Apoptosis is a specific form of cell death that is important for normal development and tissue homeostasis. Caspases are critical executioners of apoptosis, and living cells prevent their inappropriate activation through inhibitor of apoptosis proteins (IAPs). In Drosophila, caspase activation depends on the IAP antagonists, Reaper (Rpr), Head involution defective (Hid), and Grim. These proteins share a common motif to bind Drosophila IAP1 (DIAP1) and have partially redundant functions. We now show that IAP antagonists physically interact with each other. Rpr is able to self-associate and also binds to Hid and Grim. We have defined the domain involved in self-association and demonstrate that it is critical for cell-killing activity in vivo. In addition, we show that Rpr requires Hid for recruitment to the mitochondrial membrane and for efficient induction of cell death in vivo. Both targeting of Rpr to mitochondria and forced dimerization strongly promotes apoptosis. Our results reveal the functional importance of a previously unrecognized multimeric IAP antagonist complex for the induction of apoptosis.

Introduction

Apoptosis is a genetically encoded process of cell death with defined morphological features that serves to kill superfluous or unwanted cells, and abnormal regulation of this process is associated with many human diseases (Steller, 1995; Thompson, 1995; Yuan and Yankner, 2000). An evolutionarily conserved feature of apoptosis is the activation of a particular class of proteases, termed caspases (Thornberry and Lazebnik, 1998), which cleave many vital structural and regulatory proteins in the cell (Hengartner, 2000). Activation of caspases is kept in check by a conserved class of anti-apoptotic proteins, termed inhibitor of apoptosis proteins (IAPs; Reed et al., 2000; Shiozaki and Shi, 2004). IAPs can bind to both initiator and effector caspases via their BIR domains (Shi, 2002; Bergmann et al., 2003). Furthermore, many IAPs also contain a RING motif and act as E3 ubiquitin ligases to ubiquitinate cell death proteins, including caspases (Wilson et al., 2002; Tenev et al., 2005). In Drosophila, DIAP1 is strictly required to prevent caspase activation and apoptosis in virtually all somatic cells (Wang et al., 1999; Goyal et al., 2000; Lisi et al., 2000). In cells that are doomed to die, IAPs are inactivated by specific antagonists (Vucic et al., 1997; Goyal et al., 2000; Yan et al., 2004).

In Drosophila, three IAP antagonists, Reaper (Rpr), Head involution defective (Hid), and Grim are clustered together in the genome, and deleting these genes causes a severe inhibition of apoptosis (White et al., 1994; Grether et al., 1995; Chen et al., 1996). A fourth IAP antagonist, Sickle (Skl), was also identified with significant similarity to Rpr (Srinivasula et al., 2002), but due to the lack of mutants its physiological role for the induction of apoptosis is less clear. One evolutionarily conserved feature is the presence of the N-terminal IBM (IAP-binding motif), a stretch of several amino acids that interacts with the BIR domains of IAPs (Vucic et al., 1998; Shi, 2002). IAP antagonists bind IAPs and displace competitively IAP-bound caspases (Holley et al., 2002; Chai et al., 2003; Zachariou et al., 2003). Active caspases propagate a proteolytic cascade that will compromise the cell’s infrastructure and metabolism. Another aspect of IAP antagonists’ function is to stimulate IAP turnover by proteasomal degradation (Ryoo et al., 2002; Yoo et al., 2002). When expressed in human cells, Drosophila IAP antagonists preserve similar activities such as inducing cell death (McCarthy and Dixit, 1998; Haining et al., 1999) and binding and stimulating human IAP degradation (Silke et al., 2004). Humans also have IAP antagonists, among which the best characterized is...
Smac/Diablo (Du et al., 2000; Verhagen et al., 2000). Smac forms dimers and interacts with the BIR domains of XIAP (Wu et al., 2000), and yet the significance of dimer formation is not known. Other human IAP antagonists include HtrA2/Omi (also present in Drosophila; Hegde et al., 2002) and ARTS (Gottfried et al., 2004).

Rpr is a small protein of 65 amino acids (White et al., 1994). Previous reports have suggested that Rpr (Olson et al., 2003a), Hid (Haining et al., 1999), and Grim (Clavería et al., 2002) localize to the mitochondria. Rpr and Grim share a homologous motif outside of IBM, known as the GH3 (Grim helix 3), which is required for their mitochondrial localization (Clavería et al., 2002; Olson et al., 2003a). Disruption of this GH3 motif in Rpr not only impairs its mitochondrial translocation, but also disrupts Rpr’s ability to stimulate DIAP1 auto-ubiquitination and degradation (Freeel et al., 2008).

The fact that rpr, hid, and grim share homologous IBMs and that this motif binds to specific pockets in the DIAP1 BIR domains (Wu et al., 2001; Chai et al., 2003; Yan et al., 2004) has led to the idea that the IAP antagonists have partially redundant roles. Here, we provide evidence that these proteins work together as a high-order physical complex for efficient DIAP1 inactivation. Specifically, we present a structure-function analysis of Rpr that reveals the importance of a central helical domain in dimerization, the formation of multimeric complexes with other IAP antagonists, protein localization, and the ability of Rpr to promote DIAP1 degradation.

Results

Rpr self-association is essential for its apoptotic activity

To understand how Rpr interacts with DIAP1 to induce its ubiquitination, we investigated the interaction between Rpr, DIAP1, and other related apoptosis regulator proteins. In the absence of a Rpr three-dimensional structure, we have performed a secondary structure prediction to identify structural elements in the amino acid sequence. Rpr consists of three major elements, the IBM motif (residues 1–9), a central helical domain (residues 10–48) that includes the GH3 motif (Olson et al., 2003a) and adopts an α-helical conformation, and a C-terminal unstructured tail (residues 49–65) (Fig. 1 A). Because many protein helical domains are involved in protein–protein interaction, we hypothesized that Rpr might interact with self or with other proteins through this helical domain. Supporting this idea, Rpr-GST recombinant protein was able to pull down 35S radiolabeled Rpr in vitro (Fig. 1 B). The interaction proved to be specific because control GST was not able to pull down 35S-Rpr (Fig. 1 B).

