DNA damage response and MCL-1 destruction initiate apoptosis in adenovirus-infected cells

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Expression of adenovirus E1A deregulates cell proliferation to facilitate viral DNA replication, prompting the initiation of apoptosis signaled primarily through proapoptotic BAK in productively infected cells. We demonstrate here that in uninfected cells, BAK is complexed with the anti-apoptotic BCL-2 family member Myeloid Cell Leukemia 1 (MCL-1). E1A expression during infection resulted in the specific down-regulation of MCL-1 through destabilization of the protein and loss of the mRNA. Upon loss of the MCL-1-BAK complex, BAK complexed with either BAX in proapoptotic E1B mutant adenovirus-infected cells, or with the adenovirus BCL-2 homolog E1B 19K in cells infected with the wild-type virus in which apoptosis is inhibited. Loss of MCL-1 was required to initiate the apoptotic pathway in infected cells as restoration of MCL-1 expression rescued infected cells from E1A-induced apoptosis. Analogous to E1A expression, DNA damage down-regulates MCL-1, and adenovirus infection resulted in the accumulation of phosphorylated H2AX and ataxia-telangiectasia mutant protein (ATM), hallmarks of DNA double-strand breaks. Thus, MCL-1 may function by maintaining BAK in an inactive state, and the loss of MCL-1 upon activation of the DNA damage response, perhaps through replication stress induced in virus infected cells, may be required to initiate the apoptotic response.

Keywords: Apoptosis, MCL-1, E1B 19K, adenovirus, DNA damage, BAK

Received September 30, 2003; revised version accepted October 22, 2003.
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the E1B 19K protein blocks apoptosis. Yeast two-hybrid screening identified proapoptotic BAK and BAX as E1B 19K-interacting proteins, and E1B 19K expression blocks cell death induced by BAK or BAX overexpression (Farow et al. 1995; Han et al. 1996). The E1B 19K protein interacts with and inhibits the endogenous activated forms of proapoptotic BAK and BAX (Perez and White 2000; Sundararajan and White 2001; Sundararajan et al. 2001; Cuconati et al. 2002) that are required for releasing apoptogenic mitochondrial proteins [cytochrome and Smac/DIABLO, among others], which serve to activate cysteine proteases of the caspase family that implement apoptosis [Lindsten et al. 2000; Wei et al. 2001; Zong et al. 2001; Degenhardt et al. 2002]. BAK and BAX activation can occur through direct interaction with proapoptotic BH3-only proteins such as tBID, and/or through inactivation of pro-survival BCL-2 family members. Activation of BAK and BAX is thought to occur through changes in their protein conformation, followed by oligomerization into a high-molecular weight protein complex that is associated with compromised mitochondrial membrane permeability [Korsmeyer et al. 2000; Cory and Adams 2002]. For example, during tumor necrosis factor (TNF)-α-mediated apoptosis, caspase-8 activation by the receptor complex and cleavage of BID to truncated tBID propagates death signaling to mitochondria. In adenovirus-infected cells, tBID interacts with and activates both BAK and BAX, and the E1B 19K protein binds to activated BAK and BAX and prevents their co-oligomerization. This E1B 19K–BAK and E1B 19K–BAX interaction prevents the release of proapoptotic mitochondrial proteins, thereby inhibiting caspase-9 and caspase-3 activation and apoptosis [Perez and White 2000; Sundararajan and White 2001; Sundararajan et al. 2001]. The interaction of E1B 19K with BAK and BAX is sufficient to block TNF-α-mediated apoptosis as BAK and BAX-deficient cells are resistant to TNF-α (Degenhardt et al. 2002). Because BAK and BAX are general effectors of apoptotic signaling, their inhibition by E1B 19K could account for its function, not only in apoptosis mediated by TNF-α, but also by E1A.

E1A-mediated apoptosis in cells infected with an E1B 19K mutant adenovirus similarly results in BAK and BAX activation, co-oligomerization, mitochondrial release, caspase activation, and apoptosis (Cuconati et al. 2002). In cells infected with the wild-type virus, in which apoptosis is blocked by E1B 19K expression, BAK activation still occurs, but the interaction of E1B 19K with activated BAK prevents activation of BAX, co-oligomerization of BAK and BAX, mitochondrial protein release, caspase activation, and apoptosis (Cuconati et al. 2002). Thus, signaling of apoptosis by E1A in adenovirus-infected cells primarily goes through BAK, which is targeted by the E1B 19K protein. However, deficiency in both BAK and BAX is required to block apoptosis in cells infected with an E1B 19K mutant adenovirus, suggesting that in the absence of BAK, there is likely a salvage pathway for BAX activation and apoptosis induction (Cuconati et al. 2002). This varies from TNF-α death signaling, in which tBID activates both BAK and BAX and E1B 19K interacts with both. There is no caspase-8 activation or conversion of BID to tBID during E1A-mediated apoptosis in adenovirus-infected cells, suggesting that the mechanism of BAK activation by E1A was distinct from that signaled by TNF-α (Perez and White 2000; Cuconati et al. 2002). Nonetheless, BAK emerged as the key mediator of E1A-mediated apoptosis during productive virus infection, and we began a search for the mechanism of BAK activation.

