels. Together with our previous results, our data reveal that apoptotic cells inhibit the growth and proliferation of nearby PTEC responders. By acting as sentinels of environmental change, apoptotic death may allow nearby viable cells, especially nonmigratory epithelial cells, to monitor and adapt to local stresses.

**Experimental Procedures**

**Materials**—Unless otherwise stated, all chemicals were obtained from Sigma, Invitrogen, or Fisher. Cell culture medium was obtained from Mediatech (Herndon, VA).

**Antibodies**—Affinity-purified polyclonal rabbit antibodies detecting the active Thr172-phosphorylated forms of α1-AMPK and α2-AMPK, the Thr308-phosphorylated form of Akt, the Ser173-phosphorylated form of Akt, total p70S6K1 and p70S6K2, the active Thr389-phosphorylated form of p70S6K1 and p70S6K2, the active Ser240/244-phosphorylated form of S6 ribosomal protein, the active Thr34-phosphorylated form of FoxO1, the active Thr32-phosphorylated form of FoxO3a, the inactive Ser21-phosphorylated form of glycogen synthase kinase (GSK) 3α, the inactive Ser9-phosphorylated form of GSK3β, and total β-actin were obtained from Cell Signaling Technology (Beverly, MA). Rabbit mAb (57C12) detecting total β1-AMPK and β2-AMPK and rabbit mAb (14C10) detecting total GAPDH were obtained from Cell Signaling Technology. Horseradish peroxidase–linked donkey anti–rabbit F(ab’)2 from GE Healthcare was used as a secondary antibody for detection of Western blots by enhanced chemiluminescence.

**Cell Culture**—All cells were grown at 37 °C in a humidified 5% (v/v) CO2 atmosphere unless otherwise stated. The conditionally immortalized mouse kidney PTEC cell line (BU.MPT, Boston University mouse proximal tubule) was maintained in high-glucose DMEM containing 10% (v/v) heat-inactivated FBS, 2 mmL-glutamine, 10 mm HEPES, 100 units/ml penicillin/streptomycin, and 10 units/ml IFN-γ. BU.MPT cells were derived from a transgenic mouse bearing a temperature-sensitive mutation (tsA58) of the SV40 large tumor antigen (TAg) under the control of the mouse MHC H-2Kb class I promoter (30, 31). Under permissive conditions, defined as growth at 33–37 °C in the presence of IFN-γ, the tsA58 TAg transgene is expressed. Under nonpermissive temperatures, defined as growth at 39.5 °C in the absence of IFN-γ, expression of the tsA58 TAg transgene is inhibited (by >95%) and BU.MPT cells behave like primary cultures of mouse kidney PTECs. Prior to all experiments, BU.MPT cells were serum-starved and cultured under nonpermissive conditions for 24 h. DO11.10 cells, a T cell hybridoma line, were grown in RPMI 1640 medium supplemented with 10% (v/v) heat-inactivated FBS, 100 units/ml penicillin/streptomycin, and 50 μM 2-mercaptoethanol.

**Preparation of Apoptotic and Necrotic Cell Targets**—Apoptosis of BU.MPT cells was induced by incubating cells in FBS-free medium containing the nonselective protein kinase inhibitor
Apopotic Cells Activate AMPK and Inhibit Cell Growth

Staurosporine (1 μg/ml, 3 h). Apoptosis of DO11.10 cells was induced by incubating cells in FBS-containing medium containing the macromolecular synthesis inhibitor actinomycin D (200 ng/ml, overnight). After induction of apoptosis, the remaining adherent cells were detached by addition of 5 mM EDTA and pooled with floating cells, followed by three washes and resuspension in fresh FBS-free medium before use in experiments. For induction of necrosis, cells were first detached with 5 mM EDTA and suspended in the appropriate FBS-free medium. Necrosis was then induced by heating cells to 70 °C for 45 min, followed by incubation at 37 °C for 2 h. Apoptotic targets were added to responder cells either directly or after fixation for 30 min with 0.4% (v/v) paraformaldehyde in PBS, with similar results. Necrotic targets were always added directly without fixation.

Induction of apoptosis or necrosis was confirmed by flow cytometry. Early apoptotic cells (intact cell membranes) were defined as PI-negative cells with annexin V staining and decreased cell size. Necrotic cells were defined as PI-positive cells of normal or increased cell size. Late apoptotic cells (non-intact cell membranes) were defined as PI-positive cells with annexin V staining and decreased cell size. Loss of membrane integrity by necrotic cells was confirmed by trypan blue staining. By these criteria, apoptotic target preparations contained ~85% early apoptotic and ~15% late apoptotic cells. Necrotic target preparations contained ~95% necrotic cells. In all preparations, viable cells, defined as PI-negative cells of normal size without annexin V staining, composed <5% of the total cell population.

Retroviral Transfection with Akt Constructs—BU.MPT cells were infected with retroviral vectors containing either GFP alone (pBabe-GFP) or GFP plus a constitutively active (myristoylated) Akt construct (pBabe-GFP-mAkt), both of which were kind gifts of Dr. Nissim Hay (University of Illinois at Chicago). Retroviruses were generated by transient transfection of 293T cells, followed by harvesting of the retrovirus-containing culture medium. Myristoylation of Akt leads to its localization at the plasma membrane, followed by harvesting of the retrovirus-containing culture medium. Myristoylation of Akt leads to its localization at the cell membrane where Akt can be activated via phosphorylation at Thr308 by the upstream kinase 3-phosphoinositide-dependent protein kinase 1 (PDK1). All downstream signaling events, both cytoplasmic and nuclear, dependent on phosphorylation of Akt at Thr308 are replicated by this constitutively active Akt construct (32, 33). Retroviral infection of BU.MPT cells was performed in the presence of 8 μg/ml Polybrene for 24 h. The pBabe-GFP and pBabe-GFP-mAkt retroviral vectors, as well as the protocols using them, have been previously described (32, 33).

shRNA—BU.MPT cells were transduced with control scrambled or specific anti-β1-AMPK shRNA constructs cloned into the pGIPZ lentiviral vector using the trans-lentiviral shRNA packaging kit (GE Healthcare). Specific anti-β1-AMPK shRNA (TGATGATACAGTTTCAATA) targeted the β1 isoform of AMPK; control scrambled shRNA (ATCTCGCTTGGGC-GAGAAT) was nonsilencing. The pGIPZ lentiviral vector contains a puromycin resistance gene used to select successfully transduced cells.