Next, we set out to identify the amino acids involved in this interaction by introducing a number of point mutants that span the entire Rpr helical domain through site-directed mutagenesis (represented in Fig. 1 A). In support of our hypothesis, three mutants, Q23ER26A, F34AL35A, and Q22AQ23AG54E, were found to have reduced affinity for Rpr-GST (Fig. 1 C). Subsequently, we investigated the functional relevance of the mutations that disrupt Rpr self-association in vivo. Specifically, we generated fly transgenes in which Rpr-HA, mutant Q23ER26A, and the GH3 mutant F34AL35A were targeted to a defined genomic locus by Cre-mediated recombination (Oberstein et al., 2005) and compared their pro-apoptotic activity when expressed in developing Drosophila eyes, using the GMR>Gal4/UAS system. As previously reported, Rpr-HA–induced expression in the eyes produced severe eye ablation (White et al., 1996). On the other hand, the Q23ER26A variant had only mild apoptotic activity, whereas the GH3 mutant F34AL35A was completely inactive, yielding normal eyes (Fig. 1 D). These results support the idea that Rpr self-association is important for the protein’s apoptotic activity, and the disruption of self-association blocks protein activity in vivo.

Enforced dimerization of Rpr leads to efficient cell killing

Although the above experiments indicate that Rpr self-association is required for its pro-apoptotic activity, whether it is sufficient to recapitulate Rpr’s pro-apoptotic function remained unclear. To test this, we replaced the helical domain of Rpr (residues 10–46) with well-defined dimerization domains from heterologous proteins whose three-dimensional structures have been previously determined. Specifically, we used a parallel leucine zipper (LZ) from the yeast transcription factor GCN4 (O’Shea et al., 1991) and an anti-parallel coiled-coil domain from the Escherichia coli osmosensor ProP (Zoetewey et al., 2003; Fig. 2 A). When these chimeric proteins were expressed in the fly eye using the GMR>Gal4/UAS system we found that RprLZ triggered massive cell death, as evidenced by the partially ablated eye structure (Fig. 2 B), supporting the idea that Rpr dimerization is sufficient to account for its central helical domain’s function. On the other hand, RprProP did not trigger cell death under similar conditions, despite being expressed at similar levels with RprLZ (Fig. 2 C). Next, we examined whether RprLZ induces cell death through a mechanism similar to the wild-type Rpr, namely the inhibition of DIAP1 and activation of caspases. Supporting the requirement of caspase activation, coexpression of p35, a well-established caspase inhibitor of viral origin, rescued the eye morphology caused by RprLZ (Fig. 2 D, right) as well as wild-type Rpr (Fig. 2 E, right). To test the requirement of DIAP1 inactivation, we took advantage of the diapr16-3s and diapr23-4s alleles, which are endogenous alleles bearing point mutations in the IBM-binding pocket of DIAP1 BIR domains, making cells resistant to Rpr-induced cell death (Goyal et al., 2000). The presence of these diapr alleles in the background significantly suppressed apoptosis induced by the RprLZ (Fig. 2 D) as well as wild-type Rpr (Fig. 2 E). Next, we examined the ability of RprLZ to induce DIAP1 degradation. In coexpression experiments in HEK293 cells, RprLZ was able to stimulate DIAP1ΔR degradation to significant extent (Fig. 2 F, right), but lower than wild-type Rpr (Fig. 2 F, left). DIAP1ΔR was used instead of full-length DIAP1 due to its increased stability (not depicted). Ability of RprLZ to induce DIAP1 degradation was also shown in wing discs, after overexpression in the presence of p35 (Fig. 2 G). These results support the idea that RprLZ has pro-apoptotic mechanisms similar to that of wild-type Rpr.
Rpr physically interacts with other Drosophila IAP antagonists

Next, we asked whether Rpr interacts with the other Drosophila IAP antagonists. Specifically, we tested potential interactions between Rpr-GST and 35S-Hid, 35S-Grim, or 35S-Skl through in vitro pull-down assays. We found that Rpr can interact with the other Drosophila IAP antagonists Hid and Grim, but not with Skl (Fig. 3 A). Under identical conditions, a control GST protein did not interact with 35S-Hid, 35S-Grim, or 35S-Skl, indicative of the specificity of the observed interactions (Fig. 3 A). The Rpr–Rpr and Rpr–Hid interactions were further confirmed by alternative pull-down experiments using Rpr-GST as “bait” and purified ubiquitin (Ub), Rpr, and HidΔMTS as “prey”. Besides confirming the specific Rpr–Rpr interaction, this experiment also indicates that Rpr–Hid interaction is not dependent on Hid’s MTS (Fig. 3 B). The interaction between Rpr and Hid was further confirmed by a reverse pull-down assay, using purified GST-Hid as bait and purified Rpr as a prey (Fig. 3 C). Moreover, we have performed competitive displacement experiments, where preformed Rpr-GST:Rpr-GST complexes were incubated with increasing amounts of HidΔMTS or Rpr and could see displacement of proteins in these complexes (unpublished data). Next, we asked whether Rpr interacts with Hid using the same domain used for self-association. To this end, a Hid-Flag construct was cotransfected with Rpr-Myc or the GH3 mutant F34AL35A constructs in HEK293 cells followed by anti-Flag immunoprecipitation. Interestingly, Hid communoprecipitated with wild-type Rpr but not with the GH3 mutant F34AL35A (Fig. 3 D). This experiment argues that Rpr uses the same domain for protein association, either with self or with Hid. Alternatively, Rpr dimers interact with Hid in an oligomeric complex, and disrupting Rpr dimer interface blocks the formation of an oligomeric complex with Hid.
assess protein localization failed to identify any motifs for specific subcellular localization. This raises the possibility that Rpr localizes to the mitochondria through a novel mechanism. Thus, we decided to investigate Rpr localization by ectopic coexpression. Although Rpr is known to localize to mitochondria (Olson et al., 2003a; Abdelwahid et al., 2007), prediction tools used to identify motifs failed to identify any motifs for specific subcellular localization. This raises the possibility that Rpr localizes to the mitochondria through a novel mechanism. Therefore, we decided to investigate Rpr localization by ectopic coexpression.

**Figure 2.** Enforced Rpr dimers kill by apoptosis in *Drosophila*. (A) Amino acid sequences and structural elements of Rpr dimers. RprLZ is an enforced parallel Rpr dimer where Rpr helical region (residues 10–46) was replaced with a parallel leucine zipper (GCN4), whereas RprProP is an enforced anti-parallel Rpr-dimer. LZ and ProP amino acid sequences are represented in blue. Residues in brown were inserted on both sides of each dimerization domain to preserve the same length as wild-type Rpr. IBM, and Tail, are identical as in wild-type Rpr. Rpr is targeted to mitochondria via interaction with Hid.

