Metabolic Regulation of Protein N-Alpha-Acetylation by Bcl-xL Promotes Cell Survival

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SUMMARY

Previous experiments suggest a connection between the N-alpha-acetylation of proteins and sensitivity of cells to apoptotic signals. Here, we describe a biochemical assay to detect the acetylation status of proteins and demonstrate that protein N-alphaacetylation is regulated by the availability of acetyl-CoA. Because the antiapoptotic protein Bcl-xL is known to influence mitochondrial metabolism, we reasoned that Bcl-xL may provide a link between protein N-alpha-acetylation and apoptosis. Indeed, Bcl-xL overexpression leads to a reduction in levels of acetyl-CoA and N-alpha-acetylated proteins in the cell. This effect is independent of Bax and Bak, the known binding partners of Bcl-xL. Increasing cellular levels of acetyl-CoA by addition of acetate or citrate restores protein N-alpha-acetylation in Bcl-xL-expressing cells and confers sensitivity to apoptotic stimuli. We propose that acetyl-CoA serves as a signaling molecule that couples apoptotic sensitivity to metabolism by regulating protein N-alphaacetylation.

INTRODUCTION

Increasing evidence suggests that specific metabolic alterations associated with cancer cells may not be ancillary to their trans-

formation but are instrumental to their tumorigenic potential by mediating cell proliferation, growth, and survival (Vander Heiden et al., 2009). Many oncogenes and tumor suppressor genes known to promote excess cell proliferation also alter biosynthetic (or anabolic) processes. For example, Akt expression stimulates glucose uptake and glycolysis, the pentose phosphate pathway, and fatty acid synthesis. c-Myc expression promotes glutamine metabolism as well as purine and pyrimidine biosynthesis. Furthermore, mutations in genes encoding metabolic enzymes have been identified by cancer genetic association studies (Vander Heiden et al., 2009). How specific metabolites contribute to increased proliferation and apoptotic resistance in tumor cells remains a central unanswered question.

The proto-oncogene Bcl-xL has a prominent role in promoting cell survival and cancer development (Boise et al., 1993). It is well established that Bcl-xL protects against apoptosis by directly binding and inhibiting Bax/Bak oligomerization-mediated mitochondrial permeabilization. However, certain Bcl-xL mutants, such as F131V/D133A and G148E, that are unable to bind to Bax or Bak, nevertheless retain 70%–80% antiapoptotic activity of WT Bcl-xL (Cheng et al., 1996). Curiously, Bcl-xL has also been shown to regulate mitochondrial respiration and metabolism (Gottlieb et al., 2000; Vander Heiden et al., 1999). Whether the metabolic function of Bcl-xL contributes to its role in mediating apoptotic resistance is unclear.

Our unexpected identification of an N-terminal acetyltransferase, Arrest Defective 1 (dARD1), in a genome-wide RNA interference (RNAi) screen in *Drosophila* cells for apoptotic regulators (Yi et al., 2007) prompted us to posit that protein N-alpha-acetylation, a major N-terminal modification, links cell metabolism

to apoptotic induction in cancer cells. Since *dARD1* is epistatic to *diap1*, which encodes for a direct inhibitor of caspases in *Drosophila*, and ARD1 is required for caspase activation in mammalian cells (Yi et al., 2007), the role for ARD1 in mediating caspase activation is evolutionarily conserved. How ARD1 regulates caspase activation has not yet been illustrated.

In mammalian cells, protein N-alpha-acetylation is mediated by the highly conserved N-acetyltransferase protein complexes (NatA, NatB, NatC, NatD, and NatE). The NatA complex consists of the catalytic subunit, Arrest Defective 1 (hNaa10p/ ARD1), and the auxiliary subunit, N-acetyltransferase 1 (NAT1/ hNaa15p/NATH), whereas NatB consists of N-terminal acetyltransferase 3 (hNaa20p/NAT3) and mitochondrial distribution and morphology 20 (hNaa25p/Mdm20). Although the Nat complexes are implicated in regulating cell-cycle progression, cell proliferation, and tumorigenesis, the mechanisms that connect N-alpha-acetylation to the cellular protein apparatus are unknown (Ametzazurra et al., 2008; Polevoda and Sherman, 2003; Starheim et al., 2008, 2009). Recent N-acetylome studies reveal incomplete acetylation status of proteins (Arnesen et al., 2008; Goetze et al., 2009). Although a commonly accepted view is that partial acetylation results from the degenerate nature of protein N-terminal sequences, we considered the possibility that protein N-alpha-acetylation might be regulated, an alternative hypothesis that had not been tested as a result of technical limitations.

Here, we developed a biochemical approach to assess the status of endogenous levels of protein N-alpha-acetylation. Using this assay, we show that protein N-alpha-acetylation levels are sensitive to alterations in metabolism and Bcl-xL expression. Bcl-xL overexpression leads to reduced levels of acetyl-CoA and hypoacetylation of protein N termini through a Bax/Bak-independent mechanism. Conversely, *bcl-x*^{-/-} mouse embryonic fibroblasts show increased levels of acetyl-CoA as well as protein N-alpha-acetylation levels. Protein N-alphaacetylation deficiency in Bcl-xL-overexpressing cells contributes to apoptotic resistance since increasing acetyl-CoA production can rescue this deficiency in protein N-alpha-acetylation and sensitize Bcl-xL cells to cell death. Our study suggests that regulation of acetyl-CoA availability and protein N-alphaacetylation may provide a Bax/Bak-independent mechanism for Bcl-xL to regulate apoptotic sensitivity.

RESULTS

We confirmed that ARD1 is necessary for cell death induced by the DNA-damaging agent doxorubicin in multiple cell lines of different origins, including *Drosophila* Kc (Yi et al., 2007), HeLa, HT1080, and U2OS cells (Figures 1A–1D). In addition, HeLa and U2OS cells deficient for NATH were also resistant to doxorubicin treatment, recapitulating the apoptotic resistant phenotype of ARD1 knockdown cells (Figures 1A–1D). Thus, the acetylation activity of the NatA complex serves to influence the sensitivity of these cells to apoptosis. Next, we tested whether NatA influences apoptotic sensitivity to other DNA damaging agents. We found that ARD1 knockdown cells are also resistant to cisplatin and ultraviolet (UV) treatment (Figure 1E). However, these cells remained sensitive to tumor

necrosis factor (TNFalpha) and cyclohexamide treatment, which specifically activates apoptosis through the death receptor pathway (Figure 1F). We conclude that protein N-alphaacetylation regulates apoptotic sensitivity downstream of DNA damage.

Since N-alpha-acetylation has been suggested to affect protein stability (Polevoda and Sherman, 2003), we examined whether protein synthesis and/or protein turnover might be affected by acetylation status. We tested whether ARD1 substrates such as caspase-2 and Chk1 (see results below) are destabilized in ARD1 knockdown cells using cyclohexamide, an inhibitor of protein synthesis. Deficiency in ARD1 did not lead to decreases in the cellular levels of these proteins compared to that of control (Figure S1A available online). The steady-state levels of total cellular proteins in ARD1 knockdown cells were similar to the levels in control cells (Figure S1B). We also tested whether general protein stability is altered in ARD1 or NATH knockdown cells (Figure S1C). By pulse-chase ³⁵S-Met labeling experiments, we observed that neither general protein synthesis nor turnover was affected in ARD1 or NATH knockdown cells. Thus, protein N-alpha-acetylation mediated by NatA complex is not required to maintain protein stability globally. In addition, we verified that cell-cycle progression is unaffected in cells deficient for ARD1/NATH (Figure S1D). Taken together, these data suggest that the NatA complex may influence apoptotic sensitivity by mediating protein N-alpha-acetylation of key apoptotic components.