As BAK [and BAX] activation can occur through induction of and interaction with proapoptotic BH3-only proteins, or through the loss of function and interaction with anti-apoptotic BCL-2 family members, infected cells were screened for changes in the levels of these proteins and for their interaction with BAK. We report here that BAK interacted with anti-apoptotic BCL-2 family member MCL-1 in healthy, uninfected cells. Adenovirus infection and E1A expression repressed MCL-1 expression, which was associated with the induction of BAK–BAX complex formation and apoptosis that was blocked by restoring MCL-1 levels. In infected cells in which apoptosis was blocked by E1B 19K expression, MCL-1 down-regulation still occurred, but E1B 19K interacted with BAK, and BAK–BAX interaction was prevented. As MCL-1 down-regulation initiates the apoptotic response to ultraviolet and γ irradiation (Nijhawan et al. 2003), we examined infected cells for induction of a DNA-damage response. Adenovirus infection induced phosphorylation of H2AX and ATM, even when apoptosis was inhibited by E1B 19K expression in cells infected with wild-type virus. Thus, E1A expression may trigger a DNA-damage response, resulting in not only p53 stabilization, but also in MCL-1 down-regulation, which then serves as a priming event for BAK activation and apoptosis. This may necessitate that adenovirus encode a BAK and BAX inhibitor in addition to a p53 antagonist, and explain the means by which E1A expression sensitizes cells to DNA damage-mediated apoptosis.

Results

Interaction of BAK with MCL-1 is lost in adenovirus-infected cells

BAK and BAX undergo stepwise conformational changes revealed by epitope exposure that are induced by E1A expression during infection with a proapoptotic E1B 19K mutant adenovirus. Specifically, an amino-terminal BAK epitope is exposed, whereas BAX exhibits exposure of amino- and carboxy-terminal epitopes. Conformational change of BAX but not BAK, and oligomerization of BAK and BAX are prevented by expression of E1B 19K in wild-type virus-infected cells, and interaction of E1B 19K was observed predominantly with BAK rather than BAX [Cuconati et al. 2002]. These observations suggest that E1A expression signals a conformational change in BAK and the formation of a proapoptotic complex between BAK and BAX, causing a conformational change in BAX that is prevented by E1B 19K–BAK interaction. The nature of the proapoptotic signal responsible for BAK activation by
E1A led us to search for BAK-interacting proteins in vivo. Because BAK and BAX are regulated through activation by BH3-only proteins, or are sequestered by anti-apoptotic BCL-2 family members, we screened BAK for interaction with both classes of apoptotic regulators. HeLa cells were either mock infected, infected with wild-type [Ad5dl309] adenovirus, or with an E1B 19K gene-deleted adenovirus [Ad5dl337], and cells were harvested 24 h post-infection. Immunoprecipitations were carried out using antibodies directed to a variety of known prosurvival (BCL-2, MCL-1, and BCL-XL), and proapoptotic BH3-only proteins (BIM, BID, and tBID), using two different antibodies directed against BAK. Multiple antibodies were utilized for immunoprecipitations, as recognition of BCL-2 family members varies with epitope availability that is greatly influenced by protein conformation and/or protein–protein interactions. Among the antibodies tested, an anti-MCL-1 antibody [MCL-1] robustly coimmunoprecipitated BAK with MCL-1 from mock-infected cells, but not from Ad5dl309 or Ad5dl337-infected cells, whereas a second anti-MCL-1 antibody [MCL-1 Ab-1] did so weakly [Fig. 1]. Conversely, an antibody directed against BAK [BAK-TM] robustly coimmunoprecipitated MCL-1 with BAK in mock-infected, but not Ad5dl309 or Ad5dl337 infected cells, whereas a second anti-BAK antibody BAK(Ab-1) did so weakly [Fig. 1]. A third anti-BAK antibody [BAK-N] also coimmunoprecipitated MCL-1 when either used directly for immunoprecipitation or when cross-linked to sepharose prior to immunoprecipitation [data not shown]. E1B 19K coimmunoprecipitated with BAK from Ad5dl309-infected cells [Fig. 1] as expected (Cuconati et al. 2002). These observations indicated that BAK interacted with MCL-1 in uninfected cells, which was supplanted by an interaction between E1B 19K and BAK upon infection. In Ad5dl337 infected cells, in which apoptosis was induced, BAK coimmunoprecipitated with BAX, and no BAK–BAX coprecipitation was observed in mock or Ad5dl309-infected cells [Fig. 1] as expected (Cuconati et al. 2002). No coimmunoprecipitation of BAK with BCL-2, BCL-XL, BIM, BID, or tBID was observed in uninfected or infected cells [data not shown]. These results suggested that MCL-1 may sequester BAK in an inactive form in healthy, uninfected cells, and that E1B 19K may functionally substitute for MCL-1 in cells infected with the wild-type virus.