Rpr is targeted to mitochondria via interaction with Hid.

Although Rpr is known to localize to mitochondria (Olson et al., 2003a; Abdelwahid et al., 2007), prediction tools used to assess protein localization failed to identify any motifs for specific subcellular localization. This raises the possibility that Rpr localizes to the mitochondria through a novel mechanism. Thus, we decided to investigate Rpr localization by ectopic coexpression.
experiments in human BT549 cells, as well as in Drosophila S2R+ cells. We specifically followed the distribution of Rpr-HA, as well as XIAP, a human IAP member that is known to bind Rpr (Holley et al., 2002). Rpr-HA (Fig. 4 A) as well as GFP-fused XIAP (Fig. 4 B) was found to be spread diffusely throughout the cytoplasm. Similar experiments using Rpr-Myc and GFP-Rpr confirmed the broad distribution of Rpr in BT549 cells (unpublished data). In contrast, Hid, which has a mitochondrial targeting sequence, localizes exclusively to mitochondria and triggers GFP-XIAP translocation to this organelle (Fig. 4 C). This experiment confirms the ability of the IAP antagonist to interact with other proteins and recruit them to mitochondria. We also coexpressed GFP-Rpr and Hid in the BT549 cells and monitored any changes in the intracellular distribution of the proteins. Consistent with the ability of Rpr to bind Hid in vitro, the presence of Hid prompted GFP-Rpr distribution to change into a mitochondrial pattern (Fig. 4 D). The experiments above suggest that Rpr is not a mitochondrial protein, per se, but it is recruited to mitochondria by interaction with a mitochondrial-anchored protein. To further validate these results, we performed
nucleus in BT549 and S2R+ cells and triggers nuclear localization of Rpr, XIAP, and DIAP1 (Fig. S1), suggesting that indeed the observed mitochondrial localization of Rpr, XIAP, and DIAP1 is dependent on mitochondrial localization of Hid. Because Rpr and Hid overexpression induce cell death, we tested whether this might induce a change in the intracellular localization of these proteins. However, addition of the caspase inhibitor zVAD-FMK did not affect localization of Rpr or Hid (Fig. S2).

**Functional cooperativity between Rpr and Hid**

To test whether the physical interaction between Rpr and Hid has functional significance, we examined Rpr’s ability to kill cells in the absence of Hid. Although GMR->Rpr flies had rough eyes as expected, RNAi-mediated knockdown of Hid considerably suppressed this cell death phenotype of GMR->Rpr (Fig. 5 A). The cell death phenotype of GMR->Rpr is indeed caused by Rpr.

Figure 4. Rpr translocates to the mitochondria through physical interaction with Hid. (A) BT549 cells expressing Rpr-HA and GFP-XIAP. Rpr was stained with an anti-HA antibody. “Overlay” represents a composite image of Rpr (red), GFP-XIAP (green), and nuclei (DAPI, blue) staining. Bar, 20 µm. (B) BT549 cells transiently transfected with GFP-XIAP and mitochondrial RFP (mRFP) plasmids. “Overlay” indicates GFP-XIAP (green), mitochondria (red), and nuclei (blue). Bar, 20 µm. (C) BT549 cells cotransfected with Hid-HA and GFP-XIAP plasmids. “Overlay” indicates Hid (red), GFP-XIAP (green), and nuclei (blue). Bar, 20 µm. (D) BT549 cells cotransfected with GFP-Rpr and Hid-HA plasmids. “Overlay” shows Rpr (green), Hid (red), and nuclei (blue) staining. Bar, 20 µm. (E) S2R+ Drosophila cells transiently transfected with a mCherry-DIAP1 plasmid (left image) or with mCherry-DIAP1 and Hid-Myc plasmids (right image). Each image shows the overlay of DIAP1 (red) and nuclei (blue) staining. Bar, 5 µm. (F) S2R+ Drosophila cell, transiently transfected with a Rpr-HA plasmid, followed by immunostaining with anti-HA and anti-Cyt C antibodies. “Overlay” indicates Rpr (red), Cyt C (green), and nuclei (blue) staining. Bar, 5 µm. (G) S2R+ Drosophila cell cotransfected with Rpr-HA and Hid-Myc plasmids. Cells were immunostained with an anti-HA antibody and an anti-Myc antibody. “Overlay” represents Rpr (green), Hid (red), and nuclei (blue) staining. Bar, 5 µm.