In Vitro Detection of Unmodified Protein N Termini

The lack of an immunological method to detect the acetylation status of protein N termini has limited our understanding of the mechanisms that regulate protein N-alpha-acetylation. To this end, we developed a selective biotin labeling method using an engineered protein ligase, termed subtiligase (Abrahmsén et al., 1991; Tan et al., 2007) that detects nonacetylated N termini of endogenous proteins. This approach was used to capture unmodified protein N termini resulting from caspase-mediated cleavage during apoptotic cell death (Mahrus et al., 2008). Unblocked N termini can be labeled using subtiligase, which preferentially biotinylates N-terminal amine groups consistent with the specificity of NatA or NatB (Abrahmsén et al., 1991; Mahrus et al., 2008). As the N termini of up to 80%-90% of cellular proteins may be blocked by a number of different modifications (Martinez et al., 2008), very few proteins will be biotin labeled by subtiligase as previously demonstrated (Mahrus et al., 2008). Thus, any protein that is biotin labeled by subtiligase in our assays most likely results from a specific loss in N-alphaacetylation.

We utilized subtiligase to biotinylate free N termini of proteins in whole-cell lysates followed by avidin affinity purification and western blot analysis. Decreased levels of protein N-alpha-acetylation are expected to increase subtiligase-mediated protein biotinylation and conversely, increased levels of protein N-alpha-acetylation are expected to decrease subtiligase-mediated protein biotinylation (Figure 2A). First, we asked whether the subtiligase assay could be used to distinguish the N-alpha-acetylation status of protein N termini when the expression of the NatA complex is diminished by RNAi-mediated knockdown.

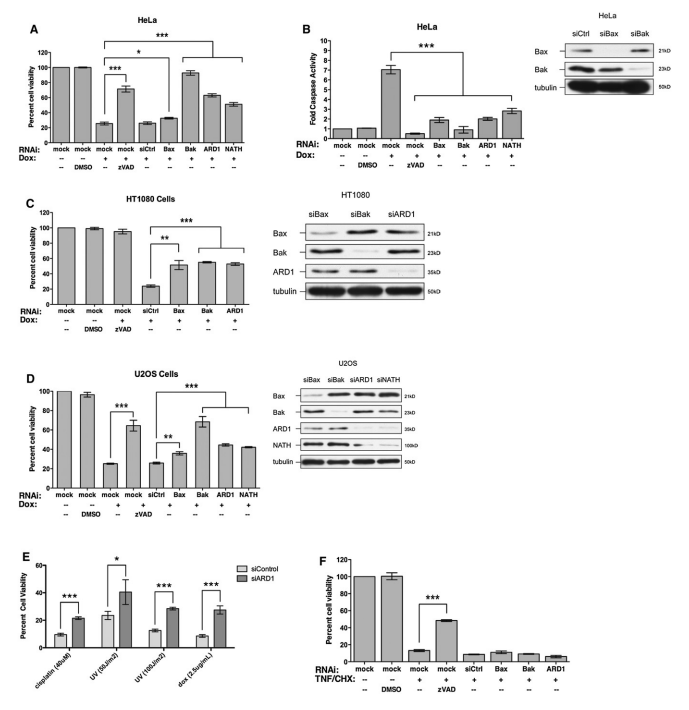


Figure 1. NatA Knockdown Suppresses Cell Death Induced by DNA Damage in HeLa, HT1080, and U2OS Cells

(A and B) HeLa cells were treated with doxorubicin (1.25 µg/ml, 20 hr for cell viability; 5 µg/ml, 8 hr for caspase activity).

- (C) HT1080 cells were treated with doxorubicin (1.25 μg/ml, 20 hr).
- (D) U2OS cells were treated with doxorubicin (1.25 μ g/ml, 20 hr).
- (E) HeLa cells were treated with cisplatin (40 μ M) or UV (50 J/m² or 100 J/m²) for 24 hr.
- (F) HeLa cells were treated with TNFalpha (10 ng/ml, 24 hr) and cyclohexamide (1 µg/ml, 24 hr) to induce death receptor mediated cell death. Immunoblots were conducted in parallel to show extent of target knockdown. Data are represented as mean ± standard deviation (SD; n = 3). (Student's t test: p < 0.05; **p < 0.01; ***p < 0.001.) See also Figure S1.

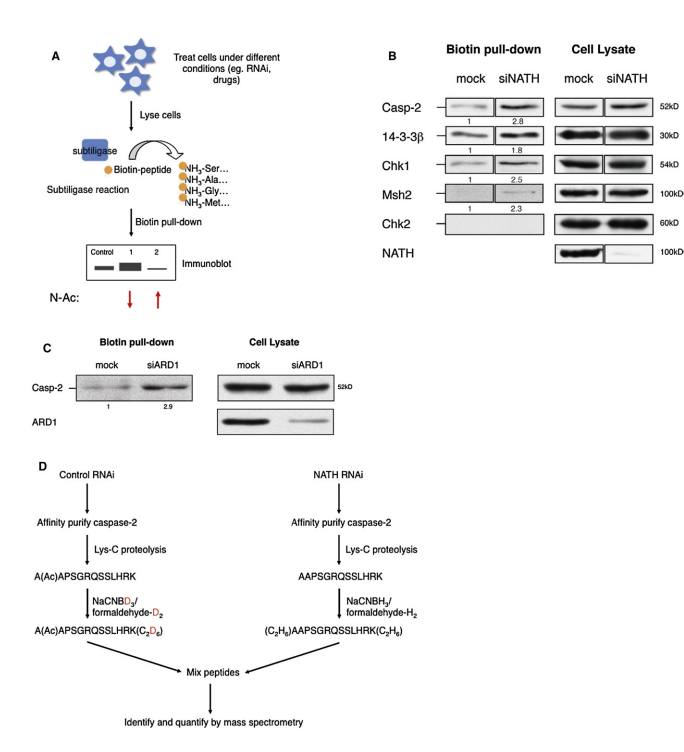


Figure 2. A Subtiligase-Based Assay to Label Unacetylated Protein N Termini

(A) A schematic of the subtiligase assay. In brief, cells were gently lysed in NP-40 lysis buffer on ice. Cell debris was pelleted, and the resulting supernatant was subjected to a subtiligase reaction (1 μ M subtiligase, 1 mM biotin-peptide, 2 mM DTT; 1 hr at room temperature). Biotinylated proteins were affinity purified with Neutravidin beads (Thermo Scientific) and analyzed by SDS-PAGE. Protein N-alpha-acetylation levels were determined by immunoblot against proteins in which the N-terminal residue corresponds to specificity of a Nat complex. Chk2 was included as a negative control, which should not be recognized by subtiligase based on sequence.

(B and C) Subtiligase reaction was conducted on HeLa cells transfected with siRNA against NATH (B) or ARD1 (C) as described in (A). Enrichment of biotin-labeled NatA substrates was observed in ARD1 or NATH-deficient cells compared to that of control. Enrichment in the amount of pulldown suggests a decrease in protein N-alpha-acetylation levels of NatA substrates in NatA-deficient cells. Blot quantification for all subtiligase assays was calculated relative to control and normalized to the corresponding lysate sample with ImageJ software.

ARD1 acetylates a subclass of proteins with Ser, Ala, or Thr as the newly exposed N-terminal residue after initiator Met (iMet) cleavage (Polevoda and Sherman, 2003). We tested 14-3-3\beta, which is known to be N-alpha-acetylated (Arnesen et al., 2009; Martin et al., 1993), and proteins that we predict to be N-alpha-acetylated based on their sequences, Chk1 and Msh2. Caspase-2, which is responsive to both DNA damage (Tinel and Tschopp, 2004) and metabolic stress (Nutt et al., 2005, 2009), is also a good candidate for acetylation by ARD1 as the second amino acid in the caspase-2 polypeptide is Ala. We observed that these proteins as well as caspase-2 were biotinylated to a higher extent by subtiligase in NATH or ARD1 knockdown cells than in control cells (Figures 2B and 2C). These data suggest that subtiligase can distinguish N-alpha-acetylation of multiple proteins that is dependent on NatA expression.

