Distinct Requirements of Adenovirus E1b55K Protein for Degradation of Cellular Substrates


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Received 4 May 2008/Accepted 3 July 2008

The E1b55K and E4orf6 proteins of adenovirus type 5 (Ad5) assemble into a complex together with cellular proteins including cullin 5, elongins B and C, and Rbx1. This complex possesses E3 ubiquitin ligase activity and targets cellular proteins for proteasome-mediated degradation. The ligase activity has been suggested to be responsible for all functions of E1b55K/E4orf6, including promoting efficient viral DNA replication, preventing a cellular DNA damage response, and stimulating late viral mRNA nuclear export and late protein synthesis. The known cellular substrates for degradation by E1b55K/E4orf6 are the Mre11/Rad50/Nbs1 DNA repair complex, the tumor suppressor p53, and DNA ligase IV. Here we show that the degradation of individual targets can occur independently of other substrates. Furthermore, we identify separation-of-function mutant forms of E1b55K that can distinguish substrates for binding and degradation. Our results identify distinct regions of E1b55K that are involved in substrate recognition but also imply that there are additional requirements beyond protein association. These mutant proteins will facilitate the determination of the relevance of specific substrates to the functions of E1b55K in promoting infection and inactivating host defenses.

The linear, double-stranded DNA genome of adenovirus serotype 5 (Ad5) is approximately 36 kb and encodes five early transcription units whose proteins perform essential functions for efficient infection. Viruses such as adenovirus employ numerous strategies to contend with host cell factors during the course of establishing a productive infection (10, 68). Examining mutant viruses has implicated E1b and E4 proteins in the modulation of the host cell environment, the promotion of viral DNA replication and viral mRNA export, the prevention of viral genome concatemerization, and the synthesis of late viral proteins (reviewed in references 3, 61, 67, and 68). In adenovirus-infected cells, most of the viral E1b55K protein is present in a complex with E4orf6 (52, 53). The E1b55K/E4orf6 complex has been implicated in the selective modulation of nucleocytoplasmic mRNA during the late phase of viral infection (reviewed in references 18 and 23). This viral complex also promotes efficient viral replication (8, 29, 33, 66) and prevents the viral genome from being concatemerized by host factors (60, 65). The precise mechanisms by which the E1b55K/E4orf6 complex achieves its many functions remain to be elucidated.

E1b55K displays a complex distribution pattern in infected cells and requires E4orf6 for nuclear localization (17, 27, 37, 42, 47) and association with viral replication centers (48). The E1b55K protein possesses a leucine-rich nuclear export signal (NES), which has been implicated in nucleocytoplasmic shuffling via the CRM1-mediated nuclear export receptor (19, 39).

Export-deficient mutant forms of E1b55K accumulate in subnuclear aggregates that also contain cellular binding partners (20, 31). In the absence of E4orf6, the expression of E1b55K produces cytoplasmic aggregates (17, 27, 37, 42, 47). In transformed cells, E1b55K from Ad2 or Ad5 accumulates in a large cytoplasmic body (4, 9, 20, 74) with characteristics of an aggresome (38, 42). Many cellular proteins that interact with E1b55K localize at the aggresome in transfected and transformed cells (4, 9, 20, 22, 42, 44, 74). Therefore, protein aggregation has been proposed previously as a potential strategy utilized by E1b55K to inactivate cellular proteins (42) and promote transformation (31).

One tactic commonly employed by viruses to inactivate inhibitory host factors is to induce the specific downregulation or degradation of cellular proteins (25, 55). Degradation can be achieved via the covalent modification of target proteins with polyubiquitin chains by an E3 ubiquitin ligase, followed by recognition and destruction by the proteasome (reviewed in reference 35). The adenoviral E1b55K and E4orf6 proteins associate with cellular proteins to form an E3 ligase complex that contains elongins B and C, cullin 5, and Rbx1 (7, 30, 49). BC-box motifs in E4orf6 have been identified previously as important for binding elongins B and C (5, 15, 43). It has been suggested that all functions of the E1b55K/E4orf6 complex are due to the degradation of cellular proteins by the ubiquitin ligase activity (6, 16, 70). Since E1b55K alone can physically associate with p53 (13, 34, 41, 56, 71, 73), DNA ligase IV (2), and the Mre11/Rad50/Nbs1 (MRN) complex (12), it is believed to mediate substrate recognition, while both E4orf6 and E1b55K are required for proteasome-mediated degradation (12, 13, 49, 50, 56, 69). Not all proteins that interact with E1b55K are downregulated, but cellular proteins so far identified as degradation substrates of the E1b55K/E4orf6 complex include p53 (13, 28, 30, 46, 49–51, 58), the MRN DNA repair...
The degradation of these cellular factors has been demonstrated by inhibition with proteasome inhibitors and by increased turnover in studies of protein half-life (12, 46, 49–51, 58, 60, 69). Exactly which substrates are relevant to specific functions of the E1b55K/E4orf6 complex is unclear. Additionally, the mechanisms for the selection and degradation of distinct proteins remain to be resolved. Understanding how the E1b55K/E4orf6 complex targets specific cellular substrates crucial to elucidating how these viral proteins perform their myriad of functions.

In the present study, we characterized some of the cellular and viral requirements for the degradation of each target. By examining the steady-state levels of E1b55K/E4orf6 substrates in mutant cell lines, we demonstrated that the downregulation of individual cellular factors can occur independently of the other factors. A series of E1b55K mutant proteins were characterized with respect to localization and the degradation of the known substrates. We identified regions of E1b55K important for cellular localization, and we demonstrated that there are independent requirements for p53, MRN, and DNA ligase IV downregulation. Our results show that affinity for substrates are independent requirements for p53, MRN, and DNA ligase IV, and we demonstrated that there is crucial to elucidating how these viral proteins perform their myriad of functions. The degradation of these cellular factors has been demonstrated by inhibition with proteasome inhibitors and by increased turnover in studies of protein half-life (12, 46, 49–51, 58, 60, 69). Exactly which substrates are relevant to specific functions of the E1b55K/E4orf6 complex is unclear. Additionally, the mechanisms for the selection and degradation of distinct proteins remain to be resolved. Understanding how the E1b55K/E4orf6 viral ubiquitin ligase targets specific cellular substrates is crucial to elucidating how these viral proteins perform their myriad of functions.