a similar set of experiments in Drosophila S2R+ cells. As in BT549 cells, we observed that mCherry-DIAP1 is distributed evenly throughout the cytoplasm of S2R+, but after cotransfection with Hid, mCherry-DIAP1 is translocated to mitochondria in a Hid-like pattern (Fig. 4 E). When Rpr was expressed transiently in S2R+ cells, it shows an occasional punctate staining that is only coincidental with cytochrome c (Cyt C) staining (Fig. 4 F). However, after cotransfection with Hid, Rpr’s colocalization with Hid becomes obvious (Fig. 4 G). In sum, our experiments suggest that Rpr is a soluble protein that displays a diffuse distribution throughout the cell, and coexpression with Hid leads to Rpr relocalization to the mitochondria. Additionally, these experiments underline Hid’s ability to recruit DIAP1 and its human homologue XIAP to the mitochondrial membrane. To test whether the recruitment of Rpr, XIAP, and DIAP1 is indeed dependent on Hid’s MTS, we have performed coexpression experiments with HidΔMTS and Rpr, XIAP, or DIAP1. HidΔMTS localizes to the nucleus in BT549 and S2R+ cells and triggers nuclear localization of Rpr, XIAP, and DIAP1 (Fig. S1), suggesting that indeed the observed mitochondrial localization of Rpr, XIAP, and DIAP1 is dependent on mitochondrial localization of Hid. Because Rpr and Hid overexpression induce cell death, we tested whether this might induce a change in the intracellular localization of these proteins. However, addition of the caspase inhibitor zVAD-FMK did not affect localization of Rpr or Hid (Fig. S2).
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Overexpression and could be suppressed by Rpr RNAi (Fig. 5 A). Furthermore, we could show that the used Hid RNAi line is effective at knocking down endogenous Hid because it rescues the GMR>Hid-induced eye ablation (Fig. 5 B). The effect of Hid knockdown on Rpr-induced cell death was also observed in cell culture. When Drosophila S2R+ cells were transiently transfected with a Rpr-HA plasmid in the presence of a 21-bp Hid RNA duplex (siRNA), the number of Rpr-HA–positive (5.65%), as identified by anti-HA immunostaining, were almost fourfold greater than those without Hid siRNA treatment (1.46%; Fig. 5, C and D). We interpret that depleting Hid mRNA in S2R+ cells gives these cells a better protection against Rpr-induced cell death, allowing a larger population of Rpr-positive cells to survive. The efficiency of the Hid siRNA was demonstrated by the ability to decrease the level of Hid-Myc in S2R+ cells in transient transfection experiments (Fig. 5 D). Hid's role in Rpr-induced cell death was further validated in the heterozygous background of hid loss-of-function mutant alleles, hid A206 and hid WR+X1 (Abbott and Lengyel, 1991; Grether et al., 1995), which suppressed the degree of Rpr-induced eye ablation (Fig. 5 E). Conversely, Hid-induced cell death in Drosophila eyes is suppressed to a degree by Rpr RNAi (Fig. S3 A). These findings are in contrast to the widely recognized view that the three Drosophila IAP antagonists Rpr, Hid, and Grim induce
reconstitute the covalent coupling of one Ub molecule on UbcD1-conjugating enzyme, in an Uba1- and Mg\(^{2+}\)-ATP–dependent fashion (Fig. 6 B). Furthermore, by using reducing agents to break down the E2-Ub thiolesters, we confirmed the presence of the UbcD1-Ub adduct (unpublished data). We next examined DIAP1 ubiquitination in the presence of Mg\(^{2+}\)-ATP, Rpr, and/or Hid\(^{6}\)MTS. Although DIAP1 does not self-ubiquitinate in the presence of Mg\(^{2+}\)-ATP (Fig. 6 C), a dramatic transfer of ubiquitin to DIAP1 could be observed when the reaction is supplemented with Rpr. Thus, we have fully reconstituted in vitro a DIAP1 auto-ubiquitinating complex from Drosophila. Because IAP antagonists bind DIAP1 with conserved motifs it is often assumed that the mechanism of DIAP1 inactivation should also be conserved. However, when Hid\(^{6}\)MTS was added in the DIAP1 ubiquitination assay instead of Rpr, no DIAP1 auto-ubiquitination could be observed. Despite good solubility, Hid\(^{6}\)MTS did not stimulate DIAP1 degradation. When Hid\(^{6}\)MTS and Rpr were added together to the DIAP1 ubiquitination assay, Hid\(^{6}\)MTS did not reconstitute the covalent coupling of one Ub molecule on UbcD1-conjugating enzyme, in an Uba1- and Mg\(^{2+}\)-ATP–dependent fashion (Fig. 6 B). Furthermore, by using reducing agents to break down the E2-Ub thiolesters, we confirmed the presence of the UbcD1-Ub adduct (unpublished data). We next examined DIAP1 ubiquitination in the presence of Mg\(^{2+}\)-ATP, Rpr, and/or Hid\(^{6}\)MTS. Although DIAP1 does not self-ubiquitinate in the presence of Mg\(^{2+}\)-ATP (Fig. 6 C), a dramatic transfer of ubiquitin to DIAP1 could be observed when the reaction is supplemented with Rpr. Thus, we have fully reconstituted in vitro a DIAP1 auto-ubiquitinating complex from Drosophila. Because IAP antagonists bind DIAP1 with conserved motifs it is often assumed that the mechanism of DIAP1 inactivation should also be conserved. However, when Hid\(^{6}\)MTS was added in the DIAP1 ubiquitination assay instead of Rpr, no DIAP1 auto-ubiquitination could be observed. Despite good solubility, Hid\(^{6}\)MTS did not stimulate DIAP1 degradation. When Hid\(^{6}\)MTS and Rpr were added together to the DIAP1 ubiquitination assay, Hid\(^{6}\)MTS did not

**Figure 6.** DIAP1 auto-ubiquitination and interaction with Rpr and Hid. (A) SDS-PAGE gel showing E1 ubiquitin-activating enzyme Uba1 (Uba1-GST), E2 ubiquitin-conjugating enzyme UbcD1 [6His-UbcD1], 6His-ubiquitin (Ub), E3 ubiquitin ligase DIAP1 [6His-Flag-DIAP1], Rpr-His6, and Hid\(^{6}\)MTS-His6, used in ubiquitination assays. Purification tags are not shown in the figure labeling. (B) In vitro coupling of Ub on UbcD1 (E2) in the absence (lane 1) or presence of Mg\(^{2+}\)-ATP (lane 2). UbcD1-Ub adduct was detected by Coomassie staining. (C) In vitro DIAP1 auto-ubiquitination. Ubiquitination reactions containing E1, E2, UbcD1, and Flag-DIAP1, in the absence of Mg\(^{2+}\)-ATP (lane 1) or in the presence of Mg\(^{2+}\)-ATP (lane 2). The reaction was supplemented additionally with Rpr (lane 3), Hid\(^{6}\)MTS (lane 4), or both (lane 5). Flag-DIAP1 was immunoprecipitated with anti-FLAG resin. Polyubiquitination species were detected in Western blot with an anti-ubiquitin antibody. (D) Coomassie-stained SDS-PAGE gel, showing the concomitant precipitation of Flag-DIAP1 with Rpr and Hid\(^{6}\)MTS. “Input” shows the amount of Flag-DIAP1 (lane 1), Rpr (lane 2), or Hid\(^{6}\)MTS (lane 3) used for co-immunoprecipitation. “IP:Flag” shows the amount of Flag-DIAP1 recovered by the anti-FLAG resin. Lane 5 shows the co-immunoprecipitation of Rpr with Flag-DIAP1. Lane 6 shows the coimmunoprecipitation of Hid\(^{6}\)MTS with Flag-DIAP1. Lane 7 shows the coimmunoprecipitation of Hid\(^{6}\)MTS and Rpr with Flag-DIAP1.