Transduction of BU.MPT cells and passaging of transduced cells were performed under permissive conditions (37 °C in the presence of IFN-γ). BU.MPT cells were seeded into 24-well tissue culture plates at a density of 5 × 10^4 cells/well. The following day, 250 μl of culture medium, containing Polybrene (8 μg/ml) and 1.8 × 10^7 transduction units of lentiviral particles, was added to each well, according to the manufacturer’s instructions. After incubation for 6 h, 1 ml of culture medium was added per well, and cells were incubated for an additional 72 h. Cells were then passaged into 60-mm dishes containing 5 ml of culture medium plus puromycin (1 μg/ml) to select BU.MPT cells successfully transduced with lentiviral constructs.

To confirm knockdown of β1-AMPK, cells surviving 12 days of puromycin selection were subjected to immunoblotting. As compared with BU.MPT cells transduced with control shRNA, β1-AMPK expression was reduced by ~75% in cells containing specific shRNA (see Fig. 5A). The β2-AMPK isoform, not knocked down by β1-AMPK shRNA, constituted <20% of total AMPK in control and nontransduced BU.MPT cells. Stably transduced BU.MPT cells were maintained in the presence of puromycin (1 μg/ml). Prior to all experiments, cells were cultured for 24 h under nonpermissive conditions in the absence of puromycin.

Western Blot Analysis—After stimulation of BU.MPT responder cells with apoptotic or necrotic targets, in the presence or absence of EGF (10 nM) (Calbiochem), responders were washed three times with ice-cold PBS and then lysed in ice-cold cell lysis buffer (TBS containing 10 mM sodium pyrophosphate, 0.5% w/v deoxycholate, 0.1% w/v SDS, 10% glycerol, 25 mM sodium fluoride, 10% Triton X-100, 1 mM DTT, 1 mM PMSF, 10 mM sodium orthovanadate with added Complete® mini EDTA-free anti-protease mixture tablet (Roche Applied Science)). Lysates were sonicated on ice with 10 pulses of 20 Hz and then centrifuged at 20,000 × g for 10 min at 4 °C. Supernatants were stored at −70 °C.

Protein concentrations of sample for gel electrophoresis were determined by the bicinchoninic acid protein assay (Pierce). Samples were then boiled in 6× reducing sample buffer for 5 min at 95 °C, and 20 μg per sample were loaded onto 4–15% Mini-PROTEAN TGX precast gels (Bio-Rad), electrophoresed at 120 V at constant voltage, and wet transferred on a Genie® blotter (Idea Scientific, Minneapolis, MN) at 12 V constant voltage to Immobilon-P (0.45 μm) PVDF membranes (Millipore, Billerica, MA). Membranes were blocked with 5% w/v dry milk in TBS plus 0.1% (v/v) Tween 20, before probing with one of the primary antibodies described above. Following incubation with secondary antibody, immunoreactive bands were visualized by the luminol reaction (GE Healthcare). Equivalent loading of protein samples was monitored by staining with Ponceau S (0.25% w/v) in 0.1% v/v acetic acid for 5 min and/or detection of GAPDH or β-actin.

Flow Cytometric Analysis of Cell Growth—Cell growth was assessed by a modification of previously described methods (27). In brief, BU.MPT responder cells were exposed to apoptotic or necrotic targets for the indicated times and then rinsed twice with PBS. After overnight culture, cells were stained with DAPI (1 μg/ml) for 30 min at 37 °C, harvested by trypsinization, and resuspended in Ca^{2+} - and Mg^{2+}-free PBS containing 2 mM...
EDTA and 10% (v/v) heat-inactivated FBS. Single-cell suspensions were run on either a BD LSRSorter flow cytometer with BD FACSDiva™ acquisition software (BD Biosciences) or a CyAn™ ADP flow cytometer with Summit™ acquisition software (Beckman Coulter, Inc., Fullerton, CA). Cells were gated to exclude cellular aggregates. Gated cells were assessed for DNA content, and the relative size of cells in G1 phase of the cell cycle was determined by measuring forward scatter. Unstained cells, both responders and targets, were used to compensate for spectral spillover. Analysis was performed on either fixed (2% paraformaldehyde for 20 min) or live unfixed responder cells with similar results.

Measurement of Intracellular AMP, ADP, and ATP—Intracellular content of AMP, ADP, and ATP was measured by modification of previously described methods (34, 35). In brief, after rinsing with ice-cold saline (phosphate-free), cells were scraped and centrifuged at 5000 × g for 4 min at 4 °C. After addition of 300 μl of acidic acetonitrile solvent (80:20 v/v acetonitrile in HPLC water containing 0.1 M formic acid), the pellet was incubated on ice for 15 min and then centrifuged at 20,000 × g for 5 min at 4 °C. The supernatant was separated, and an additional 200 μl of acidic acetonitrile solvent was added to the pellet, followed again by centrifugation at 20,000 × g for 5 min at 4 °C. This step was repeated a third time, and the supernatants were combined to a final volume of 700 μl.

Extracts were then analyzed with the 5500 QTRAP liquid chromatography-tandem mass spectrometry (LC/MS/MS) system (AB Sciex, Foster City, CA) with a 1200 series HPLC system (Agilent Technologies, Santa Clara, CA), including a degasser, an autosampler, and a binary pump. The LC separation was performed on a Phenomenex 5 μm C18(2) column (4.6 × 250 mm, 5 μm) with mobile phase A (25 mM ammonia acetate in water) and mobile phase B (methanol). The flow rate was 0.55 ml/min. The linear gradient was as follows: 0–1 min, 95% phase A; 1–6.5 min, 90% phase A; 6.5–10.5 min, 1% phase A; 10.6–19 min, 95% phase A. The autosampler was set at 5 °C. The injection volume was 10 μl. Mass spectra were acquired under negative electrospray ionization with an ion spray voltage of −4500 V. The source temperature was 350 °C. The values for the curtain gas, ion source gas 1, and ion source gas 2 were 35, 65, and 55 pounds per square inch, respectively. Multiple reaction monitoring was used for quantitation as follows: ATP mass-to-charge ratio (m/z) 506.1 → m/z 79.0; ADP m/z 426.1 → m/z 79.0; and AMP m/z 346.1 → m/z 79.0. The internal standard 2-chloro-ATP was monitored at m/z 540.0 → m/z 79.0.