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**MATERIALS AND METHODS**

**E1b55K mutagenesis and cloning and cell transfections.** Previously described E1b55K mutant genes were amplified from corresponding viruses (56, 72) with PCR primers RFV (forward primer; 5'-CCGCTCGAGATGGAGCGAAGAAACCACAT-3') and RVR (reverse primer; 5'-CCATATGCTCGAGATGGAGCGAAGAAACCACAT-3') and cloned into the Xho1 and Cla1 sites of pDC516 (see the legend to Fig. 7B), genes encoding the E1b555K mutant proteins R240A and H373A were PCR amplified and introduced in place of the wild-type (WT) E1b55K gene in either the pDC516 backbone vector or pCLNC retroviral vector, as previously described (12). Site-directed mutagenesis of the E1b55K gene in either the pDC516 backbone vector (provided by H. Young) or pCLNC was performed using a QuikChange mutagenesis kit (Stratagene) and the primers listed in Table 1. Mutant proteins were screened as described below, and interesting mutant constructs in the pDC516 vector were PCR amplified using RFV and RVR primers (or a modified RVR primer) from pDC516-T3 and cloned into the Xho1 and Cla1 sites of plasmid pCLNC (54). To generate NES mutant constructs in pDC516 (see the legend to Fig. 7B), genes encoding the E1b555K mutant proteins R240A and H373A were PCR amplified and introduced in place of the wild-type (WT) E1b55K gene in either the pDC516 backbone vector (provided by H. Young) or pCLNC was performed using a QuikChange mutagenesis kit (Stratagene) and the primers listed in Table 1. Mutant proteins were screened as described below, and interesting mutant constructs in the pDC516 vector were PCR amplified using RFV and RVR primers (or a modified RVR primer) from pDC516-T3 and cloned into the Xho1 and Cla1 sites of plasmid pCLNC (54). To generate NES mutant constructs in pDC516 (see the legend to Fig. 7B), genes encoding the E1b555K mutant proteins R240A and H373A were PCR amplified and introduced in place of the wild-type (WT) E1b55K gene in either the pDC516 backbone vector (provided by H. Young) or pCLNC was performed using a QuikChange mutagenesis kit (Stratagene) and the primers listed in Table 1. Mutant proteins were screened as described below, and interesting mutant constructs in the pDC516 vector were PCR amplified using RFV and RVR primers (or a modified RVR primer) from pDC516-T3 and cloned into the Xho1 and Cla1 sites of plasmid pCLNC (54). To generate NES mutant constructs in pDC516 (see the legend to Fig. 7B), genes encoding the E1b555K mutant proteins R240A and H373A were PCR amplified and introduced in place of the wild-type (WT) E1b55K gene in either the pDC516 backbone vector (provided by H. Young) or pCLNC was performed using a QuikChange mutagenesis kit (Stratagene) and the primers listed in Table 1. Mutant proteins were screened as described below, and interesting mutant constructs in the pDC516 vector were PCR amplified using RFV and RVR primers (or a modified RVR primer) from pDC516-T3 and cloned into the Xho1 and Cla1 sites of plasmid pCLNC (54).

### Table 1. E1b55K mutagenesis primers

<table>
<thead>
<tr>
<th>Mutant protein</th>
<th>Vector backbone</th>
<th>Round of mutagenesis</th>
<th>Primer type</th>
<th>Primer sequence (5' to 3')</th>
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<tr>
<td>L83,87,91A (NES mutant protein)</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Reverse</td>
<td>CAGCTGTAAGCGGTAGGAGAGAGAGAGAG</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Second round</td>
<td>Forward</td>
<td>GCCGTGACGGGTCCCTCCCTCTCTCTCGG</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Reverse</td>
<td>GCTGTCGACGGGTCCCTCCCTCTCTCTCGG</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Third round</td>
<td>Forward</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Reverse</td>
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<td></td>
<td></td>
<td></td>
<td>Reverse</td>
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</tr>
<tr>
<td>C348,351S</td>
<td>pDC516</td>
<td></td>
<td>Forward</td>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Reverse</td>
<td>CCCCCTGCGGCTCTGTCTTTAG</td>
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<tr>
<td>C361,366S</td>
<td>pDC516</td>
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<tr>
<td></td>
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<td></td>
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<td>Second round</td>
<td>Forward</td>
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<tr>
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<td>pDC516</td>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Reverse</td>
<td>CCCCCTGCGGCTCTGTCTTTAG</td>
</tr>
<tr>
<td>S490,491,T495A</td>
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<tr>
<td></td>
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<td>Reverse</td>
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<td></td>
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<td>Second round</td>
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<tr>
<td></td>
<td></td>
<td>Second round</td>
<td>Forward</td>
<td>GCCGTGACGGGTCCCTCCCTCTCTCTCGG</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Reverse</td>
<td>GCTGTCGACGGGTCCCTCCCTCTCTCTCGG</td>
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</table>

* Codons changed in the primer sequence are underlined. Primers were used to introduce changes into E1b55K with the backbone vector pDC516 or pCLNC. Constructs encoding interesting mutant proteins were subcloned into retroviral vector pCLNC in order to make cell lines. All mutations were verified by sequencing.
U2OS that express WT and mutant E1b55K from retrovirus vectors have been described previously (12). Immortalized A-TLD3 and A-TLD1 cells were described previously (12, 59, 60). NBS cells were provided by P. Concannon (14). FUS9 cells with mutant DNA-dependent protein kinase (DNA-PKcs) (32) were provided by T. Melendez. All cells were maintained as monolayers in Dulbecco modified Eagle’s medium (DMEM) supplemented with 10 or 20% fetal bovine serum (FBS) and penicillin-streptomycin. IMR90, A-TLD3, A-TLD1, or NBS cells were either mock treated (M) or infected with WT Ad5 (Ad; MOI of 25), and lysates were harvested at the indicated times postinfection for analysis by immunoblotting with specific antibodies. GAPDH served as a loading control. Protein levels were compared to those in mock-infected cells (M). (B) Independent degradation of p53, the MRN complex, and DNA ligase IV. Cell lines with mutant p53 (Saos2), Mre11 (A-TLD1), or DNA-PKcs (FUS9) were infected with WT Ad5 (MOIs of 25, 100, and 25, respectively). Lysates were harvested at 24 to 48 hpi and analyzed by immunoblotting for the indicated proteins. +, present; −, absent. (C) Mre11 may be the degradation target within the MRN complex. IMR90, A-TLD3, A-TLD1, or NBS cells were either mock treated (M) or infected with WT Ad5 (Ad; MOI of 75). Cells were harvested at 24 to 48 hpi and were analyzed by immunoblotting for the indicated proteins. Ku70 served as a loading control.