**Down-regulation of MCL-1 protein levels in adenovirus-infected cells**

The apparent modest reduction in MCL-1 levels in infected compared with mock-infected cells at 24 h post-infection [Fig. 1] led us to examine MCL-1 expression during a time course of infection as a possible mechanism behind the loss of the MCL-1–BAK interaction. HeLa cells were mock infected, or infected with the Ad5dl309, Ad5dl337, the E1B deletion mutant Ad5E1B−, the E1A deletion mutant Ad5E1A−, and the E1A+E1B deletion mutant Pac3. Samples were collected at 0, 12, 24, 36, 48, and 72 h post-infection and subjected to Western blot analysis for MCL-1, BCL-2, BCL-XL, and actin. Levels of MCL-1 were found to decrease starting 12–24 h post-infection in Ad5dl309, Ad5dl337, and Ad5E1B− infected cells, culminating in the substantial loss of MCL-1 protein [Fig. 2A]. In contrast, levels of MCL-1 were unaffected in mock, Ad5E1A−, and Pac3-infected cells [Fig. 2A]. Furthermore, the loss of MCL-1 occurred in Ad5dl309-infected cells, in which apoptosis was inhibited, indicating that it was not merely a consequence of cell death. Levels of BCL-2, actin [Fig. 2A], and BCL-XL [data not shown] did not change under any conditions, suggesting that the loss of MCL-1 protein was specific and occurred through an E1A-dependent mechanism. These observations suggest that the abrogation of MCL-1–BAK interaction may be prompted by a loss of MCL-1 protein levels.

**Destabilization of MCL-1 protein in adenovirus-infected cells**

We investigated the cause for the loss of MCL-1 protein levels by assessing the stability of the MCL-1 protein during adenovirus infection of HeLa cells. Cells were either mock infected or infected with Ad5dl309 or Ad5dl337 for 24 h, and then treated with a combination of the proteasome inhibitor epoxomicin (EPO) and the
translation inhibitor cycloheximide (CHX) for 6 or 20 h, with CHX alone for 0, 2, 4, 6, or 20 h, or were left untreated (UT). Inhibition of new protein synthesis under CHX treatment reduced the levels of MCL-1 protein modestly in mock-infected cells, which was more pronounced in adenovirus-infected cells (Fig. 2B). Addition of EPO for either 6 or 20 h prevented much of the loss of MCL-1 protein in both infected and uninfected cells, indicating that MCL-1 protein turnover is regulated by ubiquitin-dependent proteasome degradation. In mock-infected cells, MCL-1 was relatively stable with an apparent half-life >4 h. In contrast, adenovirus infection dramatically increases the rate of MCL-1 protein turnover with an apparent half-life of <2 h, which is reversed by addition of a proteasome inhibitor. Levels of actin remained constant under all conditions. These observations indicated that adenovirus infection stimulates the specific turnover of MCL-1, suggesting a mechanism for initiation of cell death during adenovirus infection that is upstream of E1B 19K inhibition.

Reduction of intracellular levels of mcl-1 mRNA during adenovirus infection

We ascertained the effect of infection on mcl-1 mRNA levels through real-time PCR analysis of total RNA from either mock, Ad5dl309, Ad5dl337, or Ad5E1A–infected HeLa cells at 0, 14, and 24 h post-infection. In infected cells expressing E1A, loss of mcl-1 mRNA is progressive over the course of infection, exhibiting up to sixfold repression [Fig. 2C]. In contrast, in Ad5E1A–infected cells, loss of the mcl-1 message does not progress beyond 14 h post-infection, and levels do not decrease by more than threefold. Levels of gapdh mRNA also decreased during infection, but significantly less than mcl-1 mRNA [Fig. 2C]. The data suggest that there is an E1A-specific mechanism for the repression of mcl-1 expression during infection, which could lead to reduced MCL-1 protein levels, and disruption of the BAK–MCL-1 complex.