Statistics—Data are expressed as mean ± S.E. of the averaged values obtained from each experiment. Statistical significance was determined by a two-tailed Student’s t test.

Results

Apoptotic Targets Inhibit the Growth of Live BU.MPT Responder Cells—We have shown previously that receptor-mediated recognition of apoptotic targets induces the apoptotic death and inhibits the proliferation of viable BU.MPT epithelial cell responder cells (8, 9). Both activities lead to a decrease in the magnitude of the responder cell population, thereby reducing the population’s metabolic demand. If the surge in the number of apoptotic targets is the consequence of a diminished metabolic supply, as for example from ischemia, then these two target-induced activities would tend to restore the balance between supply and demand. As cells can also reduce their metabolic demand by decreasing protein synthesis and other energy-consuming anabolic pathways, we determined the effect of apoptotic targets on cell size during the G1 phase of the cell cycle, a measure of cell mass or growth (26–29).

BU.MPT responders were exposed to apoptotic or necrotic targets for 2 h (Fig. 1). After 24 h, the relative size of responders in the G1 phase of the cell cycle was assessed by flow cytometric comparison of forward scatter (27). Apoptotic targets significantly decreased the relative size of BU.MPT responders to 64.3 ± 7.3% that of untreated responders (p < 0.02) (Fig. 1B). In contrast, necrotic targets lacked a significant effect on the relative size of BU.MPT responders (90.5 ± 5.6%; p = not significant). Although the primary source of apoptotic targets in these studies was murine DO11.10 T cells induced to undergo apoptosis by treatment with actinomycin D, similar results were obtained with BU.MPT targets induced to undergo apoptosis by treatment with staurosporine (data not shown).

The major controller of cell growth is the mTORC1-regulated ribosomal kinase, p70S6K (27–29). We therefore determined the effect of apoptotic targets on both basal and EGF-induced p70S6K phosphorylation and activity (Fig. 2A). Exposure of quiescent BU.MPT responders to apoptotic targets for 15 min strongly inhibited basal phosphorylation of p70S6K. Similarly, prior exposure to apoptotic targets strongly inhibited EGF-induced phosphorylation of p70S6K. In both cases, inhibition of p70S6K occurred in a dose-dependent manner. Inhibition of p70S6K phosphorylation correlated with inhibition of p70S6K activity, as detected by parallel inhibition of phosphorylation of the ribosomal protein S6, a downstream target of p70S6K. Remarkably, inhibition occurred at ratios below one apoptotic target per BU.MPT responder cell. Inhibition was evident at a target to responder cell ratio as low as 1:32, particularly in the absence of EGF, and it was virtually complete at ratios of 1:4 or 1:2. These effects were specific to apoptotic targets, because exposure to necrotic targets had a minimal or undetectable effect on the phosphorylation status of p70S6K and S6, whether added alone or in the presence of EGF (Fig. 2B).

Apoptotic Targets Activate AMPK, a Major Upstream Inhibitor of p70S6K—Control of cell growth and p70S6K occurs primarily through mTORC1, a multimolecular serine-threonine kinase complex (26–29). mTORC1 acts as a sensor of the adequacy of factors and conditions necessary for cell growth and proliferation. Information is derived by continuous monitoring within four major areas as follows: 1) the sufficiency of cellular energy stores; 2) the availability of growth factors such as EGF; 3) the adequacy of nutrients such as glucose and amino acids; and 4) the presence and extent of cell damage. Input from these multiple sources can either raise or lower the overall activity of mTORC1, thereby enabling each cell to adjust its metabolic activity to a level appropriate for the prevailing environment (26–29).

The sensing of energy status is dependent primarily on the activity of the intracellular kinase AMPK, a heterotrimeric protein whose activation correlates with phosphorylation of its catalytic α-subunit at Thr172 (23–25). The β- and γ-subunits are
both regulatory. When activated, AMPK inhibits mTORC1 in several ways, both direct and indirect (23–25). As a test of our hypothesis that apoptotic targets act as an extracellular stress, mimicking the effects of intracellular depletion of energy stores, we determined the degree of AMPK phosphorylation following exposure of BU.MPT responders to apoptotic targets (Fig. 3A). Consistent with their unstressed state, quiescent BU.MPT responders displayed a low basal level of AMPK phosphorylation. Exposure to apoptotic targets, either alone or in the presence of EGF, induced a dose-dependent increase in the phosphorylation of AMPK. As with inhibition of phosphorylation of p70S6K, increased phosphorylation of AMPK was observed at a target to responder ratio as low as 1:32. Exposure to necrotic targets produced no detectable change in phosphorylation of AMPK.

A second major input to mTORC1 derives from the survival kinase Akt, which is activated by growth factors like EGF. We have shown previously that apoptotic targets inhibit the phosphorylation of Akt at Ser473 (8, 9). Phosphorylation at this site is performed by mTORC2, a second mTOR-containing multimeric kinase complex (28). Because an uncertain relationship exists between the activities of mTORC1 and mTORC2, we also examined phosphorylation of Akt at Thr308, an event that occurs upstream of mTORC1. Phosphorylation of Akt at Thr308 is mediated by PDK1 and increases the activity of mTORC1 (32, 33).
Exposure to apoptotic targets induced a dose-dependent decrease in phosphorylation of Akt at both Thr\(^{308}\) and Ser\(^{473}\) (Fig. 3B). Basal phosphorylation at these two sites was low, so inhibition was most evident in the presence of EGF. Inhibition of phosphorylation of Akt correlated with inhibition of its activity, as detected by parallel inhibition of phosphorylation of the kinase GSK3\(/\beta\) (downstream of the Thr\(^{308}\) site) and the transcription factors FoxO1 and Fox3a (downstream of the Ser\(^{473}\) site). Necrotic targets had a greatly diminished effect compared with apoptotic targets and, if anything, increased slightly the phosphorylation and activity of Akt (Fig. 3B).

Taken together, these data indicate that exposure to apoptotic targets provokes two signaling events known to lower the activity of mTORC1, namely activation of AMPK and inhibition of Akt. Via subsequent decreased mTORC1-mediated phosphorylation of p70S6K, these two signaling events likely play an important role in the observed inhibition of BU.MPT responder cell growth following exposure to apoptotic targets.