Rpr can stimulate Diap1 self-conjugation in a purified in vitro system

Why does Rpr kill better in the presence of Hid? To answer this question we started from the premise that both IAP antagonists act to stimulate DIAP1 auto-ubiquitination (Hays et al., 2002; Ryoo et al., 2002; Yoo et al., 2002). To this end, we purified all components involved in DIAP1 auto-ubiquitination, namely Uba1 (E1), UbcD1 (E2), ubiquitin (Ub), DIAP1, Rpr, and Hid\(^{6}\)MTS from E. coli (Fig. 6 A). Hid\(^{6}\)MTS entails residues 1–386. The last 24 amino acids (387–410) of Hid, which constitute the membrane-inserted mitochondrial targeting sequence, were deleted to produce protein soluble for biochemical assays. When designing this construct, we inspected Hid secondary structure to avoid terminating the protein inside a secondary structure element (unpublished data). The proteins are active, as we could
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The ability of full-length Hid or Hid\textsuperscript{MTS} to induce DIAP1\textsuperscript{R} degradation in HEK293 cells. Unlike Rpr (Fig. 2 F), full-length Hid and Hid\textsuperscript{MTS} do not induce DIAP1\textsuperscript{R} degradation (Fig. S3 C).

These observations and others suggest that most probably Hid does not induce DIAP1 ubiquitination directly.

Artificial targeting of Rpr to mitochondria stimulates its activity, and Hid promotes Rpr stability

Because Hid\textsuperscript{MTS} cannot ubiquitinate DIAP1 in vitro and it does not enhance Rpr-mediated DIAP1 ubiquitination, it is possible that Hid stimulates Rpr’s activity by another mechanism. A possible scenario is that Hid recruits Rpr to the mitochondrial membrane, where Rpr is more effective in DIAP1 degradation. To test whether Rpr is more effective when present at the mitochondrial membrane, we artificially targeted Rpr at the mitochondrial membrane by appending to it the mitochondrial targeting sequence of Hid. Rpr-MTS and F34L35A-MTS constructs were used to generate transgenic animals with the constructs inserted in the same genomic location. Rpr-MTS and F34L35A-MTS were expressed in Drosophila eyes using not enhance Rpr-dependent DIAP1 ubiquitination. The inability of Hid\textsuperscript{MTS} to induce DIAP1 ubiquitination could be a result of the following reasons. First, it is possible that recombinant Hid\textsuperscript{MTS} does not reflect endogenous Hid function, perhaps due to the deletion of its C-terminal hydrophobic region. A second possibility is that Rpr and Hid have different mechanisms of DIAP1 inactivation and only Rpr induces DIAP1 ubiquitination, Hid having a different role in DIAP1 inactivation. In an attempt to address these possibilities, we examined the interactions between DIAP1, Rpr, and Hid using the purified proteins used in the ubiquitination assay. Under these conditions, Hid\textsuperscript{MTS} was able to bind DIAP1 at a roughly equimolar ratio as judged by band intensity on SDS-PAGE gel, despite Hid\textsuperscript{MTS}’s inability to stimulate DIAP1 ubiquitination (Fig. 6 D). Furthermore, we have examined the ability of Hid\textsuperscript{MTS} to form oligomers by formaldehyde cross-linking experiments. Purified Hid\textsuperscript{MTS} and Rpr appear to form oligomers under these conditions (Fig. S3 B). In addition, the interaction between Rpr and Hid\textsuperscript{MTS} was already shown in Fig. 3 B. These experiments indicate that deletion of Hid’s MTS does not block its ability to oligomerize or interact with Rpr and DIAP1. We have next examined the ability of full-length Hid or Hid\textsuperscript{MTS} to induce DIAP1\textsuperscript{R} degradation in HEK293 cells. Unlike Rpr (Fig. 2 F), full-length Hid and Hid\textsuperscript{MTS} do not induce DIAP1\textsuperscript{R} degradation (Fig. S3 C).

These observations and others suggest that most probably Hid does not induce DIAP1 ubiquitination directly.
the GMR>Gal/UAS system. Surprisingly, both constructs were lethal at late pupal stages at 21°C. When dissected from the pupal cases, Rpr-MTS fly eyes were reduced to a black spot and the F35AL35A-MTS fly eyes were severely affected (Fig. 7A), underscoring the importance of Rpr’s mitochondrial targeting for its killing activity. The GH3 F34AL35A mutant, which is unable to promote eye ablation and is deficient in Hid binding, induces significant eye ablation when artificially targeted to the mitochondria, comparable to a wild-type Rpr nontargeted to the mitochondria. Next, we asked whether the eye ablation phenotype is indeed due to apoptosis. We have isolated third instar larvae eye-antennal discs expressing Rpr-MTS and F34L35-MTS in the GMR region and immunostained them with an antibody against active caspases. Consistent with the observed rough eye phenotypes, both Rpr-MTS and F34L35-MTS showed significant caspase staining in the GMR region (Fig. 7 B). Next, we examined the ability of Rpr-MTS and F34L35-MTS to induce DIAP1 degradation. After coexpression in HEK293 cells, both Rpr-MTS and F34L35-MTS induced a decrease in DIAP1ΔR peptide (Fig. 7 C). Expression of Rpr-MTS and F34L35-MTS in HEK293 cells was driven by cotransfection of a cmv-Gal4 driver and UAS:Rpr-MTS or UAS:F34L35-MTS constructs and was lower than that achieved with mammalian expression vectors. This might explain the incomplete DIAP1ΔR degradation. In an effort to understand how Hid enhances Rpr’s cell-killing activity, we next compared Rpr protein level in the presence or absence of Hid. When expressed in HEK293 cells, we found that Rpr’s level is in fact much higher in extracts derived from rpr and hid co-transfected cells (Fig. 7 D). On the other hand, the level of Hid did not change significantly in the presence of Rpr. As an expression, protein extraction, and loading control we have used YFP-Mem (a fluorescent marker for cell membranes), which indicates that Rpr level increase is indeed dependent on Hid’s presence. These results suggest a model where Rpr is targeted to the mitochondria by interaction with Hid, and in such a complex that potentially includes other factors, Rpr is protected against degradation (Fig. 7 E).

Discussion

In this study we show that IAP antagonists undergo self-association and hetero-association that is essential for their full killing activity. Specifically, the physical association between Rpr, Hid, and Grim involves the central helical domain of Rpr. Disrupting this protein–protein interface leads to a significant loss of Rpr’s ability to induce cell death in vivo. The importance of Rpr self-association was revealed by generating enforced Rpr dimers in which the central helical domain of this protein is replaced by defined dimerization motifs. These experiments revealed that enforced parallel, but not anti-parallel dimerization of Rpr (RprLZ) can induce cell death very efficiently in transgenic Drosophila. The resulting cell death occurred by apoptosis and was rescued by the overexpression of the caspase inhibitor p35, or through Rpr-insensitive diap1 alleles. Furthermore, mutants that inhibit the self-association of Rpr have reduced pro-apoptotic activity, providing independent support for the importance of Rpr multimerization. Because an anti-parallel Rpr dimer (RprProP) was not efficiently inducing cell death in transgenic animals, it appears that the IBM motifs of multimeric Rpr have to be in a specific conformation, or at least in close proximity for efficient DIAP1 inactivation. This may occur, for example, by engaging both BIR domains of one DIAP1 molecule in a similar fashion to how SMAC can engage XIAP (Huang et al., 2003).