FIG. 1. Cellular substrates of the E1b55K/E4orf6 complex are degraded independently of one another during adenovirus infection. (A) Degradation of cellular proteins over a time course of Ad5 infection. U2OS cells were infected with WT Ad5 (MOI of 25), and lysates were harvested at the indicated times postinfection for analysis by immunoblotting with specific antibodies. GAPDH served as a loading control. Protein levels were compared to those in mock-infected cells (M). (B) Independent degradation of p53, the MRN complex, and DNA ligase IV. Cell lines with mutant p53 (Saos2), Mre11 (A-TLD1), or DNA-PKcs (FUS9) were infected with WT Ad5 (MOIs of 25, 100, and 25, respectively). Lysates were prepared at 24 to 48 hpi and analyzed by immunoblotting for the indicated proteins. +, present; −, absent. (C) Mre11 may be the degradation target within the MRN complex. IMR90, A-TLD3, A-TLD1, or NBS cells were either mock treated (M) or infected with WT Ad5 (Ad; MOI of 75). Cells were harvested at 24 to 48 hpi and were analyzed by immunoblotting for the indicated proteins. Ku70 served as a loading control.

RESULTS

Cellular substrates of the E1b55K/E4orf6 complex are degraded independently of one another. We first analyzed the kinetics of substrate degradation in U2OS cells, which express all currently known targets of the E1b55K/E4orf6 complex. The degradation of cellular proteins was examined by immunoblotting over a time course of infection with WT Ad5 (Fig. 1A). During infection, the E1b55K protein was first detected at 12 hpi. At this time point, the total steady-state levels of the Mre11, Rad50, Nbs1, p53, and DNA ligase IV proteins began to decrease. Nbs1 downregulation was slightly delayed compared to that of the other substrates, suggesting that Nbs1 may...
not be a direct target. Since many of the cellular DNA repair proteins are part of large protein complexes, we determined whether cellular proteins could be degraded independently of one another by using mutant cell lines (Fig. 1B). Mre11, Rad50, and DNA ligase IV were degraded in Saos2 cells that do not express p53. A-TLD1 cells (59) harbor a mutation in Mre11 that destabilizes the MRN complex. We found that DNA ligase IV and p53 could still be degraded in these cells, although we observed in experiments with reproducible results that p53 levels were not completely diminished. DNA ligase IV and the MRN complex are critical to DNA repair by homologous end joining. We therefore also examined whether DNA-PKcs, another key nonhomologous end-joining factor, was required for the degradation of substrates. Ad5 infection of FUS9 cells, which lack DNA-PKcs (32), still induced the downregulation of all known substrates. We have also previously shown that the ATM and ATR kinases and signaling pathways are not required for the degradation of MRN (12). Together, these data demonstrate that MRN, p53, and DNA ligase IV can undergo virus-induced downregulation independently of one another and other proteins involved in the DNA damage response.

To examine the degradation of the MRN complex by E1b55K/E4orf6, we employed mutant NBS and A-TLD cell lines. IMR90 fibroblast cells were infected as a control and showed the degradation of all MRN complex members (Fig. 1C). NBS cells harbor a mutation in the Nbs1 gene resulting in a truncated protein defective in complex formation with Mre11 and, thus, Rad50 (11). The infection of these cells with Ad5 caused the degradation of both Mre11 and Rad50 (Fig. 1C), suggesting that full-length Nbs1 is not required. Smaller proteins recognized by the Nbs1 antibody were not altered by Ad5 infection. A-TLD3 cells contain a missense mutation in Mre11 that diminishes its interaction with Nbs1 (59), while A-TLD1 cells express a prematurely truncated Mre11 protein that is unstable and defective for Rad50 and Nbs1 interaction (59). A-TLD3 cells showed reduced levels of both Mre11 and Rad50 during Ad5 infection (Fig. 1C). Although we had difficulty visualizing the truncated form of Mre11 in A-TLD1 cells, Rad50 was unaltered (Fig. 1C), suggesting that full-length Mre11 is required for the destabilization of Rad50 and Nbs1 during Ad5 infection.

Identification of additional E1b55K mutant proteins with defects in MRN degradation. Previous studies have described a large number of E1b55K mutant proteins (26, 52, 56, 63, 72), but many of these have not been assessed for the degradation of all the known substrates. We previously identified two E1b55K mutant forms that can distinguish between the degradation targets p53 and MRN (12). In the presence of E4orf6, the R240A mutant protein degrades the MRN complex but not p53, whereas the H354 insertion mutant protein degrades p53 but not MRN. We extended these studies by examining a number of E1b55K mutant viruses (56, 72) with either point mutations or small insertions across the length of E1b55K. A subset of mutations is shown in the schematic in Fig. 2.