Inhibition of AMPK, Either Pharmacologically or Molecularly, Prevents Apoptotic Target-induced Inhibition of Cell Growth—To establish a mechanistic link between activation of AMPK and inhibition of cell growth following exposure to apoptotic targets, we determined the effect of inhibition of AMPK, either pharmacologically via compound C (CC) (Fig. 4) or molecularly via shRNA (Fig. 5). CC, which inhibits AMPK by reversible competition with AMP for binding to AMPK, has been used to explore the role of AMPK in multiple tissues and cells, including BU.MPT cells (36–38). BU.MPT responders were pretreated with varying concentrations of CC for 2 h prior to exposure to apoptotic targets. As shown in Fig. 4A, pharmacological inhibition of AMPK restored phosphorylation of both p70S6K and S6 to levels observed in cells not exposed to apoptotic targets. Notably, in contrast to its dose-dependent inhibition of the effect of apoptotic targets, CC did not modulate the response to necrotic targets.

Several points merit emphasis. First, in support of the specificity of CC for AMPK, pretreatment with CC had a minimal effect on apoptotic target-induced inhibition of phosphorylation of GSK3\(/\beta\) (downstream of the Thr\(^{308}\) site) and the transcription factors FoxO1 and Fox3a (downstream of the Ser\(^{473}\) site). Necrotic targets had a greatly diminished effect as compared with apoptotic targets and, if anything, increased slightly the phosphorylation and activity of Akt (Fig. 3B).

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Several points merit emphasis. First, in support of the specificity of CC for AMPK, pretreatment with CC had a minimal effect on apoptotic target-induced inhibition of phosphorylation of GSK3\(/\beta\), a downstream target of Akt. Second, despite inhibition of the downstream effects of AMPK, CC had no detectable effect on phosphorylation of AMPK at Thr\(^{172}\). This differs from our previous results, in which CC inhibited phosphorylation of AMPK in response to ATP depletion (38), perhaps indicating a noncanonical activation of AMPK by apoptotic targets (see below). Finally, total levels of AMPK were unaffected by CC and/or apoptotic targets. BU.MPT cells express both \(\beta\)-isoforms of AMPK (\(\beta1\)- and \(\beta2\)-AMPK); of these, \(\beta1\)-AMPK is the more abundant (>80% by densitometry).
Consistent with its restoration of p70S6K phosphorylation and activity, CC (10 μM) also prevented inhibition of cell growth (Fig. 4, B and C). In the absence of CC, apoptotic targets significantly decreased the relative size of BU.MPT responders to 77.2 ± 3.6% that of untreated responders (p < 0.05) (Fig. 4C). In contrast, in the presence of CC, the relative size of BU.MPT responders was 109.9 ± 2.2% that of responders not treated with CC or exposed to apoptotic targets (p < 0.02, CC versus no CC). CC also increased the relative size of responders exposed to necrotic or no targets. This suggests that under the conditions of our studies, AMPK may be exerting a tonic basal inhibition of cell growth.

As the use of pharmacological inhibitors is subject to problems of specificity, we sought a second independent approach to inhibit AMPK. To this end, we used shRNA to knock down expression of the regulatory β-subunit of AMPK (Fig. 5). We targeted the β1 isoform of AMPK, as its expression greatly exceeds that of the β2 isoform in BU.MPT cells (Fig. 4A). The targeted shRNA reduced total AMPK by 75% compared with control shRNA (Fig. 5A).

Overall, molecular inhibition of AMPK replicated the results seen with pharmacological inhibition (Fig. 5). In accord with its knockdown of total AMPK, β1-AMPK shRNA reduced the degree of up-regulation of Thr172 phosphorylation of AMPK in response to apoptotic targets. Knockdown of AMPK restored phosphorylation of p70S6K and S6 in apoptotic target-treated responders to levels equal to or greater than those observed in untreated responder cells. As with CC, inhibition of the phos-
phorylation of GSK3α/β was unaffected by knockdown. Importantly, the responses of BU.MPT cells transduced with control scrambled shRNA were indistinguishable from those of untransduced BU.MPT cells.

Knockdown of AMPK also reduced the ability of apoptotic targets to inhibit cell growth (Fig. 5, B and C). For BU.MPT cells transduced with control shRNA, exposure to apoptotic targets significantly decreased the relative size of responders to 66.1 ± 5.9% that of unexposed responders (p < 0.01) (Fig. 4C). In contrast, for BU.MPT cells transduced with β1-AMPK shRNA, exposure to apoptotic targets reduced the relative size of responders to only 90.0 ± 3.5% that of untreated responders (p < 0.03, presence versus absence of apoptotic targets). The reduction of cell size by responders expressing β1-AMPK shRNA was significantly less than that by responders expressing scrambled shRNA (p < 0.02, β1-AMPK versus scrambled shRNA, presence of apoptotic targets).

Inhibition of Akt Also Contributes to Inhibition of Cell Growth by Apoptotic Targets—Inhibition of the phosphorylation and activity of Akt by apoptotic targets is most evident in the presence of growth factors like EGF (Fig. 3B). Because the availability of growth factors represents a second major input to mTORC1, we next determined the contribution of inhibition of Akt to inhibition of cell growth following exposure to apoptotic targets. To do so, we stably infected BU.MPT cells with a myristoylated Akt (myrAkt)-GFP retroviral construct (32, 33). Myristoylation of Akt leads to its surface membrane localization and constitutive activity (32, 33). BU.MPT cells expressing myrAkt, hereafter denoted as BU.MPT-myrAkt-GFP, were detected by co-expressed GFP. BU.MPT control cells, infected with a retroviral construct containing GFP alone, are denoted as BU.MPT-GFP.

BU.MPT-myrAkt-GFP responders demonstrated increased phosphorylation at both the PDK1 site (Thr308) and the mTORC2 site (Ser473) (Fig. 6A). In accord with the constitutive activation of Akt in BU.MPT-myrAkt-GFP cells, phosphorylation at Thr308 and Ser473 was not affected by exposure to apoptotic targets or pretreatment with CC. Control BU.MPT-GFP cells showed phosphorylation of Akt only at Ser473, which was inhibited by exposure to apoptotic targets. Inhibition was reversed by CC, suggesting that mTORC2 may lie downstream of AMPK. Alternatively, this may represent an off-target effect of CC.