We also report the association of Rpr with the other IAP antagonists Grim and Hid. Hid is the only IAP antagonist that has a defined mitochondrial targeting sequence at its C terminus and is targeted to the mitochondria by itself; therefore, we focused particularly on the interaction between Rpr and Hid. Consistent with previous reports, we find that Hid consistently localizes to the mitochondria in both human and Drosophila cells. Although it has been previously reported that Rpr localizes to the mitochondria through the GH3–lipid interaction (Olson et al., 2003a; Freel et al., 2008), our results support an alternative view that Rpr’s ability to translocate to the mitochondria is an indirect consequence of associating with Hid. Specifically, in support of our model, we show that Rpr is uniformly distributed in cells when transfected alone in heterologous cells, translocating to the mitochondria only when cotransfected with Hid. We further show that the GH3 mutant F34AL35A, unlike wild-type Rpr, does not co-immunoprecipitate with Hid. This is in agreement with previous observations that a GH3 mutant failed to localize to the mitochondria in Drosophila S2 cells (Olson et al., 2003a).

Rpr induces ubiquitination of DIAP1 in vitro and in HEK293 cells. Unlike Rpr, Hid is not able to perform this function. Thus, the significance of Rpr–Hid interaction might be to bring Rpr at the mitochondrial surface to degrade DIAP1. Although both Rpr and Hid belong to the IAP antagonists family, share a conserved IBM motif, bind DIAP1, and induce cell death, their role in induction of cell death seems to be distinct. In many paradigms Hid appears to be a more potent inducer of cell death than Rpr. It is possible that the primary role of Hid is to assemble a complex at the mitochondrial membrane that recruits Rpr as one the players. The role of Rpr in this complex is to induce DIAP1 ubiquitination. Inability of Hid itself to induce DIAP1 degradation might be related to its larger size (410 amino acids) as compared with Rpr (64 amino acids) or even Grim (138 amino acids). Potentially, the bulkier Hid might interfere with conformational changes in DIAP1 or with the ubiquitin-related transfer process.

In addition, we provide evidence that Rpr is more potent at inducing apoptosis when present at the mitochondrial membrane. When Rpr was fused to the mitochondrial targeting sequence from Hid and expressed in Drosophila eyes, we observed strong cell killing and pupal lethality. Flies dissected from the pupal cases show severely ablated eyes that are reduced to black spots. Even the inactive GH3 mutant F34AL35A, when artificially targeted to the mitochondria using the Hid MTS, induces significant eye ablation. Therefore, Rpr is more potent when present at the mitochondrial membrane. We consider two possible explanations for this enhanced pro-apoptotic activity: First, Rpr may be more active at the mitochondrial surface because of increased protein stability. Consistent with this idea, cytoplasmic Rpr is not very stable (Olson et al., 2003b) and we find that Rpr accumulates to higher protein levels when the presence of...
Hid permits mitochondrial localization. The resulting high local concentration of Rpr may be critical for DIAP1 ubiquitination. As predicted by this model, we find that Rpr-induced cell death is less efficient when Hid is depleted by RNA knockdown. Our model is also in agreement with several previous observations. For example, it has been reported that Rpr and Hid localize to mitochondria and can induce changes of the mitochondrial ultrastructure (Abdelwahid et al., 2007). This study also showed that inhibition of Rpr localization to mitochondria significantly inhibits cell killing, and that Rpr and Hid act in concert with caspases to promote mitochondrial disruption and Cyt C release. In addition, overexpression of both rpr and hid is required to induce cell death in midline cells of the nervous system, and neither of them kills well individually (Zhou et al., 1997). This is consistent with the observation that more than one IAP antagonist is expressed and they act synergistically in the dying midline glia cells (Sonnenfeld and Jacobs, 1995; Zhou et al., 1995; Dong and Jacobs, 1997; Wing et al., 1998; Bergmann et al., 2002). Finally, Drosophila salivary gland cell death is preceded by the expression of both rpr and hid, and RNAi knockdown of hid alone is sufficient to block the death of these cells (Jiang et al., 2000; Yin and Thummel, 2004). The second, and not mutually exclusive explanation is that Rpr may be more active at the mitochondria because of local concentration of apoptosis regulators that operate at this surface. It has been previously shown that Dronc and active Drcare present at the mitochondrial membrane (Dorstyn et al., 2004). The second, and not mutually exclusive explanation is that Rpr may be more active at the mitochondria because of local concentration of apoptosis regulators that operate at this surface. It has been previously shown that Dronc and active Drcare present at the mitochondrial membrane (Dorstyn et al., 2004). The second, and not mutually exclusive explanation is that Rpr may be more active at the mitochondria because of local concentration of apoptosis regulators that operate at this surface. It has been previously shown that Dronc and active Drcare present at the mitochondrial membrane (Dorstyn et al., 2004). The second, and not mutually exclusive explanation is that Rpr may be more active at the mitochondria because of local concentration of apoptosis regulators that operate at this surface.