To determine the validity of subtiligase assay, we measured the extent of protein N-alpha-acetylation by quantitative mass spectrometry using differential isotope labeling (Figure 2D). First, we tested whether we could detect the basal levels of N-alphaacetylation of caspase-2 by mass spectrometry. We observed that the mass to charge ratio (m/z) of the N-terminal peptide of caspase-2 is shifted as expected with an acetyl modification (Figures S2A and S2B). Furthermore, we found a 30% reduction in the amount of N-alpha-acetylated caspase-2 in NATH-deficient cells relative to control by subtiligase assay as well as mass spectrometry (Figure 2B and Figure S2D). These results support the conclusion that caspase-2 is N-alpha-acetylated by ARD1.

Protein N-Alpha-Acetylation Promotes the Assembly of Caspase-2 Complex

As caspase-2 is a substrate of ARD1 (Figure 2 and Figure S2) and the activation of caspase-2 is inhibited by ARD1 or NATH knockdown (Figure 3A), we asked how N-alpha-acetylation of caspase-2 might influence caspase activation.

First we conducted mutagenesis analysis of caspase-2 to disrupt protein N-alpha-acetylation. We replaced the third residue of caspase-2 with Pro (A3P) as the presence of Pro in this position inhibits protein N-alpha-acetylation. The 3P mutation has been previously demonstrated to inhibit N-alpha-acetylation of other substrates, known as the XPX rule (Goetze et al., 2009). We also replaced the second Ala for Ser as a control to maintain N-alpha-acetylation (A2S) as well as iMet removal (Arnesen et al., 2009; Goetze et al., 2009). Generation of these targeted substitutions allows us to definitively test whether subtiligase can differentiate between acetylated and unacetylated forms of caspase-2. An increase in subtiligase-mediated biotinylation of A3P was detected, while very little A2S or wildtype caspase-2 was detected after biotin pull down, consistent with acetylation as the explanation for the lower biotinylation levels (Figure 3B). A defect in N-alpha-acetylation of A3P caspase-2, but not WT and A2S caspase-2 was confirmed by mass spectrometry (Figures S2A-S2C). Thus, subtiligase is an effective tool for detecting unmodified protein N termini.

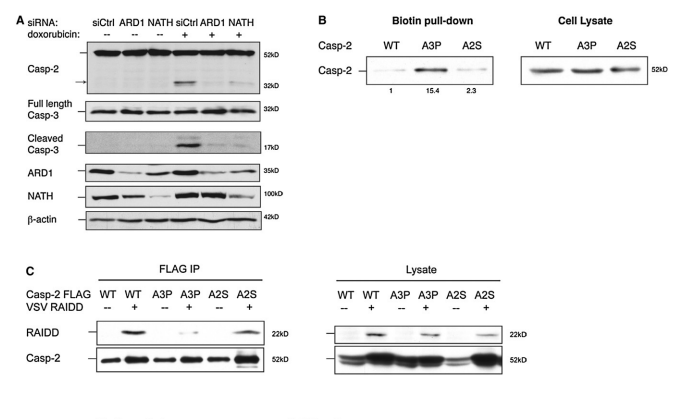
The caspase-2 scaffolding complex, which promotes caspase-2 activation, includes the adaptor protein, receptor-interacting protein (RIP)-associated ICH-1/CED-3 homologous protein with a death domain (RAIDD) (Duan and Dixit, 1997; Tinel and Tschopp, 2004). The ability of the N-terminal caspase-2 mutants to interact with RAIDD was assessed by coimmunoprecipitation. We found that RAIDD efficiently coimmunoprecipitated with WT and A2S but not with A3P caspase-2 (Figure 3C). This suggests that N-alpha-acetylation of caspase-2 facilitates its interaction with RAIDD.

Since acetyl-CoA is a key cofactor in N-alpha-acetylation, we speculated that the levels of N-alpha-acetylated caspase-2 might be dependent on expression of key metabolic enzymes that are responsible for production of cytoplasmic acetyl-CoA. To explore this question, we tested whether knockdown of ATP citrate lyase or acetyl-CoA synthetase (the two metabolic enzymes that utilize citrate and acetate, respectively) to generate acetyl-CoA, results in decreased levels of N-alpha-acetylated caspase-2. Indeed, we observed increased biotin labeling of caspase-2 in knockdown cells compared to control cells following subtiligase assay (Figure 3D). This suggests that caspase-2 is hypoacetylated when acetyl-CoA generation is reduced and therefore, protein N-alpha-acetylation is subject to metabolic regulation.

Regulation of Protein N-Alpha-Acetylation by Bcl-xL

Since decreased levels of protein N-alpha-acetylation leads to apoptotic deficiency, we reasoned that regulation of protein N-alpha-acetylation of certain apoptotic regulators might provide a mechanism to control apoptotic sensitivity. Bcl-xL, an antiapoptotic Bcl-2 family member, is known to have an effect on metabolism (Shimizu et al., 1999; Vander Heiden et al., 2001). We asked whether protein N-alpha-acetylation levels are sensitive to Bcl-xL expression using subtiligase assay. An increase in biotin labeling of caspase-2, -9, -3, and Bax was observed by Bcl-xL expression in 293T, HeLa, and Jurkat cells compared to that of control (Figures 4A, 4B, and 4C, respectively). Conversely, a decrease in biotin labeling was apparent in $bcl-x^{-/-}$ mouse embryonic fibroblasts (MEFs) compared to that of bcl-x+/+ MEFs (Figure 4D). Since Bcl-xL is known for maintaining mitochondrial integrity by blocking oligomerization of Bax/Bak, we measured the levels of protein N-alpha-acetylation in Bax/Bak-deficient cells. Surprisingly, the levels of protein N-alpha-acetylation were similar in bax-/-, bak-/-, or bax/bak^{-/-} (double knockout [DKO]) MEFs compared to that of WT MEFs by subtiligase assay (Figure 4E). This suggests that

(D) A schematic of detecting protein N-terminal peptide modifications by quantitative mass spectrometry. In brief, cells were treated with RNAi and lysed in NP-40 buffer. FLAG-tagged caspase-2 (active cysteine mutant C320G) was affinity purified with anti-FLAG agarose and eluted with FLAG peptide. Purified protein was digested by Lys-C, which cleaves after a lysine residue. Dimethyl labeling of peptides was conducted as previously described (Hsu et al., 2003; Khidekel et al., 2007). In brief, peptides were chemically modified with NaCNBH3 or NaCNBD3, followed by formaldehyde-H2 or formaldehyde-D2 treatment. Heavy or light modified peptides were mixed and analyzed by mass spectrometry. Ratios for levels of N-terminal acetylation were determined by normalization to 1:1 ratio of shared internal peptides (see Figure S1 for list of peptides). See also Figure S2.



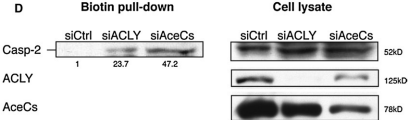


Figure 3. N-Alpha-Acetylation of Caspase-2 Is Required for Binding to RAIDD

(A) HeLa cells were transfected with siRNAs as indicated and treated with doxorubicin (5 μg/ml, 8 hr). Cell lysates were analyzed by SDS-PAGE, and caspase cleavage was assessed by immunoblot. Caspase-2 and caspase-3 activation is suppressed in ARD1 and NATH knockdown cells compared to that of control. Arrow indicates cleaved product of caspase-2.

(B) Subtiligase reaction was conducted on HEK293T cells transfected with FLAG-tagged WT, A3P, and A2S caspase-2 (active cysteine mutant C320G of caspase-2) as described in Figure 1. Enrichment of biotin-labeled A3P caspase-2 was observed compared to WT and A2S caspase-2. This suggests that A3P caspase-2 is hypoacetylated.

(C) HEK293T cells were transfected with FLAG-tagged WT, A3P, and A2S caspase-2 as well as with VSV-RAIDD as indicated. Caspase-2 was affinity purified with anti-FLAG agarose and eluted with FLAG peptide. The resulting eluants were subjected to SDS-PAGE for immunoblot analysis. RAIDD efficiently coimmuno-precipitates with WT and A2S but not with A3P caspase-2. Blots are representative of at least three independent experiments.