Inhibition of the phosphorylation of p70S6K and S6 did not occur in BU.MPT-myrAkt-GFP responders, despite potent phosphorylation of AMPK following exposure to apoptotic targets (Fig. 6B). This lack of inhibition resembles that observed with inhibition of AMPK (Figs. 4 and 5). As expected with constitutive activation of Akt, apoptotic targets did not inhibit Akt-mediated phosphorylation of GSK3α/β in BU.MPT-myrAkt-GFP responders.
Notably, despite restoration of the phosphorylation of p70S6K and S6 in BU.MPT-myrAkt-GFP responders, inhibition of cell growth was still observed (Fig. 6, C and D). Relative size following exposure to apoptotic targets was reduced to 76.0 ± 3.1% in BU.MPT responder cells pretreated with vehicle or compound C at a target/responder cell ratio of 1:1 for 30 min, washed, incubated for a further 15 min, and then harvested. The source of apoptotic targets was staurosporine-treated BU.MPT cells. In all cases, targets and nonadherent responder cells were removed by washing, and BU.MPT responder cell lysates were probed with anti-phosphorylated AMPK-α chain, p70S6K, and GSK3α/β antibodies as shown. Equal loading was confirmed by probing for total AMPKα1/β1. Shown is a representative set of blots from three separate experiments. Gaps between lanes exist for clarity of presentation or because lanes were noncontiguous on the original immunoblot.

$\text{FIGURE 4. Pharmaco logical inhibition of AMPK prevents inhibition of p70S6K and cell growth by apoptotic targets.}$ A, serum-starved BU.MPT responder cells were pretreated for 2 h with vehicle or compound C at the indicated concentrations, following which they were stimulated with either apoptotic (Apo) targets or necrotic (Necro) targets at a target/responder cell ratio of 1:1 for 30 min, washed, incubated for a further 15 min, and then harvested. The source of apoptotic targets was staurosporine-treated BU.MPT cells. In all cases, targets and nonadherent responder cells were removed by washing, and BU.MPT responder cell lysates were probed with anti-phosphorylated AMPK-α chain, p70S6K, S6, and GSK3α/β antibodies as shown. Equal loading was confirmed by probing for total AMPKα1/β1. Shown is a representative set of blots from three separate experiments. Gaps between lanes exist for clarity of presentation or because lanes were noncontiguous on the original immunoblot. B, serum-starved BU.MPT responder cells were pretreated with vehicle or compound C (10 μM) for 2 h and then exposed to no targets (Untreated), apoptotic (Apo) targets, or necrotic (Necro) targets at a target/responder cell ratio of 10:1 for 2 h. The source of apoptotic targets was actinomycin D-treated DO11.10 cells. At 24 h following exposure, relative cell size was determined by flow cytometric analysis of the forward scatter of responder cells in the G1 phase of the cell cycle. C, graph depicts the mean and S.E. from three separate flow cytometric analyses of the relative cell size of BU.MPT responder cells in the G1 phase of the cell cycle. Within each analysis, experimental values were normalized to the mean forward scatter of BU.MPT responders exposed to no targets (Control) in the absence of compound C. $p < 0.05$, apoptotic targets versus no targets, in the absence of compound C; $p < 0.02$, apoptotic targets in presence versus absence of compound C. Tar, target(s). Error bars (C) denote S.E. D, densitometric quantitation, using total AMPKα1/β1 as a loading control, is provided for the immunoblots shown in A. $p < 0.05$ for the following conditions: apoptotic targets versus no targets for AMPKα-T172 and GSK3α/β-S21/9 (in either the absence or the presence of all concentrations of compound C), and for p70S6K-T389 and S6-S240/244 (only in the absence of compound C). $p = n$ not significant for the following conditions: apoptotic targets versus no targets for p70S6K-T389 and S6-S240/244, in the presence of all concentrations of compound C.

Notably, despite restoration of the phosphorylation of p70S6K and S6 in BU.MPT-myrAkt-GFP responders, inhibition of cell growth was still observed (Fig. 6, C and D). Relative size following exposure to apoptotic targets was reduced to 76.0 ± 3.1% in BU.MPT-GFP responders ($p < 0.001$) as opposed to 82.8 ± 1.7% in BU.MPT-myrAkt-GFP responders ($p < 0.05$). Although the reduction of cell size occurring in BU.MPT-myrAkt-GFP responders was significantly less than that in BU.MPT-myrAkt-GFP responders ($p < 0.05$), the degree of improvement was still less than that observed with inhibition of AMPK, either pharmacologically by CC ($p < 0.001$) or molecularly by shRNA ($p < 0.05$). Thus, of the two signaling events contributing to growth inhibition, namely activation of AMPK and inhibition of Akt, prevention of AMPK activation largely abrogated cell growth inhibition (Figs. 4C and 5C), whereas prevention of Akt inhibition only partially abrogated cell growth inhibition (Fig. 6C). Together, these data suggest that AMPK activation plays the greater role in apoptotic target-mediated inhibition of cell growth.
Apoptotic Targets Activate AMPK Independently of Changes in Intracellular Energy Stores—AMPK is a heterotrimeric kinase activated by decreasing concentrations of ATP and increasing concentrations of AMP and ADP (23–25). Activation of AMPK depends on phosphorylation of its catalytic subunit at Thr172. Binding of either AMP or ADP promotes phosphorylation of AMPK at this site. In addition, binding of AMP, but not ADP, can directly activate AMPK. AMP sustains AMPK activity by inhibiting its dephosphorylation, whereas ATP promotes dephosphorylation. Thus, both ADP to ATP and AMP to ATP ratios contribute to AMPK regulation.

To determine whether apoptotic targets activate AMPK by inducing changes in intracellular energy stores, we measured intracellular concentrations of AMP, ADP, and ATP by LC/MS/MS in BU.MPT responders following exposure to apoptotic targets (Fig. 7). The calculated quantity of energy charge represents the fraction of adenylate-bound high energy phosphate bonds present, as compared with the maximum possible, and is defined by Equation 1.

\[
\text{Energy Charge} = \frac{[\text{ATP}]}{[\text{ATP}} + \frac{1}{2}[\text{ADP}]] + [\text{AMP}]
\]

This formula yields a value between 0 (100% AMP) and 1 (100% ATP).