Materials and methods

Plasmids for mammalian, insect, and bacterial expression

With the exception of GFP::Rpr, all tagged Rpr constructs used in this study had the epitopes fused to the C terminus of the Rpr coding sequence. For expression experiments in mammalian cells, Rpr-HA, Rpr-Myc, GFP::Rpr, Hid-HA, Hid-Flag, HidMTS-Myc (residues 1–386), GFP::XAP, RprLeHA, RprProP-Flag, and DIAP1-Flag (residues 1–292) were cloned into pcDNA3.1+Puro vector (Thomas et al., 2002). pEYFP-Mem was purchased from Takara Bio Inc. For expression in Drosophila cell culture, mCherryDIAP1, Hid-Myc, and HidMTS-Myc (residues 1–386) were cloned in pEY3-3 vector (EMD). For the expression of Rpr in fly cells, we generated Rpr-HA pUAST constructs where Rpr-HA sequence was flanked by Rpr’s 5’ and 3’ UTRs. We also made similar constructs with atto and plxP elements for targeted insertion into specific sites in the genome (Groth et al., 2004; Oberstein et al., 2005). The ploxP/UAST::Rpr-HA plasmid and its mutant variants were targeted to the [21L388, 13399006] site on the second chromosome (line A11; Bestgene, Inc.) pUASTattB::Rpr-MTS and pUASTattB::F34AL35A-MTS were targeted to a locus on the second chromosome (line 244B1, genotype: M[3xP3-RFP.cat];Z2H2A [with M[vas-n-3.Dm];Z2H2A]; Bestgene, Inc.). pUAST::RprLZ-FA construct was created by replacement of Rpr amino acids 10–48 in pUAST::Rpr-HA vector with the LC2-encoding DNA fragment, flanked by Agel and SpeI restriction sites. This vector was achieved by the insertion of two unique restriction sites, Agel and SpeI, at the above-mentioned positions in pUAST::Rpr-HA. pUAST::RprPro-PHA was generated similarly to RprLZ by insertion of the ProP PCR fragment, between the Agel and SpeI sites of the modified pUAST::Rpr-HA vector. pUAST::RprLZ-HA and pUAST::RprPro-PHA were used to generate fly transgenes (Genetic Services, Inc.). For protein production in E. coli, the following vectors were created: pETDuet:Rpr-loxI-GST, pET28b:ubiquitin, pET14b-Flag-DIAP1, pET2a-FLAG and pET11a:HidMTS, pET14b-UbcD1 (Royo et al., 2002), and pET3a::Rpr-GST (Royo et al., 2002). pET11a:Rpr, pET11a:Hid, pET12a:Grin, and pETDuet-Ski were used for expression in the in vitro transcription translation reactions. pET21a::Rpr constructs produce Rpr with a C-terminal MGGMGHHHHHHHHHHH tag. This construct was used to generate Rpr point mutants.

Protein expression and purification

For protein production we used E. coli BL21DE3 strain, transformed with appropriate plasmids. Protein production was induced with 500 µM IPTG at 25°C overnight. Purification of GST, Rpr-GST, GST-Hid, or Uba1-GST was performed following a previously described protocol (Carrington et al., 2000; Sandu et al., 2006). Purification of 6His-UbcD1 (E2), 6His-Flag-DIAP1 (E3), Rpr-His6, HidMTS-His6, and 6His-ubiquitin was performed using an adapted protocol used for Ulp1 purification (Mossessova and Lima, 2000). In brief, the pellet from one liter E. coli culture was resuspended in 70 ml of lysis buffer (50 mM Tris, pH 8.0, 350 mM NaCl, 1 mM BME, 0.2% Igepal, and 10 mM imidazole) and disrupted by sonication. After sonication the cell extract was cleared by centrifugation at 14,000 g for 15 min. The 6His-tagged protein was bound to Talon Metal Affinity resin (Takara Bio Inc.) and washed with 35 ml of lysis buffer, followed by 35 ml of washing buffer (50 mM Tris, pH 8.0, 350 mM NaCl, 1 mM BME, 0.2% Igepal, and 70 mM imidazole). After washing, the protein was eluted with 20 ml elution buffer (50 mM Tris, pH 8.0, 350 mM NaCl, 1 mM BME, 0.2% Igepal, and 50 mM imidazole). After elution, the purified proteins were concentrated using Amicon Centricon (Millipore) and flash frozen in liquid nitrogen after addition of 10% glycerol.

Protein–protein interaction studies

Interaction of Rpr-GST with 35S-Rpr, 35S-Hid, 35S-Grin, and 35S-Ski was investigated by pull-down experiments using a protocol described previously (Sandu et al., 2006). In brief, 20 µl of a Rpr-GST bead slurry were mixed in 200 µl binding buffer (50 mM Hepes, pH 7.4, 50 mM NaCl, 0.1% NP-40, and 10% glycerol) with 10 µl of the 35S-labeled Rpr, Hid, Grin, or Ski. After 2 h at 4°C, the beads were harvested by centrifugation and washed three times with 500 µl of binding buffer for 10 min at 4°C. The beads were then eluted with 15 µl SDS-PAGE loading dye at 95°C, separated on SDS-PAGE gels, and visualized by phosphorimaging of the dried gel. Interaction of Rpr with GST-Hid or GST or the interaction between Ub, Rpr, or HidMTS with Rpr-GST was tested in an alternative pull-down experiment. Approximately 5 µg GST-Hid, GST, or Rpr-GST linked on agarose beads were incubated with 5 µg of either Rpr-His6 (Rpr), His6-ubiquitin (Ub), or HidMTS-His6 (HidMTS) in 250 µl binding buffer (50 mM Hepes, pH 7.4, 50 mM NaCl, 0.1% NP-40, and 10% glycerol) and rotated overnight for 2 h at 4°C. The beads were washed three times with 500 µl binding buffer and denatured in SDS-sample buffer at 95°C. The samples were separated in SDS-PAGE gels, followed by Coomassie staining.

Cell culture, immunoprecipitation, immunostaining, and Western blotting

Human HEK293 and B1T49 or Drosophila S2R+ cell lines were used for experiments involving expression, immunolocalization, and immunoprecipitation studies. For testing the ability of Rpr-HA, RprLZ-HA, Hid-HA, or HidMTS-Myc to induce DIAP1-Flag degradation, HEK293 cells were transfected with either DIAP1-Flag construct or cotransfected in equal ratios with the DIAP1-Flag construct and each one of the above-mentioned constructs. After 16 h after transfection, cells were lysed in lysis buffer (50 mM Tris, pH 8.0, 100 mM NaCl, and 0.5% NP-40) supplemented with protease inhibitor cocktail (Roche). After centrifugation at 14,000 g, cell extracts (typically 25 µg) were separated by SDS-PAGE, blotted on nitrocellulose membranes, and proteins were detected with specific antibodies. Alternatively, HEK293 cells were cotransfected with pcDNA3.1+ProP-DIAP1-Flag, pCMV-Gal4 [a gift from Brianne Font, The Rockefeller University, New York, NY], and either pUAST::Rpr-HA-MTS or pUAST::F34L35A-HA-MTS.
of a few structural models of the Rpr helical domain. Protein models were generated using the Protein Homology/AnalogY Recognition Engine (PHyre; Kelley and Sternberg, 2009) and analyzed using Pymol. Residues with bulky, large side chains, oriented outwards, were chosen for mutagenesis. These residues were either hydrophobic or charged. Assuming that the surface of interaction is extensive, we chose to mutate two residues at a time. The two residues are oriented on the same side of the helix, would be part of the same interaction patch of the helix surface (charged or hydrophobic), and are positioned each on adjacent helix turns. The residues in the mutagenesis were spaced to cover the entire helical region.