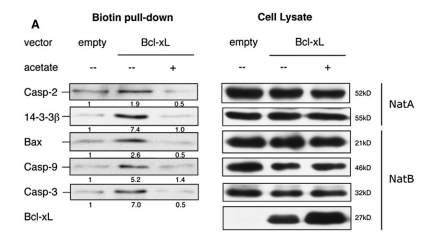
(D) Subtiligase reaction was conducted on lysates from HeLa cells transfected with siRNAs against acetyl-CoA synthetase or ATP citrate lyase as indicated. Increased biotin labeling of caspase-2 was observed in knockdown cells compared to control cells. See also Figure S2.

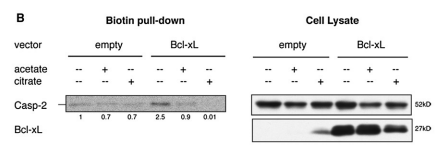
Bcl-xL-mediated regulation of protein N-alpha-acetylation is independent of Bax/Bak.

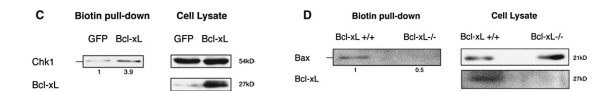
Recent studies show that histone lysine acetylation is dependent on acetyl-CoA production in yeast and mammalian cells (Takahashi et al., 2006; Wellen et al., 2009). However, we found that lysine acetylation of histone H3 and H4 were unaffected in Bcl-xL cells compared to control (Figure S3). This suggests

that histone lysine acetylation is not sensitive to the changes in acetyl-CoA levels associated with Bcl-xL expression.

We next tested whether protein N-alpha-acetylation levels in Bcl-xL cells are affected by changes in acetyl-CoA metabolism. Addition of acetate or citrate stimulates cytosolic acetyl-CoA production by acetyl-CoA synthetase or ATP citrate lyase, respectively (Sullivan et al., 1974a, 1974b). We verified that these







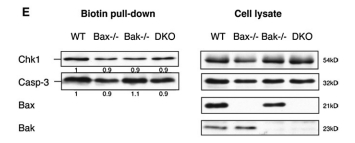


Figure 4. Metabolic Regulation of Protein N-Alpha-Acetylation by Bcl-xL

(A-C) HEK293T, HeLa, or Jurkat cells were generated to express empty GFP vector or BcI-xL. Cells were treated with acetate (50 mM, 24 hr) or citrate (10 mM, 24 hr) as indicated. Subtiligase reaction was conducted as described in Figure 1. Enrichment of biotin-labeled NatA and NatB substrates is observed in Bcl-xL cells compared to control cells. Biotin labeling is reduced to control levels by acetate or citrate treatment in Bcl-xL cells.

(D) Lysates generated from Bcl-xL^{+/+} or Bcl-xL^{-/-} MEFs were subjected to subtiligase reaction as described in Figure 1. Bax is hypoacetylated in Bcl-xL^{+/+} MEFs compared to Bcl-xL^{-/-} MEFs.

(E) Lysates generated from wild-type, Bax^{-/-}, Bak^{-/-}, or Bax/Bak^{-/-} (DKO) MEFs were subjected to subtiligase assay as described in Figure 1. Blots are representative of at least three independent experiments. See also Figure S3.

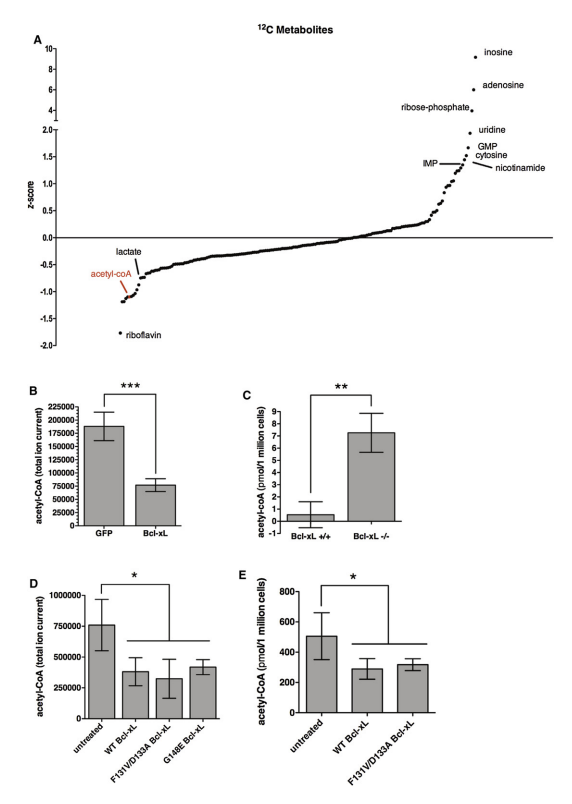


Figure 5. Bcl-xL Expression Reduces Acetyl-CoA Production

(A) Identification of metabolites using an AB/Sciex 5500 QTRAP mass spectrometer separated with normal phase HILIC NH2 chromatography using positive/ negative ion switching in Jurkat cells stably expressing GFP empty vector or Bcl-xL.

(B) Acetyl-CoA levels were measured in Jurkat cells stably expressing GFP empty vector or Bcl-xL by mass spectrometry. Peak areas of total ion current (TIC) were processed with MultiQuant 1.1 software. The average TIC for acetyl-CoA analyte is shown. Acetyl-CoA levels are lower in Bcl-xL cells relative to GFP cells. metabolites increase acetyl-CoA levels in mammalian cells (data not shown). Under metabolite treatment, protein N-alpha-acetylation levels were restored in Bcl-xL-expressing cells to that of control levels (Figures 4A and 4B). Thus, a reduction in acetyl-CoA production in Bcl-xL cells may be responsible for the observed hypoacetylation.

Metabolic Alterations Caused by Bcl-xL Expression

The expression of Bcl-xL is often elevated in tumors (Adams and Cory, 2007). To explore the effect of Bcl-xL on tumor metabolism, we conducted a systematic search using a combination of two-dimensinoal nuclear magnetic resonance (NMR) and mass spectrometry to identify metabolic changes associated with increased Bcl-xL expression. Principal component analysis (PCA) of one-dimensional proton spectra shows that the metabolome of Bcl-xL-expressing cells was significantly different from the metabolome of control cells (Figure S4).

We then used triple-quadruple mass spectrometry via selected reaction monitoring (SRM) to identify metabolite changes in Bcl-xL cells relative to GFP control cells as mass spectrometry is a more sensitive approach (Bajad et al., 2006; Lu et al., 2008). This is particularly relevant for intermediates of glucose metabolism as these metabolites are difficult to decipher by NMR due to their similar proton content. Thus, both NMR and mass spectrometry provide complementary approaches for a comprehensive understanding of the metabolite changes resulting from a specific perturbation. Indeed, we found that acetyl-CoA levels were decreased by 2-fold in Bcl-xL-expressing cells relative to GFP-expressing cells by mass spectrometry as well as an enzyme-based assay (Figures 5A and 5B). Conversely, acetyl-CoA levels were substantially increased in $bcl-x^{-/-}$ MEFs compared to $bcl-x^{+/+}$ MEFs (Figure 5C). These data provide strong evidence that Bcl-xL expression reduces the levels of acetyl-CoA, suggesting that reduced levels of acetyl-CoA in Bcl-xL-overexpressing cells leads to hypoacetylation.

Since bax/bak DKO cells are not defective in protein N-alphaacetylation, we reasoned that Bcl-xL might be able to negatively regulate the levels of acetyl-CoA independent of Bax/Bak binding. Cheng et al. reported that certain Bcl-xL mutants, such as F131V/D133A and G148E, are unable to bind to Bax or Bak but nevertheless retain 70%-80% antiapoptotic activity of WT Bcl-xL (Cheng et al., 1996). We measured acetyl-CoA levels in cells expressing WT Bcl-xL or these specific Bcl-xL mutants. A similar reduction in acetyl-coA levels was observed in cells expressing these Bcl-xL mutants and in cells expressing WT Bcl-xL (Figures 5D and 5E). Thus, Bcl-xL's metabolic function in regulating the levels of acetyl-CoA does not depend on its interaction with Bax/Bak.