Exposure to apoptotic targets for 2 h produced no detectable change in either the ADP to ATP ratio, the AMP to ATP ratio, or the energy charge at either 0.5 or 18 h following exposure \((p > 0.8)\). The first time corresponds to that at which we performed all immunoblots. The absence of a change in intracellular energy stores at a time when AMPK is strongly phosphorylated (see Figs. 3, 4A, 5A, and 6B) provides powerful evidence that exposure to apoptotic targets induces activation of AMPK via an energy-independent pathway (23–25). As further support for this idea, we also assessed intracellular energy stores at 18 h, because this later time corresponds to when we performed all flow cytometric analyses of growth inhibition. Again, there was no detectable change in intracellular levels of AMP, ADP, or ATP.

Inhibition of Cell Growth by Apoptotic Targets Depends on Cell-Cell Interaction but Is Independent of Phagocytosis—BU.MPT responders were grown in a Transwell®-permeable support system and separated from apoptotic targets by a 0.4 \(\mu\)M polycarbonate membrane. Prevention of physical interaction between BU.MPT responders and apoptotic targets abolished the ability of apoptotic targets to inhibit cell growth (Fig. 8A). Necrotic targets had no effect on relative cell volume, irrespective of the presence or absence of the Transwell®-permeable support system (data not shown).

To test the role of phagocytosis, we used the cytoskeletal inhibitor cytochalasin D to prevent phagocytosis. Inhibition of cell growth in response to apoptotic targets occurred in the absence of phagocytosis (Fig. 8B). We obtained comparable...
FIGURE 5. Molecular inhibition of AMPK prevents inhibition of p70S6K and cell growth by apoptotic targets. A, serum-starved BU.MPT responder cells, untransduced (—) or stably transduced with either control scrambled (Scrambled) or β1-AMPK (β1-AMPK) shRNA, were stimulated with apoptotic targets (Apo) or necrotic targets (Necro) at a target/responder cell ratio of 1:1 for 30 min, washed, incubated for a further 15 min, and then harvested. The source of apoptotic targets was staurosporine-treated BU.MPT cells. In all cases, targets and nonadherent responder cells were removed by washing, and BU.MPT responder cell lysates were probed with anti-phosphorylated AMPKα chain, p70S6K, S6, and GSK3α/β and anti-total AMPK and β1-AMPK antibodies as shown. Equal loading was confirmed by probing for total GAPDH. Shown is a representative set of blots from three separate experiments.

B, serum-starved BU.MPT responder cells, stably transduced with either control scrambled (Scrambled) or β1-AMPK (β1-AMPK) shRNA, were exposed to no targets (Untreated) or apoptotic targets (Apo) at a target/responder cell ratio of 10:1 for 6 h. The source of apoptotic targets was actinomycin D-treated DO11.10 cells. At 24 h following exposure, relative cell size was determined by flow cytometric analysis of the forward scatter of responder cells in the G1 phase of the cell cycle. C, graph depicts the mean and S.E. from four separate flow cytometric analyses of the relative cell size of BU.MPT responder cells in the G1 phase of the cell cycle. Within each analysis, experimental values were normalized to the mean forward scatter of BU.MPT responders, expressing either scrambled or β1-AMPK shRNA, not exposed to targets (No targets). p < 0.01, apoptotic targets versus no targets, scrambled shRNA; p < 0.03, apoptotic targets versus no targets, β1-AMPK shRNA; p < 0.02, scrambled versus β1-AMPK shRNA, apoptotic targets. Error bars (C) denote S.E. D, densitometric quantitation, using total GAPDH as a loading control, is provided for the immunoblots shown in A. p < 0.05 for the following conditions: apoptotic targets versus no targets for AMPKα-T172, p70S6K-T389, and S6-S240/244, in untransduced and scrambled shRNA-transduced responders; and for GSK3α/β-S21/9 in untransduced, scrambled shRNA-transduced, and β1-AMPK-transduced responders. p = not significant for the following conditions: apoptotic targets versus no targets for AMPKα-T172, p70S6K-T389, and S6-S240/244 in β1-AMPK-transduced responders.
data with two concentrations of cytochalasin D, both of which inhibit m\(\phi\) and BU.MPT phagocytosis by \(\geq 90\%\) (8, 9). Necrotic targets had no effect on relative cell volume, irrespective of the presence or absence of cytochalasin D (data not shown). Taken together, our results show that apoptotic target-mediated inhibition of BU.MPT growth requires direct physical interaction between targets and responder cells but is independent of phagocytosis.

**Discussion**

We show here that exposure of live kidney PTECs to apoptotic target cells inhibits PTEC cell growth, as manifested by a decreased size of cells in G\(1\) phase of the cell cycle. Apoptotic targets inhibit cell growth by modulating the activity of at least two intracellular signaling pathways, both of which converge on mTORC1. The first pathway results from inhibition of Akt, a survival kinase whose activation raises the activity of mTORC1 (28, 39–43). By inhibiting Akt, apoptotic targets lower mTORC1 activity and therefore the activity of the ribosomal kinase p70S6K, a key regulator of cell growth and an immediate downstream target of mTORC1 (26–29). The second pathway influenced by apoptotic targets results from activation of AMPK, a critical sensor of intracellular energy stores (23–25, 28). Activation of AMPK by apoptotic targets provides a second inhibitory input to mTORC1. Remarkably, activation of AMPK by apoptotic targets occurs in an energy-independent manner, without detectable changes in intracellular levels of ATP, ADP, or AMP. These events are all specific to apoptotic targets. Exposure to necrotic targets produced no detectable effect on cell growth or AMPK activity, and, if anything, weakly stimulated Akt. Consistent with our previous observations that kidney PTECs recognize apoptotic versus necrotic targets via distinct receptors (8, 9), inhibition of PTEC cell growth by apoptotic targets requires direct physical interaction between targets and PTEC responders. As in our previous studies (8, 9), inhibition of cell growth was independent of phagocytosis.