Online supplemental material

Fig. S1 shows the immunolocalization experiments of Rpr-HA, GFP-XIAP, or mCherryDIAP1 with HidMTS-Myc. Fig. S2 shows the effect of caspase inhibitor zVAD-FMK on intracellular localization of Rpr-HA and Hid-HA. Fig. S3 shows the effect of Rpr mRNA knockdown on Hid-induced cell death in Drosophila eyes (A); shows the ability of purified Rpr and HidMTS to form oligomeric species after formaldehyde cross-linking (B); and the effect of Hid-HA or HidMTS-Myc on DIAP1 ΔK Flag degradation in HEK293 cells. Online supplemental material is available at: http://www.jcb.org/cgi/content/full/jcb.201004086/DC1.

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Drosophila...


Immunolocalization of HidΔMTS and its effects on Rpr, XIAP, and DIAP1 localization.

(A) HidΔMTS localizes to the nucleus in BT549 cells. The cells were transfected with a HidΔMTS-Myc construct, and immunostained with an anti-Myc antibody coupled to a FITC-labeled secondary antibody. HidΔMTS is shown in green, nuclei are shown in blue. Bar, 20 µm.

(B) Rpr colocalizes with HidΔMTS in the nucleus of BT549 cells. The cells were cotransfected with HidΔMTS-Myc and Rpr-HA constructs, and immunostained with an anti-Myc antibody coupled to a FITC-labeled secondary and an anti-HA antibody coupled with an Alexa 546–labeled secondary antibody. HidΔMTS is shown in green, Rpr is shown red. Bar, 20 µm.

(C) GST-XIAP is targeted to the nucleus after cotransfection with HidΔMTS. BT549 cells cotransfected with HidΔMTS-Myc and GFP-XIAP constructs. HidΔMTS-Myc was stained with an anti-Myc antibody coupled with a Cy3-labeled secondary antibody. GFP-XIAP cellular distribution was visualized by GFP fluorescence. Nuclei were labeled by DAPI. Bar, 20 µm.

(D) HidΔMTS localizes to the nucleus in S2R+ cells. S2R+ cells were transfected with HidΔMTS-Myc and mtGFP constructs. HidΔMTS nuclear localization was revealed by immunostaining with an anti-Myc antibody coupled to a Cy3-labeled secondary antibody. Mitochondria were visualized by GFP fluorescence. Nuclei were labeled by DAPI. Bar, 5 µm.

(E) mCherryDIAP1 colocalizes with HidΔMTS to the nucleus. S2R+ cells were cotransfected with HidΔMTS-Myc and mCherryDIAP1 constructs. HidΔMTS localization was revealed by immunostaining with an anti-Myc antibody coupled to a FITC-labeled secondary antibody; DIAP1 localization was revealed by mCherry fluorescence. DAPI was used to label nuclei. Bar, 5 µm.
Figure S2. Effect of zVAD-FMK on Rpr or Hid intracellular localization. BT549 cells were transfected with a Rpr-HA construct in the absence (A) or presence (B) of 20 μM zVAD-FMK. 16 h after transfection, the cells were fixed and stained with an anti-HA antibody coupled with an Alexa 488-labeled secondary antibody. DAPI was used for nuclear staining. Rpr intracellular localization is not affected by addition of zVAD-FMK. In the following two panels, BT549 cells were transfected with a Hid-HA construct in the absence (C) or presence (D) of 20 μM zVAD-FMK. 16 h after transfection, the cells were fixed and stained with an anti-HA antibody coupled with an Alexa 488-labeled secondary antibody. DAPI was used for nuclear staining. Hid mitochondrial localization is not affected by addition of zVAD-FMK. Bar, 20 μm.
Figure S3. Hid’s ability to induce cell death, ubiquitinate DIAP1, and form oligomeric species. (A) Rpr mRNA knockdown rescues to degree Hid-induced rough eye in Drosophila. (Left) A rough eye phenotype caused by overexpression of Hid in the eye. Genotype: GMR-Gal4/+; Sco/GMR-Hid; Sb/TM6B. (Right) Partial rescue of the Hid-induced rough eye phenotype by Rpr mRNA knockdown. Genotype: GMR-Gal4/+; Sco/GMR-Hid; Sb/UAS:Rpr RNAi. Crosses were incubated at 21°C. (B) Formaldehyde cross-linking reveals formation of oligomeric species by Rpr and HidΔMTS. (Left) Rpr forms oligomeric species after formaldehyde cross-linking. 5 µg purified Rpr protein was incubated for 10 min at room temperature in the absence or presence of 0.1% or 1.0% formaldehyde. The samples were neutralized with 3 M Tris, pH 8.0, and separated on SDS-PAGE. Appearance of dimeric species is visible after Coomassie staining. (Right) Formation of oligomeric species by HidΔMTS after formaldehyde cross-linking. 5 µg purified HidΔMTS was incubated at room temperature in the absence or presence of 0.1% or 1.0% formaldehyde. After incubation the reaction was neutralized by addition of 3 M Tris, pH 8.0, and samples prepared and separated by SDS-PAGE. Appearance of dimeric and oligomeric species is visible after Coomassie staining. (C) Hid full-length or HidΔMTS cannot effectively induce DIAP1ΔR degradation in HEK293 cells. (Left) Western blots on cell extracts derived from HEK293 cells, transfected with either DIAP1ΔR-Flag or DIAP1ΔR-Flag and Hid-HA constructs. Expression of Hid full-length does not induce degradation of DIAP1ΔR. Actin was used as a loading control. Expression level of DIAP1ΔR, Hid, and Actin was assessed with anti-Flag, anti-HA, and anti-Actin antibodies. (Right) Western blots on cell extracts derived from HEK293 cells, transfected with either DIAP1ΔR-Flag or DIAP1ΔR-Flag and HidΔMTS-Myc. Expression of HidΔMTS does not cause effective DIAP1ΔR degradation. Actin was used as a loading control. Expression level of DIAP1ΔR, HidΔMTS, and Actin was assessed with anti-Flag, anti-Myc, and anti-Actin antibodies.