Restoration of MCL-1 expression rescued apoptosis induced by adenovirus infection

To determine whether MCL-1 is functionally capable of rescuing infected cells from adenovirus-induced apoptosis, we utilized transient transfection to overexpress MCL-1 in infected HeLa cells. Cells were transfected with plasmids encoding MCL-1L fused to a FLAG epitope (FLAG–MCL-1L), E1B 19K fused to a V5 epitope (E1B 19K–V5), and β-galactosidase fused to an Xpress epitope (β-gal–Xpress). All three sets of transfected cells were
mock, Ad5dl309, or Ad5dl337-infected at 24 h post-transfection. Cells were fixed 36 h post-infection on glass coverslips, stained for immunofluorescence with antibodies directed to the cognate epitope tag, and then stained with Hoescht’s dye for determination of chromatin condensation. The percentage of cells undergoing apoptosis (exhibiting condensed chromatin) in the transfected population was determined visually by counting the number of condensation-positive cells in 100–300 transfected cells per sample. As depicted in Figure 3, A and B, cells exhibiting condensed chromatin were largely seen in the Ad5dl337-infected sample. Of those that were positive for transfection, the β-gal-Xpress-transfected sample under Ad5dl337 infection was found to contain ~40% apoptotic cells, as compared with <2% and 1% apoptotic cells in the Ad5dl337-infected E1B 19K-V5 and FLAG-MCL-1 expressing cells, respectively [Fig. 3B]. As expected, Ad5dl309 infection did not cause measurable levels of apoptosis, due to expression of E1B 19K, and introduction of E1B 19K into cells infected with Ad5dl337 rescued the cells from apoptosis [Fig. 3B]. Interestingly, the introduction of MCL-1 into Ad5dl337-infected cells blocked apoptosis similarly to E1B 19K, clearly indicating that MCL-1 overexpression was able to complement E1B 19K deletion. These observations showed that MCL-1 was functionally capable of inhibiting adenovirus-induced apoptosis, and suggest that under normal conditions, endogenous MCL-1 must be lost in order for adenovirus-induced apoptosis to occur.

**MCL-1 down-regulation is not sufficient for BAK activation**

Adenovirus infection and E1A expression results in exposure of an amino-terminal epitope on BAK revealed by enhanced immunoprecipitation by BAK(AB-1) [Fig. 1; Cuconati et al. 2002]. To test whether MCL-1 down-regulation is sufficient for exposure of this BAK epitope indicative of conformational activation of BAK, we utilized RNA interference to down-regulate MCL-1. Despite significant reduction of MCL-1 by transfection with one or two siRNAs specific for mcl-1, no increase in BAK immunoprecipitation was observed with BAK(AB-1) or two other conformation-sensitive anti-BAK antibodies [Fig. 3C], nor was there any apparent affect on cell viability [data not shown]. BAK–MCL-1 immunoprecipitation was observed with the BAK–TM antibody as expected [Fig. 3C]. Thus, MCL-1 down-regulation by E1A may be required for apoptosis, but it is not sufficient for BAK activation and cell death.

**Induction of H2AX and ATM phosphorylation by E1A expression during adenovirus infection**

The E1A-dependent down-regulation of MCL-1 observed during adenovirus infection was strikingly similar to the mechanism of apoptosis induction in response to DNA damage. Ultraviolet or γ irradiation stimulate the elimination of MCL-1 protein expression, which is required,
but not sufficient, for apoptosis [Nijhawan et al. 2003]. Indication of induction of a DNA-damage response in adenovirus-infected cells was explored by testing for accumulation of the phosphorylated forms of ATM and H2AX. ATM responds to double-strand DNA breaks by undergoing autophosphorylation at serine 1981, and a downstream substrate in the ATM pathway, H2AX, becomes phosphorylated on serine 139. Antibodies specific for the phosphorylated forms of ATM (ATM1981S-P) and H2AX (γH2AX) were utilized in Western blots, IP-Westerns, and indirect immunofluorescence to determine whether their levels were induced by adenovirus infection. Whereas levels of γH2AX were low as expected in uninfected cells, the levels accumulated dramatically in cells infected with the wild-type virus [Ad5dl309; Fig. 4A]. γH2AX levels also accumulated in cells infected with the two E1B mutant viruses [Ad5dl337 and Ad5E1B−], however, this was expected, as these viruses induce apoptosis, the DNA fragmentation of which creates double-strand DNA breaks. Interestingly, infection with the two E1A deletion mutants (E1A− and Pac3) did not induce γH2AX accumulation [Fig. 4A], indicating the requirement for E1A expression. E1A and E1B 19K were expressed only by the viruses that encoded their genes, and cellular actin levels were comparable among the samples as an indication of equal protein leading [Fig. 4A].

To test for ATM activation by phosphorylation upon infection, ATM was immunoprecipitated from mock, Ad5dl309, or Ad5dl337-infected cells and subjected to Western blotting using either ATM or and ATM phospho-serine 1981-specific antibodies. Only Ad5dl309 and Ad5dl337, but not mock-infected cells, displayed evidence of ATM phosphorylation on serine 1981 [Fig. 4B]. Double-label indirect immunofluorescence for γH2AX, and the adenovirus 72 kD DBP as a marker for infected cells, revealed similar pronounced induction of serine 139 phosphorylated H2AX [Fig. 4C]. Indirect immunofluorescence with the phospho-serine 1981 ATM antibody also demonstrated its induction upon infection with both the wild-type Ad5dl309 and proapoptotic Ad5dl337 adenoviruses [Fig. 4C]. This suggested that cells responded to E1A expression during virus infection in a manner that resembled the cellular response to DNA damage, not only with respect to p53 induction, but also evidenced by accumulation of phosphorylated forms of H2AX and ATM, which may be linked to the loss of MCL-1, which facilitates apoptosis.