As the majority of the cellular acetyl-group in acetyl-CoA is produced from glucose (DeBerardinis et al., 2007), we asked whether glucose metabolism might be altered in Bcl-xLexpressing cells. We fed Bcl-xL cells uniformly labeled ¹³C-glucose to differentiate glucose-derived metabolites from those derived from other carbon sources (Figure 6A and Table S1). We found that the levels of glucose-derived citrate were decreased by approximately 25% in Bcl-xL-expressing cells relative to control (Student's t test, p value < 0.05; Figure 6B and Table S1). As citrate is the direct precursor of cytoplasmic pools of acetyl-CoA, the lower levels of glucose-derived citrate might explain the decrease in acetyl-CoA levels observed in Bcl-xL-expressing cells. Consistent with this hypothesis, levels of alpha-ketoglutarate, which is also derived from citrate, were lower in Bcl-xL-expressing cells relative to control (Figure 6B and Table S1).

Since metabolite addition rescues the defect on protein N-alpha-acetylation by Bcl-xL (Figures 4A and 4B), we asked whether these metabolites could alter cell survival that is supported by Bcl-xL expression. Bcl-xL expression effectively protects against a wide range doses of doxorubicin (Figures 7A-7D). Remarkably, increasing levels of citrate or acetate sensitized HeLa cells stably expressing Bcl-xL to doxorubicininduced cell death compared to that of untreated cells (Figure 7B). This corresponds with a 2-fold increase in caspase activity (Figure 7D). Importantly, RNAi against acetyl-CoA synthetase or ATP citrate lyase completely suppressed the sensitization to doxorubicin elicited by addition of acetate or citrate, respectively (Figure 7E). This indicates that metaboliteinduced apoptotic sensitization of cells expressing Bcl-xL specifically results from changes in acetyl-CoA production.

The above data suggest that Bcl-xL may mediate apoptosis resistance through two parallel pathways by inhibiting Bax/Bak oligomerization and by downregulating protein N-alpha-acetylation. We therefore directly tested whether the effects of inhibiting Bax and ARD1 are additive in protecting against apoptosis. We found that double knockdown of both ARD1 and Bax indeed provided enhanced protection against apoptosis compared to that of knockdown individually, which was especially significant at higher concentrations of doxorubicin (Figure 7F). This finding supports the notion that Bcl-xL has dual functions in regulating protein N-alpha-acetylation levels and Bax/Bak oligomerization.

DISCUSSION

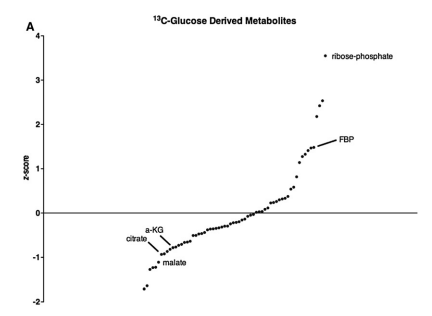
The ability to rapidly assess protein modifications immunologically has been essential for exploring the significance and regulation of multiple protein posttranslational modifications such as phosphorylation, histone methylation, and acetylation. Since an antibody for protein N-alpha-acetylation does not exist, the ability to assess this modification was severely limited. In this regard, the subtiligase assay as described in the present study provides a powerful tool to allow us to rapidly assess the endogenous levels of protein N-alpha-acetylation. Using this assay,

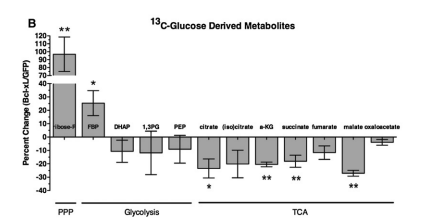
⁽C) Acetyl-CoA levels were measured in Bcl-xL+/+ or Bcl-xL-/- MEFs with an enzyme-based assay (Abcam). Metabolite concentration was determined by generation of a standard curve. Bcl-xL deficiency results in higher acetyl-CoA levels compared to control.

⁽D) 293T cells were transiently transfected with empty vector, WT Bcl-xL, or specific Bcl-xL mutants (F131V/D133A or G148E). Acetyl-CoA levels were measured by mass spectrometry. WT or mutant Bcl-xL cells show a decrease in acetyl-CoA levels compared to empty vector cells.

⁽E) Acetyl-CoA levels were measured in 293T cells transiently expressing WT or mutant Bcl-xL as described in (C).

Data are represented as mean \pm SD (n = 3). (Student's t test: *p < 0.05; **p < 0.01; ***p < 0.001.) See also Figure S4.





we discovered that protein N-alpha-acetylation status is reduced in cells overexpressing Bcl-xL. Furthermore we show that protein N-alpha-acetylation is sensitive to acute changes in acetyl-CoA availability.

Our study directly links a specific metabolite, acetyl-CoA, to apoptotic sensitivity and supports a growing number of studies that describe a role for cell metabolism in controlling apoptosis (Nutt et al., 2005, 2009; Schafer et al., 2009; Vaughn and Deshmukh, 2008). Interestingly, the cellular levels of acetyl-CoA are sensitive to Bcl-xL status in a Bax/Bak-independent manner because expression of Bcl-xL mutants that are unable to bind to Bax or Bak (Cheng et al., 1996) can also affect acetyl-CoA levels to the same extent as that of wild-type Bcl-xL. Hardwick and colleagues demonstrated that these Bcl-xL mutants preserve 70%-80% antiapoptotic activity of WT Bcl-xL despite their inability to bind to Bax or Bak (Cheng et al., 1996). Thus, inhibition of acetyl-CoA production might provide an additional mechanism for Bcl-xL to protect against apoptosis in a Bax/Bak-independent manner. Taken together, these data

Figure 6. Metabolite Profiling of Glucose-Derived Intermediates in Bcl-xL Cells by Mass Spectro-

(A) Jurkat cells stably expressing GFP empty vector or Bcl-xL were fed uniformly labeled 13C-glucose, and metabolites were isolated by methanol extraction. Lysates were subjected to mass spectrometry as described in Figure 3A.

(B) Percent change in ¹³C-glucose derived metabolites of the pentose phosphate pathway (PPP), glycolysis, and TCA cycle. Fructose-1,6 bisphosphate (FBP), dihydroxyacetone-phosphate (DHAP), 1,3-bis-phosphoglycerate (1,3PG), phosphoenolpyruvate (PEP), α-ketoglutarate (a-KG). Data are represented as mean ± SD (n = 3). (Student's t test: *p < 0.05; **p < 0.01; ***p < 0.001.) See also Table S1.

suggest that Bcl-xL might protect against apoptosis through two parallel mechanisms: by directly binding and inhibiting Bax/Bak oligomerization and by regulating mitochondrial metabolism, which leads to reduced levels of acetyl-coA and protein N-alpha-acetylation. We conclude that Bcl-xL integrates metabolism to apoptotic resistance by modulating acetyl-CoA levels.