Infections with mutant E1b55K viruses were compared to those with WT Ad5 and the E1b55K deletion virus dl110 with respect to the degradation of p53 and MRN over the time course of infection. In addition to the previously described H354 and R240A viruses, we found separation-of-function phenotypes for E1b55K mutant viruses R443 (which has a 4-amino-acid insertion at residue 443) (72) and Y444A (ONYX85) (56). While both viruses induced the degradation of p53, in agreement with previous reports (56, 58), the kinetics of p53 degradation was delayed for R443. We found that neither the R443 virus nor the Y444A virus exhibited complete downregulation of all three MRN complex members (Fig. 3). The R443 virus also produced more E1b55K than WT Ad5. It is interesting that the insertion creating the R443 mutation also results in a Y444D mutation (72), which may contribute to a compound phenotype. In the course of our survey, we found that the R443A mutation (in virus ONYX84) (56) did not affect the degradation of MRN (data not shown), suggesting that the region around position 443, rather than the actual residue at this position, is important to MRN downregulation.

Given that the MRN complex is required for the DNA damage response to mutant adenoviruses (12), we tested whether these mutants induced damage signaling. We observed that the partial degradation of MRN by R443 and Y444A was not sufficient to prevent signaling during viral infection (Fig. 3; also data not shown), as shown by Nbs1 phosphorylation (at S343). This finding suggests that infection with adenovirus mutants unable to neutralize MRN completely can induce DNA damage signaling. Together with our previous observations (12), these data indicate that multiple regions of E1b55K are important for MRN degradation. The complete survey results for mutant proteins are presented in Table 2, together with a summary of observations previously reported in the literature.
The phosphorylation of E1b55K is required for correct cellular localization and the degradation of substrates. Multiple posttranslational modifications of E1b55K have been reported, including SUMO-1 conjugation at K104 (21, 36, 40) and phosphorylation at the C terminus (62, 63). We wanted to address whether these modifications affected E1b55K localization or substrate degradation. E1b55K proteins harboring mutations at the SUMO modification site (K104) (21, 36, 40) and the known phosphorylation sites in the C terminus (S490, S491, and T495) (62, 63) were expressed from plasmids and first assessed for localization in the absence and presence of E4orf6 (52). H373A behaved similarly to H354 (12) in cellular localization and the degradation of p53 but not MRN (Fig. 4). R443 also appeared to be a separation-of-function mutant protein upon immunoblotting, although considering the levels of E1b55K, p53 degradation was not as robust as that by the WT. Together with our previous data (12), the results for these mutant proteins illustrate that the requirements for E1b55K-mediated degradation of MRN are distinct from those for the degradation of p53, and they identify two cysteine regions as critical for the proper localization of E1b55K and MRN degradation.

Using the stable E1b55K-expressing cell lines, we also examined E1b55K/E4orf6-mediated degradation of targets by immunoblotting (Fig. 4B). The previously characterized cell lines expressing green fluorescent protein (GFP), WT E1b55K, and mutant proteins R240A and H354 (12) were included as controls. The immunoblotting results confirmed our initial immunofluorescence observations and indicated that, with the exception of Y444A, most of the isolated mutant proteins were defective in MRN degradation (Fig. 4B and Table 2). C361,366S and C454,456S did not appear to degrade either p53 or MRN (Fig. 4B). This phenotype is unlikely to be due to defective E4orf6 binding since these mutant forms exhibited an interaction with E4orf6 similar to that of H224 (Fig. 4A; data not shown), a mutant protein competent to degrade all substrates but previously reported to exhibit reduced interaction with E4orf6 (52). H373A behaved similarly to H354 (12) in cellular localization and the degradation of p53 but not MRN (Fig. 4). R443 also appeared to be a separation-of-function mutant protein upon immunoblotting, although considering the levels of E1b55K, p53 degradation was not as robust as that by the WT. Together with our previous data (12), the results for these mutant proteins illustrate that the requirements for E1b55K-mediated degradation of MRN are distinct from those for the degradation of p53, and they identify two cysteine regions as critical for the proper localization of E1b55K and MRN degradation.

The changes in the E1b55K mutant proteins that we identified as defective for MRN degradation (H354, R443, and Y444A) are located in or near two regions that contain cysteine residues and may resemble zinc fingers (26). To analyze E1b55K further, we cloned the genes for these and other site-specific mutant forms into retroviral vectors (Fig. 2). Stable cell lines expressing individual E1b55K proteins were generated by retrovirus transduction as described previously (12). Mutant proteins were examined by immunofluorescence analysis in the presence and absence of E4orf6 for cellular localization and the degradation of the MRN complex. In the absence of E4orf6, WT E1b55K was localized in cytoplasmic aggregates (Fig. 4A), as described previously (27, 37). In the presence of E4orf6, E1b55K accumulated in a diffuse nuclear pattern with few remaining cytoplasmic foci and this outcome was accompanied by the degradation of Nbs1. The E1b55K mutant proteins exhibited a variety of patterns, which are shown in Fig. 4A and summarized in Table 2. Those with patterns differing from the WT E1b55K pattern included the R443 protein, which displayed predominantly diffuse cytoplasmic staining, and the Y444A protein, which showed more elongated cytoplasmic foci (Fig. 4A). With the exception of the Y444A and C348,351S proteins, the mutant forms with changes near cysteine regions were unable to degrade MRN in the presence of E4orf6 (Fig. 4A and Table 2), regardless of their localization patterns. Many mutant proteins were not retained in the nucleus to the same extent as WT E1b55K in the presence of E4orf6 (Fig. 4A and Table 2). However, this phenotype did not correlate completely with the lack of MRN degradation, since mutant forms H224 and Y444A were still able to degrade MRN (Fig. 4A). This finding suggests that nuclear retention by E4orf6 is not absolutely required for the degradation of cellular substrates.