Discussion

MCL-1 regulates adenovirus-induced apoptosis

The mitochondrial checkpoint, regulated by proapoptotic BCL-2 family members BAK and BAX, plays an important role in the regulation of death signaling prompted by E1A expression in adenovirus-infected cells. Specifically, E1A expression in E1B 19K mutant virus-infected cells induces conformational changes in BAK and BAX, and the formation of a high-molecular weight BAK–BAX protein complex, which coincides with the release of cytochrome c from the mitochondria, caspase-9 and caspase-3 activation and cell death. In the presence of E1B 19K in cells infected with the wild-type virus, BAK displays a change in conformation, which is associated with interaction of BAK with the E1B 19K protein, but conformational changes in BAX and the formation of the BAK–BAX complex are abrogated. This interruption of apoptosis signaling through mitochondria is manifested by failure to release cytochrome c and activate caspase-9 and caspase-3, which extends the viability of infected cells.

Proof of the requirement for BAK and BAX in E1A-mediated apoptosis of adenovirus-infected cells was re-
revealed when BAK and BAX deficiency prevented apoptosis induction upon infection with and E1B 19K mutant adenovirus, and greatly enhanced virus production (Cuconi et al. 2002). Furthermore, this validated BAK and BAX as the functional targets for E1B 19K in inhibition of apoptosis during productive adenovirus infection. Thus, BAK and BAX function as part of an antiviral response to limit virus replication through apoptosis induction, which is thwarted by E1B 19K–BAK and E1B 19–BAX interaction to sustain the viability of infected cells until the replication cycle is complete. We expected that identification of the signal by which infection and E1A expression activates BAK upstream of E1B 19K should provide insight into the virus–host cell interactions that initiate the apoptotic defense cascade. BAK is normally sequestered by antiapoptotic MCL-1, and what is likely a DNA damage response to virus infection represses MCL-1 expression, priming infected cells to apoptosis induction.

BAK has been reported recently to interact with the mitochondrial outer-membrane protein VDAC2, which acts to restrict BAK activation (Cheng et al. 2003). There may be multiple BAK protein complexes that respond to different death stimuli to regulate BAK activation and apoptosis. In addition to MCL-1 and VDAC2, BAK also interacts with E1B 19K and tBID, and BAK protein interactions may be required for both BAK activation and inhibition.

In healthy, uninfected cells, 24-kD BAK migrates with a native molecular weight of at least 60–70 kD by gel filtration chromatography, suggesting that it may be complexed with other cellular proteins (Sundararajan et al. 2001). In contrast, BAX appears predominantly monomeric, fractionating with an expected native molecular weight of ~21 kD [Sundararajan and White 2001; Sundararajan et al. 2001]. The interaction of BAK with 42-kD MCL-1 reported here may explain the aberrant native molecular weight of BAK, and suggests that MCL-1 may be a BAK chaperone to regulate the proapoptotic activity of BAK. MCL-1 fractionates by gel-filtration chromatography with a native molecular weight of at least 60 kD with significant overlap with the BAK fractionation profile consistent with BAK–MCL-1 complex formation as well as complex formation with other proteins (R. Sundararajan and E. White, unpubl.). MCL-1 has potent antiapoptotic function (Kozopas et al. 1993; Reynolds et al. 1994; Craig 2002), and has been reported to interact with BAK in the yeast two-hybrid system [Bae et al. 2000]. As shown here, MCL-1 may function in mammalian cells by interacting with and inhibiting BAK, and loss of MCL-1 upon E1A expression in adenovirus-infected cells may enable the apoptotic response [Fig. 5A]. Restoration of MCL-1 is sufficient to block apoptosis induced by E1A expression in adenovirus-infected cells, and the E1B 19K protein apparently functionally substitutes for the absence of MCL-1 by interacting with and inhibiting BAK once the MCL-1–BAK complex is disrupted [Fig. 5B]. In that way, E1B 19K can be thought of as a functional homolog of MCL-1 more so that BCL-2, which may predominantly function through interaction with and inhibition of BH3-only proteins rather than BAK and BAX (Cheng et al. 2001).