Previous studies show that Bcl-xL directly binds to the voltage-dependent anion channel (VDAC), a component of the mitochondrial permeability transition pore, which controls mitochondrial metabolite exchange (Shimizu et al., 1999; Vander Heiden et al., 2001). It is possible that Bcl-xL expression may alter levels of acetyl-coA by regulating mitochondrial membrane permeability. Citrate carrier (CiC), a nuclear-encoded protein located in the mitochondrial inner membrane and a member of the mitochondrial carrier family, is responsible for the efflux of acetyl-CoA from the mitochon-

dria to the cytosol in the form of citrate (Gnoni et al., 2009; Kaplan et al., 1993). We found that the levels of glucose-derived citrate were decreased by approximately 25% in Bcl-xL-expressing cells relative to the control. This reduction in citrate levels could explain the observed decrease in acetyl-CoA levels in Bcl-xLexpressing cells and contribute to the antiapoptotic function of Bcl-xL. Indeed, addition of citrate to Bcl-xL-expressing cells leads to increased protein N-alpha-acetylation and sensitization of these cells to apoptosis. Perturbations in acetyl-CoA production may extend to other oncogenic contexts beyond that of Bcl-xL. For example, the levels of glucose-derived acetyl-CoA were found to be approximately 20% higher in myc+/+ cells relative to $myc^{-/-}$ cells (Morrish et al., 2009). An increase in acetyl-CoA levels might contribute to enhanced apoptotic sensitivity of cells overexpressing c-Myc (Evan et al., 1992). We propose that the basal levels of acetyl-CoA may influence the apoptotic threshold in multiple oncogenic contexts.

The ability of Bcl-xL to control the levels of acetyl-CoA and protein-N-acetylation provides a clear example by which

metabolism is mechanistically linked with apoptotic sensitivity. Loss-of-function ard1 mutant yeast are specifically defective in alpha-factor response but not to a-factor (Whiteway and Szostak, 1985), indicating that protein N-alpha-acetylation status can dictate a specific cellular behavior or process. Since protein N-alpha-acetylation affects a large number of cellular proteins, we speculate that metabolic regulation of this process exerts its control on cellular processes through regulating a group(s) of proteins rather than individual proteins. ARD1deficient mammalian cells are defective in the activation of caspase-2, caspase-3, and caspase-9 in response to DNA damage (Yi et al., 2007). Consistently, N-alpha-acetylation of multiple caspases, including caspase-2, caspase-3, and caspase-9, was reduced in Bcl-xL-overexpressing cells. It is possible that defects in N-alpha-acetylation of multiple caspases, which may negatively regulate their activation, contribute to apoptotic resistance of ARD1-deficient cells as well as Bcl-xL-overexpressing cells. Thus, the N-alpha-acetylation status of multiple proteins that are involved in a particular pathway may collectively determine a specific physiological outcome. In this regard, the cofactor for the Nat complexes, acetyl-CoA, serves as a signaling molecule that functions as an important liaison between metabolism and multiple cellular processes.

EXPERIMENTAL PROCEDURES

Cell Death Assays

For RNAi studies, low-passage HeLa cells were transiently transfected (5 x 10⁵/well in 96-well plates) with a pool of four small interfering RNAs (siRNAs) (predesigned ON-TARGETplus siRNAs available from Dharmacon, 50 nM) with Oligofectamine transfection reagent (Invitrogen). After a 48 hr incubation, cells were treated with doxorubicin (concentrations and incubation times are noted in figure legends). siRNAs were tested in triplicate for each independent experiment. For detection of caspase cleavage by western blot, HeLa cells were transfected as described above (1.5 \times 10⁵/well in 6-well plates) followed by treatment with doxorubicin. Cells were lysed directly in SDS samples buffer and subjected to SDS-PAGE and western blot analysis via standard procedures.

For metabolite sensitization experiments, HeLa cells stably expressing GFP or Bcl-xL (3 \times 10⁵/well in 96-well plates) were pretreated with acetate (50 mM) or citrate (10 m) for 24 hr, followed by treatment with doxorubicin as indicated. Cell viability was determined by measurement of cellular ATP levels (CellTiter-Glo Luminescent Cell Viability Assay, Promega). Caspase-3/7 activity was quantified with a luciferin-labeled DEVD peptide substrate (Caspase-Glo 3/7 Assay, Promega). Luminescence was measured with a Wallac Victor2 plate reader.

Subtiligase Biotinylation Experiments Synthesis of Peptide Substrate

The biotinylated peptide substrate for subtiligase with a TEV protease cleavage site, biotin-ahx-ahx-GGTENLYFQSY-glc-Y-NH2, was synthesized manually on a rink amide MBHA resin according to standard 9-Fluorenylmethoxycarbonyl (Fmoc) chemistry-based solid-phase peptide synthesis protocols (Mahrus et al., 2008). The crude peptide was purified with a C₁₈ semipreparative reverse-phase column on a Waters HPLC system. The identity of the purified product was confirmed by ESI-MS (isotopic molecular weight [MW] calculated 1949.9 Da, m/z [M+H]⁺ found 1951.0).

The peptide substrate could be made more soluble by incorporating D-arginine residues into the sequence (Yoshihara et al., 2008). So a more soluble form of the substrate, biotin-ahx-ahx-dRdRdR-ahx-ahx-GGTENLYFQSYglc-Y-NH2, was also synthesized, and its integrity was verified by HPLC and ESI-MS (MW calculated 2644.4 Da, m/z [M+H]+ found 2645.8).

Expression and Purification of Subtiligase

The expression construct for subtiligase was prepared with the plasmid pBS42 (ATCC 37279) according to published procedures (Wells et al., 1983), except that a His₆ tag was added to the C terminus (Tan et al., 2007). The gene of the subtiligase variant contains point mutations S221C, P225A, M124L, and S125A on wild-type subtilisin BPN'. The recombinant protein was expressed in B. subtilis RIK1285, which is deficient in the production of neutral and alkaline proteases. Purification of subtiligase was performed as previously described (Abrahmsén et al., 1991), except that a Co2+ affinity chromatography step was used instead of ion-exchange chromatography (Tan et al., 2007). The affinity-purified subtiligase was desalted using a PD-10 column (Amersham) with deionized H₂O. Aliquots of the desalted enzyme solution were flash frozen, lyophilized, and stored at -80°C until utilized. The prepared subtiligase was analyzed by SDS-PAGE gel and MALDI-TOF MS, which confirmed its purity and identity. The ligase activity of the enzyme preparation was confirmed in model ligation reactions using known peptide substrates for subtiligase.

Subtiligase Ligation Reaction

Cells used for subtiligase assay were plated at 50%-60% confluency to support exponential growth, followed by pretreatment with acetate (50 mM) or citrate (10 mM) for 24 hr as indicated. Cells were lysed in 0.2% Tween 20 and 0.2% Triton X-100 buffer, and the resulting lysates were used for subtiligase reaction that includes 1 mM purified subtiligase, 1 mM purified biotinpeptide, and 2 mM DTT as previously described (Mahrus et al., 2008). Reactions were allowed to proceed for 1 hr at room temperature. Biotinylated proteins were affinity purified with Neutravidin agarose at 4° overnight (Thermo Scientific). The following day, agarose was pelleted and washed three times in lysis buffer. Purified proteins were eluted directly in 2× SDS sample buffer, and eluants were analyzed by SDS PAGE.

¹³C Mass Spectrometry Sample Preparation

Three replicates of Jurkat cells stably expressing GFP or Bcl-xL (5 \times 10 6 for each replicate) were washed twice in PBS and resuspended in RPMI medium with glutamine, 10% dialyzed NCS, and 10 mM uniformly labeled ¹³C-glucose (Cambridge Isotope Laboratories), followed by incubation for 24 hr. Cells were washed twice, and metabolites were extracted in 3 ml 80% ice-cold methanol. Insoluble material in lysates was pelleted at 13.000 q for 10 min, and methanol from the resulting supernatant was evaporated. Samples were resuspended with 20 µl HPLC grade water for mass spectrometry.