Two cysteine-containing regions of E1b55K are important for substrate degradation. The changes in the E1b55K mutant proteins that we identified as defective for MRN degradation (H354, R443, and Y444A) are located in or near two regions that contain cysteine residues and may resemble zinc fingers (26). To analyze E1b55K further, we cloned the genes for these and other site-specific mutant forms into retroviral vectors (Fig. 2). Stable cell lines expressing individual E1b55K proteins were generated by retrovirus transduction as described previously (12). Mutant proteins were examined by immunofluorescence analysis in the presence and absence of E4orf6 for cellular localization and the degradation of the MRN complex. In the absence of E4orf6, WT E1b55K was localized in cytoplasmic aggregates (Fig. 4A), as described previously (27, 37). In the presence of E4orf6, E1b55K accumulated in a diffuse nuclear pattern with few remaining cytoplasmic foci and this outcome was accompanied by the degradation of Nbs1. The E1b55K mutant proteins exhibited a variety of patterns, which are shown in Fig. 4A and summarized in Table 2. Those with patterns differing from the WT E1b55K pattern included the R443 protein, which displayed predominantly diffuse cytoplasmic staining, and the Y444A protein, which showed more elongated cytoplasmic foci (Fig. 4A). With the exception of the Y444A and C348,351S proteins, the mutant forms with changes near cysteine regions were unable to degrade MRN in the presence of E4orf6 (Fig. 4A and Table 2), regardless of their localization patterns. Many mutant proteins were not retained in the nucleus to the same extent as WT E1b55K in the presence of E4orf6 (Fig. 4A and Table 2). However, this phenotype did not correlate completely with the lack of MRN degradation, since mutant forms H224 and Y444A were still able to degrade MRN (Fig. 4A). This finding suggests that nuclear retention by E4orf6 is not absolutely required for the degradation of cellular substrates.

Using the stable E1b55K-expressing cell lines, we also examined E1b55K/E4orf6-mediated degradation of targets by immunoblotting (Fig. 4B). The previously characterized cell lines expressing green fluorescent protein (GFP), WT E1b55K, and mutant proteins R240A and H354 (12) were included as controls. The immunoblotting results confirmed our initial immunofluorescence observations and indicated that, with the exception of Y444A, most of the isolated mutant proteins were defective in MRN degradation (Fig. 4B and Table 2). C361,366S and C454,456S did not appear to degrade either p53 or MRN (Fig. 4B). This phenotype is unlikely to be due to defective E4orf6 binding since these mutant forms exhibited an interaction with E4orf6 similar to that of H224 (Fig. 4A; data not shown), a mutant protein competent to degrade all substrates but previously reported to exhibit reduced interaction with E4orf6 (52). H373A behaved similarly to H354 (12) in cellular localization and the degradation of p53 but not MRN (Fig. 4). R443 also appeared to be a separation-of-function mutant protein upon immunoblotting, although considering the levels of E1b55K, p53 degradation was not as robust as that by the WT. Together with our previous data (12), the results for these mutant proteins illustrate that the requirements for E1b55K-mediated degradation of MRN are distinct from those for the degradation of p53, and they identify two cysteine regions as critical for the proper localization of E1b55K and MRN degradation.
of E4orf6 expressed after transfection (Fig. 5A) or during infection (Fig. 5B) with the 3XPA mutant virus, pm490/1A (62), we observed increased nuclear staining, although a large amount of E1b55K also remained in the cytoplasm. The 2XPA double phosphorylation site mutant virus, pm490/1A (62), displayed a more predominantly nuclear E1b55K staining pattern (Fig. 5B). To address whether phosphorylation was required for the proper localization of E1b55K, we constructed a phosphomimic mutant protein in which the amino acids at positions 490, 491, and 495 were changed to aspartic acid (3XPD). After transient transfection with the construct expressing this mutant protein, we observed a cellular localization pattern more similar to WT E1b55K-expressing cell line. The localization pattern in these stable cell lines was similar to that seen after transient transfection, although less nuclear staining in the presence of E4orf6 was noted (data not shown). The degradation of cellular targets in these cell lines was assessed with E4orf6 alone (Fig. 5A) or during infection (Fig. 5B) with the 3XPA mutant virus, pm490/1A (62). We then examined how the SUMO and phosphorylation site mutations affected substrate downregulation. Immunofluorescence analyses revealed that the mutant proteins were competent to degrade MRN in the presence of E4orf6 (Fig. 5A and Table 2). To analyze further the requirement of E1b55K phosphorylation for degradation, we generated stable U2OS cell lines expressing the 3XPA and 3XPD proteins and compared them to the WT E1b55K-expressing cell line. The localization pattern in these stable cell lines was similar to that seen after

### Table 2. Summary of E1b55K mutant phenotypes

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<tr>
<th>Protein</th>
<th>Substrate degradation&lt;sup&gt;a&lt;/sup&gt; phenotype (during coexpression with E4orf6 alone) for:</th>
<th>Substrate degradation&lt;sup&gt;a&lt;/sup&gt; phenotype (during virus infection) for:</th>
<th>Substrate binding&lt;sup&gt;b&lt;/sup&gt; phenotype for:</th>
<th>Aggregate colocalization&lt;sup&gt;c&lt;/sup&gt; phenotype for:</th>
<th>Cytoaggregate phenotype</th>
<th>Nuclear retention by E4orf6&lt;sup&gt;d&lt;/sup&gt;</th>
<th>Concatemer formation&lt;sup&gt;e&lt;/sup&gt; phenotype</th>
<th>Additional references&lt;sup&gt;f&lt;/sup&gt;</th>
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<sup>a</sup> Unless otherwise indicated, symbols and abbreviations are as follows: +, proficient (may not reflect WT E1b55K activity); ±, defective; −, very defective; ND, not determined; NA, not applicable.

<sup>b</sup> Degradation of endogenous proteins by E1b55K mutant forms in the presence of rAdE4orf6 as assessed by Western blotting or immunofluorescence analysis.

<sup>c</sup> Degradation of endogenous proteins assessed in the context of viruses expressing E1b55K mutant proteins or E1b55K mutant cell lines infected with d3112.

<sup>d</sup> Interaction measured by coinmunoprecipitation experiments.

<sup>e</sup> MRN or p53 colocalization with E1b55K aggregates as assessed by immunofluorescence.

<sup>f</sup> E1b55K staining in the absence of rAdE4orf6 was assayed by using immunofluorescence. +, presence of E1b55K aggregates; diffuse, diffuse cytoplasmic E1b55K. See Fig. 4A and the corresponding legend for data and details.

<sup>g</sup> E1b55K staining in the presence of rAdE4orf6 was assayed by using immunofluorescence. See Fig. 4A and the corresponding legend for data and details.

<sup>h</sup> Concatemer formation by mutant virus d3112 measured in mutant E1b55K-expressing cells.