**E1A provokes a DNA damage response that results in MCL-1 elimination**

The elimination of MCL-1 by E1A expression during adenovirus infection occurred by both increased ubiquitin-dependent turnover of the protein and repression of expression of the mRNA. The expression of MCL-1 and its prosurvival function is regulated by growth factors and is essential for early mammalian development (Craig 2002). Proapoptotic stimuli are also known to repress MCL-1 expression. E2F-1 activation, for example, represses mcl-1 expression, and E1A activates E2F-1 through binding and inhibition of RB (Croxton et al. 2002a,b). DNA damage, induced by either ultraviolet or γ irradiation, down-regulates MCL-1 protein levels in all, or in part, by stimulating proteasome-mediated turnover of MCL-1 protein [Nijhawan et al. 2003]. Although E1A-mediated, E2F-1 activation could account for down-regulation of the mcl-1 mRNA, the enhanced turnover of the

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**Figure 5.** Schematic representation of the regulation of adenovirus-induced apoptosis in infected cells. See text for explanation.
MCL-1 regulates adenovirus-induced apoptosis

MC1-1 protein in adenovirus-infected cells resembled a DNA-damage response. Remarkably, H2AX, a known substrate of the PI3 kinases ATM and ATR that initiate the cellular response to DNA damage [Rogakou et al. 1998, 2000; Ward and Chen 2001; Redon et al. 2002; Bakkenist and Kastan 2003], and ATM itself, become dramatically phosphorylated upon adenovirus infection [Fig. 4]. The induction of γH2AX (the phosphorylated form of H2AX) and ATM 1981S-P occurred in both wild-type virus-infected cells, in which apoptosis is blocked by E1B 19K expression, and in cells infected with a pro-apoptotic E1B 19K mutant virus. Thus, induction of γH2AX and ATM 1981S-P upon infection occurred upstream of apoptosis induction.

E1A expression also resembles a DNA-damage response by inducing p53 accumulation [Lowe and Ruley 1993], which is seen during productive adenovirus infection as long as the E1B 55K gene is deleted [Chiou and White 1997]. In infected HeLa cells, p53 induction by E1A maps primarily to the p300 and not the RB-binding region [Chiou and White 1997], and E1A induces the polyubiquitination of p53 by p300, which promotes, along with MDM-2, p53 degradation in proteasomes [Grossman et al. 2003]. It is not known whether regulation of p53 turnover by p300 is responsible p53 accumulation in response to DNA damage. Likely, a combination of events initiated by E1A expression signals a cellular stress response that primes cells for apoptosis. As we can only correlate activation of a DNA-damage response by E1A to initiation of apoptosis, it will be of great interest to interfere with upstream components of the DNA-damage signaling [ATM and ATR, for example] to determine whether they are necessary or sufficient for apoptosis induced by E1A.

Although adenovirus encodes within E1B 19K and E1B 55K mechanisms to disable this cellular response to stress, the upstream pathways to activate p53, ATM, and H2AX in response to E1A expression appear intact. How adenovirus infection and E1A expression induces a DNA-damage response is not clear, but could be explained by the induction of double-strand DNA breaks resulting from either replication stress mediated by deregulation of cell cycle control by E1A, or by viral DNA-replication intermediates that may be perceived as damaged DNA, or by changes in chromatin structure that are known to activate ATM [Bakkenist and Kastan 2003]. When expressed in transformed cells, the human papilloma virus E7 gene product, which functions analogously to E1A by antagonizing RB and deregulating cell cycle control, also induces γH2AX that is associated with genomic instability [Duensing and Munger 2002]. H2AX is required to prevent genomic instability and tumorigenesis in vivo [Bassing et al. 2003; Celeste et al. 2003]. This raises the possibility that transformation by viral oncogenes may initiate a DNA-damage response triggering down-regulation of MCL-1 and susceptibility to apoptosis. E1A expression sensitizes cells to apoptosis induced by DNA-damaging agents [Sanchez-Prieto et al. 1996; Brader et al. 1997; Shao et al. 1997; Stiewe et al. 2000; Viniegra et al. 2002], as does specific knock-down of MCL-1 [Nijhawan et al. 2003]. This priming of apoptosis by E1A may require adenovirus to encode an antiapoptotic function to sustain viability during oncogenesis.

Finally, although elimination of MCL-1 is required for apoptosis in response to DNA damage, it is not sufficient [Nijhawan et al. 2003]. Eliminating MCL-1 may render BAK susceptible to activation by BH3-only proteins that are coordinately induced in infected cells. Interestingly, the antiapoptotic latent membrane protein-1 of Epstein-Barr virus induces MCL-1 accumulation [Wang et al. 1996]. Taken together, MCL-1 may be part of an antiviral response to render infected cells susceptible to apoptosis. This may have necessitated the evolution of viral countermeasures to either prevent the destruction of MCL-1, or for the viruses themselves to encode functional substitutes of MCL-1 to maintain viability of infected cells until the replication cycle is complete. It will be of great interest to determine whether other classes of viruses possess similar activities.