¹²C and ¹³C Mass Spectrometry Data Acquisition and Analysis

Seven microliters of 20 µl were injected using a 5500 QTRAP mass spectrometer (Applied Biosystems/MDS Sciex) equipped with a Prominence UFLC HPLC system (Shimadzu) via SRM of a total of 249 endogenous metabolites for ¹²C analyses of GFP and BcL-xL samples. For analyses of ¹³C-labeled GFP and BcL-xL samples, 153 endogenous metabolites were targeted via SRM. Reliable quantitative data are only acquired from approximately 60% of the targeted metabolites (~160 metabolites from the ¹²C method and \sim 55% from the 13 C method (\sim 80 metabolites). Some metabolites were targeted in both positive and negative ion mode, including acetyl-CoA. ESI voltage was 5000 V in positive ion mode and -4500 V in negative ion mode. The dwell time was 5 ms per SRM transition, and the collision energy was optimized for each SRM transition. Total cycle time was 2.09 s for the ¹²C method and 1.24 s for the ¹³C method. Scheduled SRMs were not utilized. Samples were delivered to the MS via normal phase chromatography using a 2.0 mm i.d × 10 cm HILIC Luna NH2 column (Phenomenex) at 250 µl/min at basic pH using positive and negative ion switching within the same 30 min LC/MS/MS analytical run. Gradients were run starting from 85% buffer B (HPLC grade acetonitrile) to 42% B from 0-5 min; 42% B to 0% B from 5-16 min; 0% B was held from 16-24 min; 0% B to 85% B from 24-25 min; 85% B was held for 7 min to re-equilibrate the column. Buffer A was composed of 20 mM ammonium hydroxide/20 mM ammonium acetate in 95/5 water/acetonitrile. All metabolomic measurements were performed in triplicate. Peak areas from the total ion current (TIC) for each metabolite SRM transition were integrated using MultiQuant v1.1 software (Applied Biosystems).

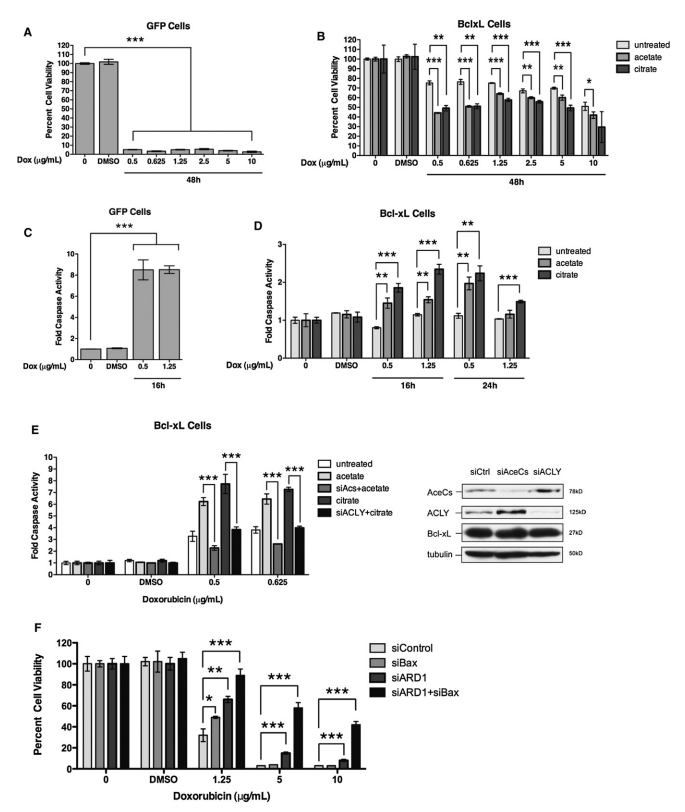


Figure 7. Metabolic Function of Bcl-xL Contributes to Cell Survival

(A–D) HeLa cells stably expressing empty GFP vector or Bcl-xL were treated with doxorubicin as indicated. Cell viability was determined by measurement of cellular ATP levels with a luminescence-based assay (Promega). Caspase activity was determined by measurement of the cleavage of a luciferin substrate containing the caspase cleavage site, DEVD (Promega).

Statistical analysis was conducted with an unpaired, two-tailed Student's t test.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, four figures, and one table and can be found with this article online at doi:10. 1016/i.cell.2011.06.050.

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REFERENCES

Abrahmsén, L., Tom, J., Burnier, J., Butcher, K.A., Kossiakoff, A., and Wells, J.A. (1991). Engineering subtilisin and its substrates for efficient ligation of peptide bonds in aqueous solution. Biochemistry 30, 4151–4159.

Adams, J.M., and Cory, S. (2007). The Bcl-2 apoptotic switch in cancer development and therapy. Oncogene 26, 1324-1337.

Ametzazurra, A., Larrea, E., Civeira, M.P., Prieto, J., and Aldabe, R. (2008). Implication of human N-alpha-acetyltransferase 5 in cellular proliferation and carcinogenesis. Oncogene 27, 7296-7306.

Arnesen, T., Thompson, P.R., Varhaug, J.E., and Lillehaug, J.R. (2008). The protein acetyltransferase ARD1: a novel cancer drug target? Curr. Cancer Drug Targets 8, 545-553.

Arnesen, T., Van Damme, P., Polevoda, B., Helsens, K., Evjenth, R., Colaert, N., Varhaug, J.E., Vandekerckhove, J., Lillehaug, J.R., Sherman, F., and Gevaert, K. (2009). Proteomics analyses reveal the evolutionary conservation and divergence of N-terminal acetyltransferases from yeast and humans. Proc. Natl. Acad. Sci. USA 106, 8157-8162.

Bajad, S.U., Lu, W., Kimball, E.H., Yuan, J., Peterson, C., and Rabinowitz, J.D. (2006). Separation and quantitation of water soluble cellular metabolites by hydrophilic interaction chromatography-tandem mass spectrometry. J. Chromatogr. A 1125, 76-88.

Boise, L.H., González-García, M., Postema, C.E., Ding, L., Lindsten, T., Turka, L.A., Mao, X., Nuñez, G., and Thompson, C.B. (1993). bcl-x, a bcl-2-related gene that functions as a dominant regulator of apoptotic cell death. Cell 74, 597-608

Cheng, E.H., Levine, B., Boise, L.H., Thompson, C.B., and Hardwick, J.M. (1996). Bax-independent inhibition of apoptosis by Bcl-XL. Nature 379,

DeBerardinis, R.J., Mancuso, A., Daikhin, E., Nissim, I., Yudkoff, M., Wehrli, S., and Thompson, C.B. (2007). Beyond aerobic glycolysis: transformed cells can engage in glutamine metabolism that exceeds the requirement for protein and nucleotide synthesis. Proc. Natl. Acad. Sci. USA 104, 19345-19350.

Duan, H., and Dixit, V.M. (1997). RAIDD is a new 'death' adaptor molecule. Nature 385, 86-89

Evan, G.I., Wyllie, A.H., Gilbert, C.S., Littlewood, T.D., Land, H., Brooks, M., Waters, C.M., Penn, L.Z., and Hancock, D.C. (1992). Induction of apoptosis in fibroblasts by c-myc protein. Cell 69, 119-128.

Gnoni, G.V., Priore, P., Geelen, M.J., and Siculella, L. (2009). The mitochondrial citrate carrier: metabolic role and regulation of its activity and expression. IUBMB Life 61. 987-994.

Goetze, S., Qeli, E., Mosimann, C., Staes, A., Gerrits, B., Roschitzki, B., Mohanty, S., Niederer, E.M., Laczko, E., Timmerman, E., et al. (2009). Identification and functional characterization of N-terminally acetylated proteins in Drosophila melanogaster. PLoS Biol. 7, e1000236.

Gottlieb, E., Vander Heiden, M.G., and Thompson, C.B. (2000). Bcl-x(L) prevents the initial decrease in mitochondrial membrane potential and subsequent reactive oxygen species production during tumor necrosis factor alphainduced apoptosis. Mol. Cell. Biol. 20, 5680-5689.

Hsu, J.L., Huang, S.Y., Chow, N.H., and Chen, S.H. (2003). Stable-isotope dimethyl labeling for quantitative proteomics. Anal. Chem. 75, 6843-6852.

Kaplan, R.S., Mayor, J.A., and Wood, D.O. (1993). The mitochondrial tricarboxylate transport protein. cDNA cloning, primary structure, and comparison with other mitochondrial transport proteins. J. Biol. Chem. 268, 13682-13690.