<sup>i</sup> Observation from the present study based on E1b55K expression in human cells.

<sup>j</sup> This table summarizes data from this paper as well as the additional references indicated.
MRN and p53 degradation, the proteins C361,366S and C454,456S were defective for the degradation of DNA ligase IV (Fig. 6A). The 3XPA mutant protein displayed reduced DNA ligase IV downregulation (Fig. 5C and D and 6A). Interestingly, we found that the separation-of-function mutant proteins could all degrade DNA ligase IV, regardless of p53 or MRN specificity (Fig. 6A). This pattern was also true for most of the E1b55K mutant viruses (Fig. 3 and Table 2), although the R443 virus had a slight defect (Fig. 3A). These data indicate that the requirements of E1b55K for DNA ligase IV degradation are distinct from those for both p53 and MRN degradation.

The concatemerization of mutant viral genomes requires both the MRN complex and DNA ligase IV (60). Therefore, E1b55K proteins unable to degrade MRN may still be able to prevent the concatemerization of a mutant virus if DNA ligase IV is downregulated. We examined whether concatemers formed during the infection of E1b55K mutant cell lines (Fig. 6B). Since the E4orf3 protein can independently prevent concatemers (60), we utilized the d3112 mutant virus, which has both E1b55K and E4orf3 deleted (1, 57). As expected, mutant viral genomes were concatemerized in GFP-expressing control cells but not in cells expressing WT E1b55K protein. We also found that concatemers were inhibited in mutant cell lines proficient in the degradation of DNA ligase IV alone (those expressing H354 or H373A) or in cell lines in which both MRN and DNA ligase IV were degraded (those expressing H224, R240A, Y444A, or 3XPD) (Fig. 6B and Table 2). There was no correlation between p53 degradation and concatemers (Table 2). Additionally, the cell lines expressing C361,366S and C454,456S, which were unable to downregulate any substrates, could not prevent viral genome concatemerization (Fig. 6B and Table 2). In these experiments, we found that mutant proteins R443 and 3XPA could not fully complement d3112 to prevent concatemer formation and degrade MRN or DNA ligase IV sufficiently (Table 2). These data confirm the require-
FIG. 5. The phosphorylation of E1b55K is required for correct cellular localization and substrate degradation. (A) U2OS cells were transfected with constructs expressing WT E1b55K, E1b55K mutant protein 3XPA, or E1b55K phosphomimic 3XPD. After 16 h, cells were superinfected with rAdE4orf6 (MOI of 20) for 24 h before being processed for immunofluorescence analysis with the indicated antibodies. Cell nuclei are stained with DAPI in the merged images. –E4orf6, without E4orf6; +E4orf6, with E4orf6. (B) U2OS cells were infected with WT Ad5, the ΔE1b55K mutant d/l10, or E1b55K mutant virus pm490/1/5A (expressing 3XPA) or pm490/1A (expressing 2XPA) for approximately 24 h. Cells were fixed and analyzed for E1b55K localization by using immunofluorescence. DAPI indicates cell nuclei in the merged images. (C) Cell lines expressing E1b55K were infected with rAdE4orf6 (MOI of 50) for 24 h. Cells were harvested, and lysates were analyzed by immunoblotting with the indicated antibodies. +, present; –, absent. (D) U2OS cells were infected with WT Ad5, the ΔE1b55K mutant d/l10, or E1b55K phosphomutant virus pm490/1/5A (expressing 3XPA; left panel) or pm490/1A (expressing 2XPA; right panel), all at an MOI of 50. Cells were harvested at the times postinfection indicated above the gels and processed for immunoblotting with the indicated antibodies. GAPDH served as a cellular loading control.
E1b55K separation-of-function mutant proteins still degrade DNA ligase IV to prevent concatemer formation. (A) U2OS-based cell lines stably expressing WT and mutant E1b55K proteins were either mock treated or infected with rAdE4orf6 (MOI of 50) for 24 h. Lysates were analyzed by immunoblotting for DNA ligase IV and GAPDH (loading control). +, present; −, absent. (B) E1b55K mutant proteins that cannot degrade MRN still retain the ability to prevent concatemer formation. U2OS cells expressing E1b55K mutant forms were infected with df3112 at an MOI of 10 for 30 h, and viral DNA was analyzed by pulsed-field gel electrophoresis as described in Materials and Methods. Controls included a GFP-expressing U2OS cell line that was either mock treated (M) or infected with df3112 to produce concatemers. An arrowhead indicates the position of the linear viral genome.

ment of DNA ligase IV for concatemer formation (60). They also demonstrate that the downregulation of DNA ligase IV is sufficient to prevent the joining of viral genomes even when MRN is not degraded.

E1b55K separation-of-function mutant proteins have defects in substrate binding. We next examined how mutations in E1b55K affect interactions with cellular degradation substrates. The expression of E1b55K in cells can induce a large, perinuclear accumulation of the protein with characteristics of an aggresome (42). Many cellular proteins interacting with E1b55K have been observed to colocalize with E1b55K aggresomes (4, 22, 44, 74). We therefore examined the ability of E1b55K mutant proteins to concentrate MRN and p53 into cytoplasmic aggregates (Fig. 7A). We found that mutant proteins which degraded MRN (H224, R240A, Y444A, and 3XPD) could all relocalize Nbs1 to cytoplasmic aggregates, although some mutant forms were less efficient than the WT E1b55K protein (Fig. 7A and Table 2). Interestingly, we found that all E1b55K mutant forms concentrated p53 into cytoplasmic aggregates to some extent (Fig. 7A and Table 2). Surprisingly, despite their inability to downregulate the p53 substrate, even the C361,366S and C454,456S proteins relocalized p53, R240A, however, was extremely defective in this activity (Fig. 4B), consistent with its diminished capacity to degrade p53 (12, 56).