Materials and methods

Virus infection and Western blotting

HeLa cells were mock infected, or infected with adenovirus type 5 (Ad5) wild-type Ad5d1309 [Jones and Shenk 1979], the E1B 19K gene deletion mutant Ad5d1337 [Pilder et al. 184], the E1B gene-deletion mutant Ad5E1B, the E1A gene deletion mutant Ad5E1A, or the E1A and E1B gene-deletion mutant Pac3 [Chiou et al. 1994], at a multiplicity of infection [MOI] of 100 by standard methods [White et al. 1984]. Attached and floating cells were harvested at 0 (for mock only), 12, 24, 36, 48, and 72 h post-infection by scraping and centrifugation. All cell pellets were resuspended in 2× Laemmli buffer and vortexed briefly. Each lysate sample was subjected to 17% SDS-PAGE, and analyzed by Western blotting [Perez and White 1998] with the following primary antibodies: anti-MCL-1 rabbit polyclonal [Stressgen Biotechnologies]; anti-BCL-2 hamster monoclonal polyclonal [PharMingen]; anti-E1A and anti-actin mouse monoclonal [Oncogene Research]; anti-E1B 19K rabbit polyclonal [Sundararajan and White 2001]; and anti-γH2AX (anti-H2AX phosho Ser139) rabbit polyclonal [Upstate Biotechnology] antibodies. Blots were developed with horseradish peroxidase-conjugated secondary antibodies using the ECL system [Amer sham-Pharmacia Biotech].

Immunoprecipitation

For each immunoprecipitation reaction, HeLa cells were infected as indicated and harvested at 24 and 48 h post-infection by scraping and lysed in 2.0% 3-[3-cholamidopropyl]-dimethylammonio]-1-propanesulfonate [CHAPS]-containing lysis buffer and immunoprecipitated as described previously [Perez and White 2000]. Immunoprecipitations were carried out with the following primary antibodies: anti-p19 rabbit polyclonal and anti-PCNA mouse monoclonal [Santa Cruz Biotechnology]; anti-BAK (Ab-1) mouse monoclonal [Oncogene Research]; anti-BAK (minus transmembrane domain, or ~TM) rabbit polyclonal [PharMingen]; anti-BAK-NT rabbit polyclonal [Upstate Biotechnology]; anti-MCL-1 rabbit polyclonal [Stressgen]; and anti-MCL-1 mouse monoclonal [Oncogene Research]; anti-MYC rabbit polyclonal [Santa Cruz Biotechnology]; anti-BAX (1-113) rabbit polyclonal [Santa Cruz Biotechnology]; anti-BAX (150-165) rabbit polyclonal [Santa Cruz Biotechnology].
rabit polyclonal (Oncogene Research), anti-BAX(43-61) rabbit polyclonal (PharMingen), and anti-BAK G-23 rabbit polyclonal (Santa Cruz Biotechnology). Immunoprecipitates were resuspended in 2x Laemmli buffer, and a sample of each immunoprecipitation reaction was subjected to 17% SDS-PAGE. Western blotting of immunoprecipitates was carried out with the following primary antibodies: anti-MCL-1 rabbit polyclonal (Stressgen), anti-BAX[11-30] rabbit polyclonal (Santa Cruz), anti-BAK[23-37] (Upstate Biotechnology); and anti-E1B 19K rabbit polyclonal (Sundarajan and White 2001) antibodies. The secondary antibody used was horseradish peroxidase-conjugated goat anti-rabbit (Southern Biotech). For immunoprecipitation of ATM, HeLa cells were Mock, Ad5dl309, or Ad5dl337 infected as described above, and cells were harvested and lysed in 50 mM Tris (pH 7.5), 150 mM NaCl, 1% Tween-20, 0.05% NP-40, 50 mM NaF, with protease inhibitors. Immunoprecipitations were carried out with an anti-ATM rabbit polyclonal antibody (Rockland Immunocemical Research) and subjected to 7.5% SDS-PAGE, and Western blotted with either anti-ATM or anti-ATM1981S-P rabbit polyclonal antibodies (Rockland Immunocemical Research, Bakkenist and Kastan 2003).

Determination of MCL-1 protein stability
HeLa cells were infected as indicated at an MOI of 100 and treated with 50 µg/mL of cycloheximide (CHX, Sigma) solubilized in ethanol starting at 24 h post-infection for durations of 0, 2, 4, 6, or 20 h, or CHX and 200 nM of epoxomicin (EPO, Alexis) solubilized in dimethyl sulfoxide for durations of 6 or 20 h, or mock treated with dimethyl sulfoxide and ethanol alone. Cells were harvested at the indicated time point by scraping and centrifugation, and were resuspended in 2x Laemmli buffer and vortexed briefly. Lysate samples were subjected to 17% SDS-PAGE, and analyzed by Western blotting with the following primary antibodies: anti-MCL-1 rabbit polyclonal (Stressgen Biotechnoloe) and anti-actin mouse monoclonal (Oncogene Research). 