Khidekel, N., Ficarro, S.B., Clark, P.M., Bryan, M.C., Swaney, D.L., Rexach, J.E., Sun, Y.E., Coon, J.J., Peters, E.C., and Hsieh-Wilson, L.C. (2007). Probing the dynamics of O-GlcNAc glycosylation in the brain using quantitative proteomics. Nat. Chem. Biol. 3, 339-348.

Lu, W., Bennett, B.D., and Rabinowitz, J.D. (2008). Analytical strategies for LC-MS-based targeted metabolomics. J. Chromatogr. B Analyt. Technol. Biomed. Life Sci. 871, 236-242.

Mahrus, S., Trinidad, J.C., Barkan, D.T., Sali, A., Burlingame, A.L., and Wells, J.A. (2008). Global sequencing of proteolytic cleavage sites in apoptosis by specific labeling of protein N termini. Cell 134, 866-876.

Martin, H., Patel, Y., Jones, D., Howell, S., Robinson, K., and Aitken, A. (1993). Antibodies against the major brain isoforms of 14-3-3 protein. An antibody specific for the N-acetylated amino-terminus of a protein. FEBS Lett. 331, 296-303.

(A and B) Doxorubicin treatment efficiently kills GFP cells but not Bcl-xL cells. Pretreatment with acetate (50 mM) or citrate (10 mM) for 24 hr sensitizes Bcl-xL cells to doxorubicin-induced cell death.

(C and D) Doxorubicin treatment induces substantial caspase activity in GFP cells but is completely suppressed in Bcl-xL cells. Pretreatment of Bcl-xL cells with acetate or citrate enhances caspase activity induced by doxorubicin treatment.

(E) Bcl-xL cells were transiently transfected with a pool of siRNAs (50 nM, Dharmacon) as indicated, followed by pretreatment with acetate or citrate prior to doxorubicin treatment. Metabolite enhancement of doxorubicin-induced caspase activity in Bcl-xL cells is suppressed to control levels by RNAi against acetyl-CoA synthetase (Acs) or ATP citrate lyase (ACLY), respectively. Immunoblot shows extent of knockdown.

(F) HeLa cells were transiently transfected with a pool of siRNAs as indicated. Double knockdown of ARD1 and Bax provides additive protection against doxorubicin-induced cell death compared to single knockdown of these genes.

Graphs are representative of three independent experiments. Data are represented as mean ± SD (n = 3). (Student's t test: *p < 0.05; **p < 0.01; ***p < 0.001.)

Martinez, A., Traverso, J.A., Valot, B., Ferro, M., Espagne, C., Ephritikhine, G., Zivy, M., Giglione, C., and Meinnel, T. (2008). Extent of N-terminal modifications in cytosolic proteins from eukaryotes. Proteomics 8, 2809-2831.

Morrish, F., Isern, N., Sadilek, M., Jeffrey, M., and Hockenbery, D.M. (2009). c-Myc activates multiple metabolic networks to generate substrates for cellcycle entry. Oncogene 28, 2485-2491.

Nutt, L.K., Margolis, S.S., Jensen, M., Herman, C.E., Dunphy, W.G., Rathmell, J.C., and Kornbluth, S. (2005). Metabolic regulation of oocyte cell death through the CaMKII-mediated phosphorylation of caspase-2. Cell 123, 89-103.

Nutt, L.K., Buchakjian, M.R., Gan, E., Darbandi, R., Yoon, S.Y., Wu, J.Q., Miyamoto, Y.J., Gibbons, J.A., Andersen, J.L., Freel, C.D., et al. (2009). Metabolic control of oocyte apoptosis mediated by 14-3-3zeta-regulated dephosphorylation of caspase-2. Dev. Cell 16, 856-866.

Polevoda, B., and Sherman, F. (2003). Composition and function of the eukaryotic N-terminal acetyltransferase subunits. Biochem. Biophys. Res. Commun. 308, 1-11.

Schafer, Z.T., Grassian, A.R., Song, L., Jiang, Z., Gerhart-Hines, Z., Irie, H.Y., Gao, S., Puigserver, P., and Brugge, J.S. (2009). Antioxidant and oncogene rescue of metabolic defects caused by loss of matrix attachment. Nature 461, 109-113,

Shimizu, S., Narita, M., and Tsujimoto, Y. (1999). Bcl-2 family proteins regulate the release of apoptogenic cytochrome c by the mitochondrial channel VDAC. Nature 399, 483-487.

Starheim, K.K., Arnesen, T., Gromyko, D., Ryningen, A., Varhaug, J.E., and Lillehaug, J.R. (2008). Identification of the human N(alpha)-acetyltransferase complex B (hNatB): a complex important for cell-cycle progression. Biochem. J. 415, 325-331.

Starheim, K.K., Gromyko, D., Velde, R., Varhaug, J.E., and Arnesen, T. (2009). Composition and biological significance of the human Nalpha-terminal acetyltransferases. BMC Proc 3 (Suppl 6), S3.

Sullivan, A.C., Triscari, J., Hamilton, J.G., and Miller, O.N. (1974a). Effect of (-)-hydroxycitrate upon the accumulation of lipid in the rat. II. Appetite. Lipids 9, 129-134.

Sullivan, A.C., Triscari, J., Hamilton, J.G., Miller, O.N., and Wheatley, V.R. (1974b). Effect of (-)-hydroxycitrate upon the accumulation of lipid in the rat. I. Lipogenesis. Lipids 9, 121-128.

Takahashi, H., McCaffery, J.M., Irizarry, R.A., and Boeke, J.D. (2006). Nucleocytosolic acetyl-coenzyme a synthetase is required for histone acetylation and global transcription, Mol. Cell 23, 207-217.

Tan, X.H., Wirjo, A., and Liu, C.F. (2007). An enzymatic approach to the synthesis of peptide thioesters: mechanism and scope. ChemBioChem 8, 1512-1515

Tinel, A., and Tschopp, J. (2004). The PIDDosome, a protein complex implicated in activation of caspase-2 in response to genotoxic stress. Science 304. 843-846.

Vander Heiden, M.G., Chandel, N.S., Schumacker, P.T., and Thompson, C.B. (1999). Bcl-xL prevents cell death following growth factor withdrawal by facilitating mitochondrial ATP/ADP exchange. Mol. Cell 3, 159-167.

Vander Heiden, M.G., Li, X.X., Gottleib, E., Hill, R.B., Thompson, C.B., and Colombini, M. (2001). Bcl-xL promotes the open configuration of the voltage-dependent anion channel and metabolite passage through the outer mitochondrial membrane. J. Biol. Chem. 276, 19414-19419.

Vander Heiden, M.G., Cantley, L.C., and Thompson, C.B. (2009). Understanding the Warburg effect: the metabolic requirements of cell proliferation. Science 324, 1029-1033.

Vaughn, A.E., and Deshmukh, M. (2008). Glucose metabolism inhibits apoptosis in neurons and cancer cells by redox inactivation of cytochrome c. Nat. Cell Biol. 10, 1477-1483.

Wellen, K.E., Hatzivassiliou, G., Sachdeva, U.M., Bui, T.V., Cross, J.R., and Thompson, C.B. (2009). ATP-citrate lyase links cellular metabolism to histone acetylation. Science 324, 1076-1080.

Wells, J.A., Ferrari, E., Henner, D.J., Estell, D.A., and Chen, E.Y. (1983). Cloning, sequencing, and secretion of Bacillus amyloliquefaciens subtilisin in Bacillus subtilis. Nucleic Acids Res. 11, 7911-7925.

Whiteway, M., and Szostak, J.W. (1985). The ARD1 gene of yeast functions in the switch between the mitotic cell cycle and alternative developmental pathwavs. Cell 43, 483-492.

Yi, C.H., Sogah, D.K., Boyce, M., Degterev, A., Christofferson, D.E., and Yuan, J. (2007). A genome-wide RNAi screen reveals multiple regulators of caspase activation. J. Cell Biol. 179, 619-626.

Yoshihara, H.A., Mahrus, S., and Wells, J.A. (2008). Tags for labeling protein N-termini with subtiligase for proteomics. Bioorg. Med. Chem. Lett. 18, 6000-6003.