To confirm our aggresome immunofluorescence results, we generated mutant E1b55K constructs that also contained a mutation (L83,87,91A) in the NES (39). These E1b55K proteins were expressed by transfection in U2OS cells and analyzed by immunofluorescence (Fig. 7B). Cells expressing E1b55K with a mutated NES displayed nuclear aggregates of E1b55K staining, as previously reported (20). The endogenous cellular Nbs1 and p53 proteins appeared to colocalize with E1b55K foci, suggesting possible interaction (Fig. 7B). When our mutations were combined with the NES mutation, all proteins tested formed distinct nuclear foci, but there were differences in their abilities to redistribute cellular proteins. Consistent with the degradation data, NES-R240A relocalized Nbs1 efficiently but not p53, whereas NES-H373A and NES-H354 recruited p53 efficiently but not Nbs1. Together with the aggresome immunofluorescence data, these results suggest that most mutant proteins with defects in degradation also have defects in binding substrates. However, the cysteine mutant forms (C361,366S and C454,456S) may have additional defects related to p53 degradation, since they can still bind and relocalize this substrate.

To validate the immunofluorescence results, we performed coimmunoprecipitation experiments with E1b55K mutant proteins in the absence of E4orf6 (Fig. 7C and Table 2). E1b55K was immunoprecipitated from lysates of U2OS E1b55K mutant cells with the 2A6 antibody and analyzed for MRN and p53 interaction. Because it was difficult to resolve p53 from the background immunoglobulin band in E1b55K immunoprecipitates, we performed a reverse immunoprecipitation against p53 and probed for E1b55K interaction (Fig. 7C, bottom panels). Most mutant forms that failed to degrade either MRN or p53 also failed to interact with that particular substrate (Fig. 7C and Table 2), confirming the immunofluorescence results. In the absence of E4orf6, MRN binding to Y444A and 3XPA was significantly weaker than that to WT E1b55K. This interaction, however, was sufficient to induce the degradation of MRN in the presence of E4orf6 (Fig. 4 and 5). This finding suggests that E4orf6 may enhance E1b55K interaction with MRN by relocalizing E1b55K to the nucleus. We also found that the cysteine mutant proteins C361,366S and C454,456S were defective for MRN interaction but still retained the ability to bind p53, although not to the same extent as WT E1b55K (Table 2). Unfortunately, we were unable to examine endogenous DNA ligase IV by using immunofluorescence and could not detect an interaction between E1b55K and endogenous DNA ligase IV in our immunoprecipitation experiments (data not shown). Together, our data suggest that the E1b55K separation-of-function mutant proteins described here are defective in degradation due to diminished substrate binding. The ability of the cysteine mutant forms to bind p53 in the absence of downregulation suggests that there are additional E1b55K requirements for substrate degradation.
FIG. 7. Defects in substrate binding of E1b55K mutant proteins. (A) E1b55K mutant proteins colocalize with cellular substrates in cytoplasmic aggresomes/aggregates. U2OS cells were transfected with E1b55K mutant constructs to induce cytoplasmic aggresome/aggregate formation. Cells were processed for immunofluorescence analysis after 24 to 36 h by using antibodies to the indicated proteins. DAPI staining marks cell nuclei. Arrowheads indicate the colocalization of substrates with E1b55K in cytoplasmic aggregates. (B) E1b55K NES mutant proteins relocalize cellular targets into nuclear aggregates. U2OS cells were transfected with constructs expressing E1b55K proteins with NES mutations. Cells were processed for immunofluorescence analysis after 24 h by using antibodies to the indicated proteins. DAPI staining marks cell nuclei. (C) Lysates from mutant E1b55K cell lines were subjected to immunoprecipitation (IP) with the 2A6 antibody to E1b55K or a p53 antibody as described in Materials and Methods. Approximately half the immunoprecipitate was analyzed by immunoblotting alongside 5% of the input lysate.
DISCUSSION

In this study, we examined requirements for adenovirus-mediated degradation of the known cellular substrates of the E1b55K/E4orf6 ubiquitin ligase complex: the MRN proteins, p53, and DNA ligase IV. We first addressed whether the degradation of each of these cellular factors required other proteins in the DNA damage response. While our previous use of E1b55K mutant proteins suggested that MRN degradation and p53 degradation are separable events (12), it was not clear whether the prior absence of cellular factors would affect the degradation of the other substrates. Using mutant cell lines, we found that adenovirus downregulates each substrate independently of the other substrates and does not require other DNA damage response proteins (Fig. 1B). Additionally, after examining adenovirus infections in A-TLD and NBS mutant cell lines, we suggest that Mre11 is the degradation target of E1b55K/E4orf6 (Fig. 1C). The downregulation of Rad50 and Nbs1, therefore, may be an indirect consequence of complex destabilization, as noted in other studies using RNA interference against particular members of the MRN complex in the absence of virus infection (64, 77). It is possible, however, that Rad50 is the target but that the protein is not recognized by E1b55K in the absence of Mre11 due to conformational changes. Further analysis of protein-protein interactions will be required to resolve the target.

We analyzed a large collection of E1b55K mutant forms in order to define further the viral requirements for degradation. We found that the downregulation of the MRN complex is separable from p53 and DNA ligase IV degradation, as discerned by using a number of E1b55K mutant proteins. The results for these mutant proteins indicate that at least two C-terminal cysteine-containing regions are critical to MRN degradation (Fig. 2 and Table 2). These regions include a proposed zinc finger motif (23) and a transcriptional repression domain (71, 73). In the context of viral infection, mutants with defects in MRN degradation cannot completely prevent DNA damage signaling (Fig. 3 and 5D) (12). However, some of these mutants can still prevent viral genome concatenation. This effect appears to occur through the degradation of DNA ligase IV (Fig. 6), although we cannot exclude targeting of additional, unidentified substrates. Many of the mutant proteins analyzed could degrade DNA ligase IV independently of their p53 and MRN phenotypes, which suggests that there are distinct E1b55K requirements for DNA ligase IV destabilization.