Analysis by real-time PCR
A total of 2 x 10^6 HeLa cells per condition were infected as indicated at an MOI of 100, and were harvested by scraping and centrifugation at 0, 14, and 24 h post-infection. Total RNA was isolated through the use of the RNeasy Mini kit (QIAGEN), and 100 ng of total RNA from each condition was subjected to real-time RT–PCR with the Taqman EZ-RT PCR kit (PE Applied Biosystmes) using recommended conditions, in the ABI Prism 7700 Sequence Detector. Optimal primer and probe sequences for mcl-1 real-time PCR were determined with the Primer Express software version 1.5 (Applied Biosystems), and were supplied compliments of Integrated DNA Technologies; probe sequence was supplied with 5’ linkage to the reporter dye 6-carboxy-fluorescein [FAM], and 3’ linkage to Black Hole quencher (Integrated DNA technologies). MCL-1 primer sequences were 5’TGAAATCTGTTGCTCTGAAGCTATG-3’ and 5’TTCACAA TCGCCCACCTT-3’. MCL-1 probe sequence was 5’FAM-5’TCCATGTTTTCAGCGACGGCGTAA-3’. 

Indirect immunofluorescence
HeLa cells transfected by electroporation with plasmid pCDNA3-FLAG-MCL-1L (Bae et al. 2000), kindly provided by J. Bae and A.J. Hsueh, Stanford University, plasmid pCDNA3.1-E1B-V5 (Kasof et al. 1998), or plasmid pCDNA3.1-ß-Gal-Xpress [Invitrogen] were grown on glass coverslips and were mock infected with Ad5dl309, Ad5dl337, or were mock infected for 30 h at an MOI of 100. Cells on coverslips were fixed in 4% paraformaldehyde, and indirect immunofluorescence was performed as described previously (Perez and White 1998), except that coverslips were blocked with 4% BSA/PBS for 1 h at 37°C. Coverslips were washed with anti-FLAG mouse monoclonal [Sigmal, V-5 mouse monoclonal [Invitrogen], anti-Xpress mouse monoclonal [Invitrogen], anti-yH2AX rabbit polyclonal [Upstate Biotechnology], anti-phosphorylated ATM [1981S-P] polyclonal [Bakkenist and Kastan 2003, Rockland Immunocemical Research], or anti-tiadenovirus 72-kd DNA-binding protein [DBP, generously provided by Dr. Peter Yacuik, St. Louis University] antibodies. In some cases, following secondary antibody staining, coverslips were washed with PBS and stained with Hoechst’s dye. Staining was visualized by epifluorescence microscopy as described previously (Perez and White 1998), and percentages of cells positive for both transgene expression and or adenovirus structural proteins were determined by scoring 100–200 cells on each coverslip.

DNA interference
A total of 200 nM of annealed, purified, and desalted double-stranded siRNA targeted against mcl-1, MCL-1 [1] [Nijhawan et al. 2003], and MCL-1 [2] (AAGGACACACAAAGCCAATGG GC; Dharmacon Research) were mixed with 175 µL of OPTI-MEM (Invitrogen) and incubated for 10 min. A mixture of 3 µL of Oligofectamine Reagent [Invitrogen] and 12 µL of OPTI-MEM was then added to the above and incubated for 20 min. Medium was removed from adherent HeLa cells, which were plated in DMEM without antibiotics and with 10% FBS in a 6-well dish on day 0 to achieve about 30%–50% confluence on day 1. A total of 900 µL of antibiotic-free medium with 10% FBS was added per well and overlaid with 200 µL of the siRNA mixture. Cells were incubated at 37°C for 4 h, and 500 µL of DMEM with 30% serum was then added to each well. Immunoprecipitations using Bca2 and BAK antibodies was performed in CHAPS buffer as described above at 24 h post-transfection. The BAK antibodies used were the BAK carboxy-terminal polyclonal antibody [PharMingen], and conformation-specific BAK antibodies BAK/Ab-1 [Oncogene] and BAK G-23 (Santa Cruz Biotechnology). Western blots were performed with BAK polyclonal antibody [Upstate Biotechnology], MCL-1 polyclonal antibody [Stressgen].

Acknowledgments
We thank Dr. Deirdre Nelson for critical reading of the manuscript, Guanghua Chen for technical assistance, Drs. Jechyeon Bae and Aaron J. Hsueh (Stanford University) for the plasmid pCDNA3-FLAG-MCL-1L, and Dr. Peter Yacuik (St. Louis University) for the anti-adenovirus DBP monoclonal antibody. This work was funded by NCI grant R37-CA53370 and the Howard Hughes Medical Institute.
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