Of the two signaling events leading to apoptotic target-mediated inhibition of cell growth, namely activation of AMPK and inhibition of Akt, AMPK activation appears to be the more potent. The dominance of AMPK activation as a mediator of growth inhibition by apoptotic targets is seen by comparison of Figs. 4C and 5C with Fig. 6D. Reversal of AMPK activation almost completely abrogated cell growth inhibition by apoptotic cells, whereas reversal of Akt inhibition only partially abrogated cell growth inhibition. Thus, pharmacological inhibition of AMPK with CC increased relative cell size from 77.2 \(\pm\) 3.6 to 109.9 \(\pm\) 2.2\% \((p < 0.02)\), and similarly, molecular inhibition of AMPK increased relative cell size from 66.1 \(\pm\) 5.9 to 90.0 \(\pm\) 3.5\% \((p < 0.02)\). In marked contrast, constitutive activation of Akt only increased relative size from 76.0 \(\pm\) 3.1 to 82.8 \(\pm\) 1.7\%. The degree of improvement with activation of Akt was significantly less than that observed with inhibition of AMPK, either pharmacologically by CC \((p < 0.001)\) or molecularly by shRNA
Apoptotic Cells Activate AMPK and Inhibit Cell Growth

FIGURE 6. Constitutive activation of Akt prevents inhibition of p70S6K and cell growth by apoptotic targets. A and B, serum-starved BU.MPT responder cells, infected with retroviral constructs containing either GFP alone (GFP) or GFP plus constitutively active Akt (myrAkt-GFP), were stimulated with no targets (−) or apoptotic (Apo) targets at a target/responder cell ratio of 1:1 or 1:4 for 30 min in the absence or presence of compound C (20 μM), washed, incubated for a further 15 min, and then harvested. The source of apoptotic targets was staurosporine-treated BU.MPT cells. In all cases, targets and nonadherent responder cells were removed by washing, and BU.MPT responder cell lysates were probed with anti-phosphorylated Akt antibodies (A) or anti-phosphorylated AMPKα chain, p70S6K, S6, and GSK3α/β antibodies (B) as shown. Equal loading was confirmed by probing for either total GAPDH (A) or total AMPKβ1/β1 (B). Shown is a representative set of blots from three separate experiments. Gaps between lanes exist for clarity of presentation or because lanes were noncontiguous on the original immunoblot. C, serum-starved BU.MPT responder cells, infected with retroviral constructs containing either GFP alone (GFP) or GFP plus constitutively active Akt (myrAkt-GFP), were exposed to no targets, apoptotic (Apo) targets, or necrotic (Necro) targets at a target/responder cell ratio of 10:1 for 24 h in the absence (Control) or presence of compound C (10 μM). The source of apoptotic targets was actinomycin D-treated DO11.10 cells. 24 h following exposure, relative cell size was determined by flow cytometric analysis of the forward scatter of responder cells in the G1 phase of the cell cycle. D, graph depicts the mean and S.E. from six separate flow cytometric analyses of the relative cell size of BU.MPT responder cells in G1 phase of the cell cycle. Within each analysis, experimental values were normalized to the mean forward scatter of BU.MPT responders expressing GFP alone (GFP) immediately prior to target exposure at 0 h. p < 0.001, apoptotic targets versus no targets, GFP; p < 0.05, apoptotic targets versus no targets, myrAkt-GFP; p < 0.05, GFP versus myrAkt-GFP, apoptotic targets. Tar, target(s). Error bars (D) denote S.E. Densiometric quantitation, using total AMPKβ1/β1 as a loading control, is provided for the immunoblots shown in B. p < 0.05 for the following conditions: apoptotic targets versus no targets in control GFP-transfected responders for S6-S240/244 and GSK3α/β-21 at apoptotic target/responder cell ratios of 1:1 and 1:4; and in control GFP-transfected responders for AMPKα-T172 and p70S6K-T389 at an apoptotic target/responder cell ratio of 1:1. p = not significant for the following conditions: apoptotic targets versus no targets in myrAkt-GFP-transfected responders for AMPKα-T172, p70S6K-T389, S6-S240/244, and GSK3α/β-219 at all apoptotic target/responder cell ratios, regardless of the presence or the absence of compound C.
These data indicate that activation of AMPK in response to apoptotic targets exerts a greater growth inhibitory effect than does inhibition of Akt. A potential explanation may lie in the mechanisms by which Akt and AMPK modulate mTORC1 activity. Whereas Akt raises the activity of mTORC1, AMPK lowers its activity (28). In the case of Akt, mTORC1 activation involves a release of inhibition through phosphorylation of two negative regulators of mTORC1. The first target of Akt is the tuberous sclerosis complex (TSC) component tuberin (TSC2), an upstream inhibitor of mTORC1 (39–41). The second target of Akt is PRAS40, a negative regulator within the mTORC1 complex. Phosphorylation of PRAS40 by Akt causes its dissociation from mTORC1 (42, 43). Relief of inhibition from these two sources, the first upstream of mTORC1 and the second within the mTORC1 complex, leads to activation of mTORC1. Like Akt, AMPK targets both TSC2 and a protein within the mTORC1 complex, but phosphorylation by AMPK has opposite effects to those of Akt. Thus, phosphorylation of TSC2 by AMPK leads to the activation of this upstream inhibitor of mTORC1 (41, 45). The second AMPK target is Raptor, a scaffold protein necessary for the assembly of mTORC1 as well as for interaction of mTORC1 with its various substrates and regulators. Phosphorylation of Raptor by AMPK leads to Raptor’s sequestration by 14-3-3 proteins, rendering Raptor incapable of fulfilling its role as a scaffolding molecule (46). Because Akt and AMPK have overlapping targets, the greater potency of AMPK in inhibiting cell growth may reflect the dominance of inhibitory over stimulatory signals. For example, dissociation of the negative regulator PRAS40 from mTORC1 may be of lesser relevance if Raptor, the scaffold upon which mTORC1 assembles, is sequestered and unavailable. Alternatively, the seemingly greater potency of AMPK activation over Akt inhibition may reflect off-target effects of CC and/or differences in the penetrance and effectiveness between the myrAkt and β1-AMPK shRNA constructs.