Further analysis of the panel of E1b55K mutant proteins also identified cysteine mutant forms (C361,366S and C454,456S) with multiple defects that may affect degradation. In our studies, these mutant proteins were unable to downregulate any of the known cellular substrates (Fig. 4 and 6). We found that these cysteine mutant forms were defective in their association with MRN, but not with p53 (Fig. 7). This result is consistent with the location of the mutations outside the previously defined p53 interaction domain in E1b55K (71). In contrast to our findings, Hartl et al. recently observed that the C454,456S protein degraded exogenously expressed p53 and mislocalized Rad50 but not Mre11 in transformed baby rat kidney cells (31). The discrepancies may reflect different functions of E1b55K in transformed rodent cells versus human cells and the fact that we examined degradation specifically for endogenous proteins. From our data, it appears that the cysteine mutant proteins have an additional defect related to p53 degradation. The cysteine amino acids mutated in these E1b55K proteins may be important for modifying degradation substrates with ubiquitin or another marker required for ubiquitination and degradation. Unfortunately, we could not detect E1b55K interactions with endogenous DNA ligase IV, so it is unclear why the cysteine mutant proteins do not downregulate this substrate. Further insights will come from studies that delineate regions in E1b55K important for binding and degrading DNA ligase IV.

We examined a number of mutant proteins to address whether posttranslational modifications of E1b55K are important for the degradation of substrates. We found that the sumoylation site in E1b55K could be modified without impairing MRN degradation (Table 2), in agreement with a previous report (36). Phosphorylation sites in the C terminus appeared to be important for both the degradation of cellular substrates and the correct cellular localization of E1b55K (Fig. 5 and 6). Consistent with our data, a similar Ad12 E1b55K mutant protein (S476,477A) was also found previously to be defective in the formation of cytoplasmic aggregates (76). Phosphorylation may alter protein conformation, thereby affecting intramolecular E1b55K interactions or association with partner proteins. Along these lines, we found by coimmunoprecipitation that the interaction between 3XPA and p53 was severely defective or unstable (Table 2), consistent with the inability of this mutant protein to prevent p53-mediated transactivation and promote cellular transformation (62). The phosphorylation of all three amino acids may be required, however, since 2XPA still degraded p53 (Fig. 5D) (50) and other cellular substrates (Fig. 5D) and retained the ability to bind p53 (63). The interaction between 3XPA and MRN was significantly weaker than that between WT E1b55K and MRN (Fig. 7), although it was sufficient to induce MRN degradation in the presence of rAdE4orf6 (Fig. 5A and C). The phosphorylated amino acids in E1b55K lie within consensus casein kinase I and casein kinase II motifs (62, 63). However, it is still unclear whether these kinases are responsible for the phosphorylation of E1b55K in cells. The identification of kinases that modify E1b55K will provide more insight into the effects of phosphorylation on localization, structure, and protein-protein interactions important to function. It is interesting that the R443 protein also displayed a diffuse cytoplasmic staining pattern in U2OS stable cell lines (Fig. 4A), although it still formed some small, perinuclear foci. Therefore, the C terminus of E1b55K may be important to its structure and localization. While we do not know whether phosphorylation is affected for this mutant protein, R443 retained stable p53 binding (Fig. 7) (34, 71), unlike 3XPA.

Our analysis of E1b55K mutant proteins illustrated differences in degradation by proteins coexpressed with E4orf6 alone compared to degradation in the context of virus infection. We noticed a particular difference with proteins mutated in the C terminus (Y444A, R443, and 3XPA), which showed greater defects than the other proteins in the degradation of substrates during viral infection (compare Fig. 3 and 5D to Fig. 4B). Higher levels of E4orf6 expressed from the recombinant adenovirus than in the context of virus infection may partly
contribute to these observations. The results of experiments with E1b55K mutant proteins and E4orf6 alone are useful in delineating the absolute requirements for degradation. For example, the phosphorylation of E1b55K may not be absolutely required for MRN degradation but may contribute more to the degradation of p53 and DNA ligase IV (Fig. 5C). Interactions between viral proteins are likely to be far more complex in the context of virus infection, and therefore, it will be informative to incorporate some of the E1b55K mutant proteins analyzed here into the virus to characterize further their phenotypes. It will also be interesting to examine degradation in primary human cells, in order to determine whether the same requirements hold true.

Although the degradation of p53, MRN, and DNA ligase IV induced by E1b55K/E4orf6 appears to be proteasome mediated, only p53 has been shown to be a direct ubiquitination target (30, 49). It will be important to determine whether the ubiquitination of the MRN complex and DNA ligase IV by the E1b55K/E4orf6 ubiquitin ligase is directly involved in degradation. This is likely to be the case, since the downregulation of all these substrates is prevented by the expression of transdominant cullin 5 (43) and by small interfering RNA knockdown of cullin 5 (R. A. Schwartz and M. D. Weitzman, unpublished data; 2, 43, 70). Other proteins that interact with E1b55K have been identified by genetic, biochemical, and proteomic studies (22, 24, 30, 44, 75); however, not all of these interacting proteins are targets for downregulation. It will be interesting to ascertain what determines the ubiquitination and interacting proteins are targets for downregulation. It will be informative to incorporate some of the E1b55K mutant proteins analyzed here into the virus to characterize further their phenotypes. It will also be interesting to examine degradation in primary human cells, in order to determine whether the same requirements hold true.

ACKNOWLEDGMENTS

We thank A. Berk, P. Branton, P. Concannon, G. Ketner, A. Levine, T. Melendy, D. Ornelles, J. Petrini, Y. Shen, and H. Young for generous gifts of reagents. We thank Daniel Linfesty and Darwin Lee for technical assistance. We thank members of the Weitzman lab for technical assistance. We thank members of the Weitzman lab for technical assistance. We thank members of the Weitzman lab for technical assistance. We thank members of the Weitzman lab for technical assistance. We thank members of the Weitzman lab for technical assistance. We thank members of the Weitzman lab for technical assistance. We thank members of the Weitzman lab for technical assistance.

REFERENCES

34. Kao, C. C., P. R. Yew, and A. J. Berk. 1990. Domains required for in
38. Kratzer, F., O. Rosorius, P. Heger, N. Hirschmann, T. Dobner, J. Hauber,
68. Wenzel, S., J. Roth, and M. Dobbelstein. 2000. E1B 55-kilodalton onco-