The lack of a detectable change in the ADP to ATP ratio, AMP to ATP ratio, or energy charge following exposure to apoptotic targets implies that activation of AMPK occurs in an energy-independent manner. Thus, despite strong phosphorylation and activation of AMPK at Thr172 increases AMPK activity by more than 100-fold (23–25). The major upstream kinase responsible for phosphorylating AMPK in response to energy stress is liver kinase B1 (LKB1) (23–25, 47). Phosphorylation of the catalytic α-subunit of AMPK at Thr172 increases AMPK activity by more than 100-fold (23–25). The major upstream kinase responsible for phosphorylating AMPK in response to energy stress is liver kinase B1 (LKB1) (23–25, 47). Phosphorylation of AMPK by LKB1 is enhanced by binding of ADP and AMP to the γ-subunit of AMPK. Binding of ADP and AMP also inhibits dephosphorylation of AMPK (23–25). AMP has an additional effect not shared by ADP. Allosteric changes induced by binding of AMP, but not ADP, further increase AMPK activity by as much as 10-fold (23). However,
activation of AMPK by LKB1 has been described only under circumstances of energy stress. Therefore, unless exposure to apoptotic targets induces signaling events that permit LKB1 to phosphorylate AMPK in the absence of a change in intracellular energy stores, LKB1 is unlikely to mediate AMPK activation in our system.

A second kinase capable of phosphorylating AMPK at Thr\(^{172}\) of its \(\alpha\)-subunit is Ca\(^{2+}\)/calmodulin-activated protein kinase \(\beta\) (CaMKK\(\beta\), also known as CAMKK2) (23, 48). In LKB1-deficient cells, CaMKK\(\beta\) phosphorylates and activates the \(\alpha\)-subunit of AMPK at Thr\(^{172}\) in response to an increase of intracellular Ca\(^{2+}\) (49). Interestingly, purified CaMKK\(\beta\) forms a stable multiprotein complex \textit{in vitro} with the \(\alpha\)- and \(\beta\)-subunits of AMPK. Because the AMP- and ADP-binding \(\gamma\)-subunit of AMPK is absent, phosphorylation and activation of AMPK within this complex are regulated by Ca\(^{2+}\) but not AMP or ADP (50, 51). The \textit{in vivo} significance of this observation remains uncertain (52). Besides increased intracellular Ca\(^{2+}\), several other signaling molecules, such as protein kinase A (PKA), are known to modulate the activity of CaMKK\(\beta\) (53).

In addition to LKB1 and CaMKK\(\beta\), a number of other proteins, second messengers, and intracellular stresses have been reported to modulate the activity of AMPK. Although their mechanisms of activation remain unclear, these include second messengers like cAMP (53, 54), proteins and kinases such as p53 (55), PKA (54), and ataxia telangiectasia mutated (56), and a variety of intracellular stresses such as oxidative (23, 28, 44, 56) and genotoxic (23, 28, 55). Future studies will address whether any of these molecules participate in the activation of AMPK by LKB1.
AMPK by apoptotic targets or whether apoptotic targets activate AMPK via an as yet unrecognized and novel pathway.

Regardless of the pathway by which apoptotic targets activate AMPK, it appears that PTECs respond to the extracellular presence of apoptotic targets as if subjected to a metabolic or other stress. Together with previous work from our and others’ laboratories (3–9, 16–18), our results suggest that the network of signaling responses to dead targets, as observed in viable responder cells of multiple lineages, represents a critical and ubiquitous dimension of tissue homeostasis. We hypothesize that, throughout the local, alterations in the mode and extent of cell death alert viable resident cells in the vicinity to environmental change or stress (9, 22). A local increase in the number of dead or dying cells may be triggered by diverse stresses such as ischemia, infection, aging, or acute injury. Viable resident cells can then extract information about the nature of the stress and the possibility for adaptation, via their interaction with adjacent dead or dying cells. Responding cells gauge the severity of the environmental disturbance not simply from the number of dead cells but also from the mode of death (apoptotic versus necrotic), pattern of distribution of dead cells, and timing of their appearance. For example, in the case of necrotic targets, the disturbance is likely to be sudden and catastrophic, whereas in the case of apoptotic targets, it may be more gradual and potentially adaptable. Such considerations may underlie the observation that, in general, the responses elicited by apoptotic versus necrotic targets are oppositely directed (3–6, 8, 9, 22). For example, we have previously shown that apoptotic targets inhibit the proliferation and survival of PTECs, although necrotic targets promote these activities (8, 9). In this study, we show that apoptotic targets activate AMPK and inhibit cell growth, although the effect of necrotic targets is minimal or neutral.

It is important to emphasize that the cellular response to dead targets is varied, robust, and complex. This breadth of response permits cells and organs to tailor their behavior to function and need. Thus, within an organ the responses of nearby resident cells may vary considerably, depending on their lineage or differentiative stage (6–9, 17–19, 22). Even for responder cells of similar lineage, responses can differ depending on their origin or state of activation (9, 20).

Seen in light of this hypothesis, inhibition of PTEC cell growth following exposure to apoptotic targets may represent one aspect of an overall adaptive strategy, in response to a threatened reduction in metabolic supply. Activation of AMPK initiates a wide range of intracellular events that affect intermediary cell metabolism. The net result is the conservation or augmentation of intracellular energy stores, through increased energy production, decreased energy consumption, and facilitated cellular uptake of nutrients (23, 24). Metabolic demand by PTECs is further decreased following exposure to apoptotic targets by a reduction in population size, brought about through the combined effects of decreased proliferation and increased cell death (8, 9). Thus, in the case of PTECs, exposure to apoptotic targets produces a stress response, whose ultimate effect is a drastic reduction in metabolic demand, achieved via a decrease not only in the number of live cells but also in the energy expenditure per cell.

In summary, we have shown that exposure to apoptotic targets inhibits the growth of live PTEC responders. At least two signaling events cooperate to produce this inhibition, namely inhibition of Akt and activation of AMPK. Activation of AMPK occurs without detectable change in intracellular energy stores. These data offer further support to the hypothesis that the recognition of apoptotic targets by live responder cells represents an important source of information for responder cells about their environment. In the case of PTECs, a highly metabolic lineage, apoptotic cells serve as an extracellular stress, mimicking the effects of depletion of intracellular energy stores. By acting as sentinels of environmental change, apoptotic targets permit nearby cells to monitor their environment and adapt to local changes. Adaptation involves the modulation of multiple critical cell activities. These include the proliferation, survival, and, as shown, here the growth of responder cells.

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